IHE Patient Care Coordination (PCC)

Technical Framework
Volume 2
Revision 5.0

Final Text
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1 Preface to Volume 2

1.1 Intended Audience

The intended audience of this document is:

- Technical staff of vendors planning to participate in the IHE initiative
- IT departments of healthcare institutions
- Experts involved in standards development
- Anyone interested in the technical aspects of integrating healthcare information systems

1.2 Related Information for the Reader

The reader of volume 2 should read or be familiar with the following documents:

- HL7 Clinical Document Architecture Release 2: Section 1, CDA Overview.
- Care Record Summary – Implementation Guide for CDA Release 2 (US Realm): Section 1
- Presentations from IHE Workshop: Effective Integration of the Enterprise and the Health System - June 28–29, 2005: http://www.ihe.net/Participation/workshop_2005.cfm, June 2005:
- For a RHIO-3.ppt Leveraging IHE to Build RHIO Interoperability
- Cross-Enterprise Document Sharing (XDS)
- Notification of Document Availability (NAV)
- Use Cases for Medical Summaries
- Ovrw.ppt Patient Care Coordination - Overview of Profiles
1.2.1 How this Document is Organized

Section 1 is the preface, describing the intended audience, related resources, and organizations and conventions used within this document.

Section 2 provides an overview of the concepts of IHE actors and transactions used in IHE to define the functional components of a distributed healthcare environment.

Section 3 defines transactions in detail, specifying the roles for each actor, the standards employed, the information exchanged, and in some cases, implementation options for the transaction.

Section 4 defines a set of payload bindings with transactions.

Section 5 defines the content modules that may be used in transactions.

1.2.2 Conventions Used in this Volume

This document has adopted the following conventions for representing the framework concepts and specifying how the standards upon which the IHE Technical Framework is based should be applied.

1.2.2.1 The Generic IHE Transaction Model

Transaction descriptions are provided in section 4. In each transaction description, the actors, the roles they play, and the transactions between them are presented as use cases.

The generic IHE transaction description includes the following components:

- Scope: a brief description of the transaction.
- Use case roles: textual definitions of the actors and their roles, with a simple diagram relating them, e.g.:

```
Figure 1.2-1 Use Case Role Diagram
```

- Referenced Standards: the standards (stating the specific parts, chapters or sections thereof) to be used for the transaction.
- Interaction Diagram: a graphical depiction of the actors and transactions, with related processing within an actor shown as a rectangle and time progressing downward, similar to:

- **Message definitions**: descriptions of each message involved in the transaction, the events that trigger the message, its semantics, and the actions that the message triggers in the receiver.

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### 1.4 How to Contact Us

IHE Sponsors welcome comments on this document and the IHE initiative. They should be directed to the discussion server at [http://forums.rsna.org](http://forums.rsna.org) or to:

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Senior Director Informatics  
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Chicago, IL 60611  
Email: [ihe@himss.org](mailto:ihe@himss.org)
2 Introduction

This document, the IHE Patient Care Coordination Technical Framework (PCC TF), defines specific implementations of established standards. These are intended to achieve integration goals that promote appropriate exchange of medical information to coordinate the optimal patient care among care providers in different care settings. It is expanded annually, after a period of public review, and maintained regularly through the identification and correction of errata. The latest version of the document is always available via the Internet at http://www.ihe.net/Technical_Framework/index.cfm, where the technical framework volumes specific to the various healthcare domains addressed by IHE may be found.

The IHE Patient Care Coordination Technical Framework identifies a subset of the functional components of the healthcare enterprises and health information networks, called IHE actors, and specifies their interactions in terms of a set of coordinated, standards-based transactions.

The other domains within the IHE initiative also produce Technical Frameworks within their respective areas that together form the IHE Technical Framework. Currently, the following IHE Technical Framework(s) are available:

- IHE IT Infrastructure Technical Framework
- IHE Cardiology Technical Framework
- IHE Laboratory Technical framework
- IHE Radiology Technical Framework
- IHE Patient Care Coordination Technical Framework

Where applicable, references are made to other technical frameworks. For the conventions on referencing other frameworks, see the preface of this volume.

2.1 Relationship to Standards

The IHE Technical Framework identifies functional components of a distributed healthcare environment (referred to as IHE actors), solely from the point of view of their interactions in the healthcare enterprise. At its current level of development, it defines a coordinated set of transactions based on standards (such as HL7, IETF, ASTM, DICOM, ISO, OASIS, etc.) in order to accomplish a particular use case. As the scope of the IHE initiative expands, transactions based on other standards may be included as required.

Each transaction may have as its payload one or more forms of content, as well as specific metadata describing that content within the transaction. The specification of the payload and metadata about it are the components of a Content Integration Profile. The payload is specified in a Content Module, and the impacts of any particular payload on a transaction are described within a content binding. The payloads of each transaction are also based on standards (such as HL7, IETF, ASTM, DICOM, ISO, OASIS, etc.), again, in order to meet the needs of a specific use case.

In some cases, IHE recommends selection of specific options supported by these standards. However, IHE does not introduce technical choices that contradict
conformance to these standards. If errors in or extensions to existing standards are identified, IHE's policy is to report them to the appropriate standards bodies for resolution within their conformance and standards evolution strategy.

300 IHE is therefore an implementation framework, not a standard. Conformance claims for products must still be made in direct reference to specific standards. In addition, vendors who have implemented IHE integration capabilities in their products may publish IHE Integration Statements to communicate their products' capabilities. Vendors publishing IHE Integration Statements accept full responsibility for their content. By comparing the IHE Integration Statements from different products, a user familiar with the IHE concepts of actors and integration profiles can determine the level of integration between them. See PCC TF-1: Appendix C for the format of IHE Integration Statements.

2.2 Relationship to Product Implementations

The IHE actors and transactions described in the IHE Technical Framework are abstractions of the real-world healthcare information system environment. While some of the transactions are traditionally performed by specific product categories (e.g. HIS, Clinical Data Repository, Electronic Health record systems, Radiology Information Systems, Clinical Information Systems or Cardiology Information Systems), the IHE Technical Framework intentionally avoids associating functions or actors with such product categories. For each actor, the IHE Technical Framework defines only those functions associated with integrating information systems. The IHE definition of an actor should therefore not be taken as the complete definition of any product that might implement it, nor should the framework itself be taken to comprehensively describe the architecture of a healthcare information system.

320 The reason for defining actors and transactions is to provide a basis for defining the interactions among functional components of the healthcare information system environment. In situations where a single physical product implements multiple functions, only the interfaces between the product and external functions in the environment are considered to be significant by the IHE initiative. Therefore, the IHE initiative takes no position as to the relative merits of an integrated environment based on a single, all-encompassing information system versus one based on multiple systems that together achieve the same end.
2.3 Relation of this Volume to the Technical Framework

The IHE Technical Framework is based on actors that interact through transactions using some form of content.

Actors are information systems or components of information systems that produce, manage, or act on information associated with operational activities in the enterprise.

Transactions are interactions between actors that transfer the required information through standards-based messages.

The implementation of the transactions described in this PCC TF-2 support the specification of Integration Profiles defined in PCC TF-1. The role and implementation of these transactions require the understanding of the Integration profile they support.

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.

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 Integration Profile.

Content Integration Profiles specify how the payload of a transaction fits into a specific use of that transaction. A content integration profile has three main parts. The first part describes the use case. The second part is binding to a specific IHE transaction, which describes how the content affects the transaction. The third part is a Content Module, which describes the payload of the transaction. A content module is specified so as to be independent of the transaction in which it appears.

2.3.1 Content Modules

The Patient Care Coordination Technical Framework organizes content modules categorically by the base standard. At present, the PCC Technical Framework uses only one base standard, CDA Release 2.0, but this is expected to change over time.

Underneath each standard, the content modules are organized using a very coarse hierarchy inherent to the standard. So for CDA Release 2.0 the modules are organized by document, section, entry, and header elements.

Each content module can be viewed as the definition of a "class" in software design terms, and has associated with it a name. Like "class" definitions in software design, a content module is a "contract", and the PCC Technical Framework defines that contract in terms of constraints that must be obeyed by instances of that content module. Each content module has a name, also known as its template identifier. The template identifiers are used to identify the contract agreed to by the content module. The PCC Technical Committee is responsible for assigning the template identifiers to each content module.
Like classes, content modules may inherit features of other content modules of the same type (Document, Section or Entry) by defining the parent content module that they inherit from. They may not inherit features from a different type. Although information in the CDA Header is in a different location that information in a CDA Entry, these two content modules are considered to be of the same type, and so may inherit from each other when necessary.

The PCC Technical Framework uses the convention that a content module cannot have more than one parent (although it may have several ancestors). This is similar to the constraint in the Java™ programming language, where classes can derive from only one parent. This convention is not due to any specific technical limitation of the technical framework, but does make it easier for software developers to implement content modules.

Each content module has a list of data elements that are required (R), required if known (R2), and optional (O). The presentation of this information varies with the type of content module, and is described in more detail below. Additional data elements may be provided by the sender that are not defined by a specific content module, but the receiver is not required to interpret them.

Required data elements must always be sent. Data elements that are required may under exceptional circumstances have an unknown value (e.g., the name of an unconscious patient). In these cases the sending application is required to indicate the reason that the data is not available.

Data elements that are marked required if known (R2) must be sent when the sending application has that data available. The sending application must be able to demonstrate that it can send all required if known elements, unless it does not in fact gather that data. When the information is not available, the sending application may indicate the reason that the data is not available.

Data elements that are marked optional (O) may be sent at the choice of the sending application. Since a content module may include data elements not specified by the profile, some might ask why these are specified in a content module. The reason for specifying the optional data elements is to ensure that both sender and receiver use the appropriate semantic interpretation of these elements. Thus, an optional element need not be sent, but when it is sent, the content module defines the meaning of that data element, and a receiver can always be assured of what that data element represents when it is present. Senders should not send an optional data element with an unknown value. If the value is not known, simply do not send the data element.

Other data elements may be included in an instance of a content module over what is defined by the PCC Technical Framework. Receivers are not required to process these elements, and if they do not understand them, must ignore them. Thus, it is not an error to include more than is asked for, but it is an error to reject a content module because it contains more than is defined by the framework. This allows value to be added to the content modules delivered in this framework, through extensions to it that are not defined or profiled by IHE. It further allows content modules to be defined later by IHE that are refinements or improvements over previous content modules.
For example, there is a Referral Summary content module defined in this framework. In later years an ED Referral content module can be created that inherits the constraints of the Referral Summary content module, with a few more use case specific constraints added. Systems that do not understand the ED Referral content module but do understand the Referral Summary content module will be able to interoperate with systems that send instances of documents that conform to the ED Referral content module. This interoperability, albeit at a reduced level of functionality, is by virtue of the fact that ED Referrals are simply a refinement of the Referral Summary.

In order to retain this capability, there are a few rules about how the PCC Technical Committee creates constraints. Constraints that apply to any content module will always apply to any content modules that inherit from it. Thus, the "contracts" are always valid down the inheritance hierarchy. Secondly, data elements of a content module will rarely be deprecated. This will usually occur only in the cases where they have been deprecated by the base standard. While any specific content module has a limited scope and set of use cases, deprecating the data element prevents any future content module from taking advantage of what has already been defined when a particular data element has been deprecated simply because it was not necessary in the original use case.

### 2.3.1.1 Document Content Module Constraints

Each document content module will define the appropriate codes used to classify the document, and will also describe the specific data elements that are included. The code used to classify it is specified using an external vocabulary, typically LOINC in the case of CDA Release 2.0 documents. The set of data elements that make up the document are defined, including the whether these data elements must, should or may be included in the document. Each data element is typically a section within the document, but may also describe information that is contained elsewhere within of the document (e.g., in the header). Each data element is mapped into a content module via a template identifier, and the document content module will further indicate whether these are data elements are required, required if known or optional. Thus, a document content module shall contain as constraints:

- The template identifier of the parent content module when there is one.
- The LOINC code or codes that shall be used to classify the document.
- A possibly empty set of required, required if known, and optional section content modules, and their template identifiers.
- A possibly empty set of required, required if known, and optional header content modules, and their template identifiers.
- Other constraints as necessary.

The template identifier for the document will be provided in the narrative, as will the legal LOINC document type codes and if present, any parent template identifier.

The remaining constraints are presented in two tables. The first table identifies the relevant data elements as determined during the technical analysis, and maps these data elements to one or more standards. The second table actually provides the constraints, wherein each data element identified in the first table is repeated, along with whether it is
required, required if known, or optional. Following this column is a reference to the specification for the content module that encodes that data element, and the template identifier assigned to it. The simple example below completes the content specification described above. A simplified example is shown below.

<table>
<thead>
<tr>
<th>Data Element Name</th>
<th>Opt</th>
<th>Template ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Section</td>
<td>R</td>
<td>SampleSectionOID</td>
</tr>
<tr>
<td>Comment on section</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample Entry</td>
<td>R2</td>
<td>SampleEntryOID</td>
</tr>
<tr>
<td>Comment on entry</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 2.3.1.1.1-1**

### 2.3.1.2 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate `<templateId>` elements in the header of the document. This is shown in the sample document below.

```xml
<ClinicalDocument xmlns='urn:hl7-org:v3'>
  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='SampleDocumentOID'/>
  <id root=' ' extension=' '/>
  <code code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <title>Sample Document</title>
  <effectiveTime value='20080601012005'/>
  <confidentialityCode code='N' displayName='Normal' codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality'/>
  <languageCode code='en-US'/>
  <component><structuredBody>
    <component>
      <section>
        <templateId root='SampleSectionOID'/>
        <!-- Required Sample Section Section content -->
      </section>
    </component>
  </structuredBody></component>
</ClinicalDocument>
```

**2.3.1.2 Section Content Module Constraints**

Section content modules will define the content of a section of a clinical document. Sections will usually contain narrative text, and so this definition will often describe the information present in the narrative, although sections may be wholly comprised of subsections.

Sections may contain various subsections, and these may be required, required if known or optional. Sections may also contain various entries, and again, these may be required,
required if known, or optional. A section may not contain just entries; it must have at least some narrative text or subsections to be considered to be valid content.

Again, sections can inherit features from other section content modules. Once again, sections are classified using an external vocabulary (again typically this would be LOINC), and so the list of possible section codes is also specified. Sections that inherit from other sections will not specify a LOINC code unless it is to restrict the type of section to smaller set of LOINC codes specified by one of its ancestors.

Thus, a section content module will contain as constraints:

- The template identifier of the parent content module when there is one.
- The LOINC code or codes that shall be used to classify the section.
- A possibly empty set of required, required if known, and optional section content modules, and their template identifiers for the subsections of this section.
- A possibly empty set of required, required if known, and optional entry content modules, and their template identifiers.
- Other constraints as necessary.

These constraints are presented in this document using a table for each section content module, as shown below.

<table>
<thead>
<tr>
<th>Sample Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>SampleSectionOID</td>
</tr>
<tr>
<td>foo (SampleParentOID)</td>
</tr>
<tr>
<td>Description of this section</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>XXXXX-X</td>
<td>R</td>
<td>SECTION NAME</td>
</tr>
</tbody>
</table>

| OID | R | Sample Entry |

<table>
<thead>
<tr>
<th>Subsections</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>OID</td>
<td>R</td>
<td>Sample Subsection</td>
</tr>
</tbody>
</table>

Table 2.3.1.1.2-1
Table 2.3.1.1.2-2
Table 2.3.1.1.2-3
Table 2.3.1.1.2-4 LOINC Codes
Table 2.3.1.1.2-5 General Description
Table 2.3.1.1.2-6 Parent Template

2.3.1.2.1 Parent Template
The parent of this template is foo.
2.3.1.3 Entry and Header Content Modules Constraints

Entry and Header content modules are the lowest level of content for which content modules are defined. These content modules are associated with classes from the HL7 Reference Information Model (RIM). These "RIM" content modules will constrain a single RIM class. Entry content modules typically constrain an "Act" class or one of its subtypes, while header content modules will normally constrain "Participation", "Role" or "Entity" classes, but may also constrain an "Act" class.

Entry and Header content modules will describe the required, required if known, and optional XML elements and attributes that are present in the CDA Release 2.0 instance. Header and Entry content modules may also be built up using other Header and Entry content modules. An entry or header content module may also specify constraints on the vocabularies used for codes found in the entry, or data types for the values found in the entry. Thus, an entry or header content module will contain as constraints:

- The template identifier of the parent content module when there is one.
- A description of the XML elements and attributes used in the entry, along with explanations of their meaning.
- An indication of those XML elements or attributes that are required, required if known, or optional.
- Vocabulary domains to use when coding the entry.
- Data types used to specify the value of the entry.
- Other constraints as necessary.
An example is shown below:

Sample Entry

Some text describing the entry.

```xml
<observation classCode='OBS' moodCode='EVN'>
    <templateId root='foo'/>
</observation>
```

2.3.1.4 `<observation classCode='OBS' moodCode='EVN'>`

Some details about the observation element

2.3.1.5 `<templateId root='foo'/>`

Some details about the template id element
3  IHE Transactions

This section defines each IHE transaction in detail, specifying the standards used, and the information transferred.

3.0 Cross Enterprise Document Content Transactions

At present, all transactions used by the PCC Content Profiles appear in ITI TF-2. General Options defined in content profiles for a Content Consumer are described below.

3.0.1 View Option

A Content Consumer that supports the View Option shall be able to:

1. Use the appropriate XD* transactions to obtain the document along with associated necessary metadata.

2. Render the document for viewing. This rendering shall meet the requirements defined for CDA Release 2 content presentation semantics (See Section 1.2.4 of the CDA Specification: Human readability and rendering CDA Documents). CDA Header information providing context critical information shall also be rendered in a human readable manner. This includes at a minimum the ability to render the document with the stylesheet specifications provided by the document source, if the document source provides a stylesheet. Content Consumers may optionally view the document with their own stylesheet, but must provide a mechanism to view using the source stylesheet.

3. Support traversal of links for documents that contain links to other documents managed within the sharing framework.

4. Print the document to paper.

3.0.2 Document Import Option

This Option requires that the View Option be supported. In addition, the Content Consumer that supports the Document Import Option shall be able to support the storage of the entire document (as provided by the sharing framework, along with sufficient metadata to ensure its later viewing) both for discharge summary or referral documents. This Option requires the proper tracking of the document origin. Once a document has been imported, the Content Consumer shall offer a means to view the document without the need to retrieve it again from the sharing framework. When viewed after it was imported, a Content Consumer may chose to access the sharing framework to find out if the related Document viewed has been deprecated, replaced or addended.

Note: For example, when using XDS, a Content Consumer may choose to query the Document Registry about a document previously imported in order to find out if this previously imported document may have been replaced or has received an addendum. This capability is offered to Content Consumers by this Integration Profile, but not required, as the events that may justify such a query are extremely implementation specific.
3.0.3 Section Import Option

This Option requires that the View Option be supported. In addition, the Content Consumer that supports the Section Import Option shall be able to support the import of one or more sections of the document (along with sufficient metadata to link the data to its source) both for discharge summary or referral. This Option requires the proper tracking of the document section origin. Once sections have been selected, a Content Consumer shall offer a means to copy the imported section(s) into local data structures as free text. This is to support the display of section level information for comparison or editing in workflows such as medication reconciliation while discrete data import is not possible. When viewed again after it is imported, a Content Consumer may chose to access the sharing framework to find out if the related information has been updated.

Note: For example, when using XDS, a Content Consumer may choose to query the Document Registry about a document whose sections were previously imported in order to find out if this previously imported document may have been replaced or has received an addendum. This capability is offered to Content Consumers by this Integration Profile, but not required, as the events that may justify such a query are extremely implementation specific.

This Option does not require, but does not exclude the Content Consumer from offering a means to select and import specific subsets of the narrative text of a section.

3.0.4 Discrete Data Import Option

This Option does not require that the View, Import Document or Section Import Options be supported. The Content Consumer that supports the Discrete Data Import Option shall be able to support the storage of the structured content of one or more sections of the document. This Option requires that the user be offered the possibility to select among the specific sections that include structured content a set of clinically relevant record entries (e.g. a problem or an allergy in a list) for import as part of the local patient record with the proper tracking of its origin.

Note: The Discrete Data Import Option does not require the support of the View, Import Document or Import Sections Options so that it could be used alone to support implementations of Content Consumers such as Public Health Data or Clinical Research systems that might aggregate and anonymize specific population healthcare information data as Document Consumer Actors, but one where no care provider actually views the medical summaries.

When discrete data is accessed after it was imported, a Content Consumer may choose to check if the document related to the discrete data viewed has been deprecated, replaced or addended.

A Content Consumer Actor grouped with the XDS Document Source Actor may query the Document Registry about a document from which discrete data was previously imported in order to find out if this previously imported document may have been replaced or has received an addendum. This capability is offered to Content Consumers by this Integration Profile, but not required, as the events that may justify such a query are extremely implementation specific.


### 4 IHE Patient Care Coordination Bindings

This section describes how the payload used in a transaction of an IHE profile is related to and/or constrains the data elements sent or received in those transactions. This section is where any specific dependencies between the content and transaction are defined.

A content integration profile can define multiple bindings. Each binding should identify the transactions and content to which it applies.

The source for all required and optional attributes have been defined in the bindings below. Three tables describe the three main XDS object types: XDSDocumentEntry, XDSSubmissionSet, and XDSFolder. XDSSubmissionSet and XDSDocumentEntry are required. Use of XDSFolder is optional. These concepts are universal to XDS, XDR and XDM.

The columns of the following tables are:

- **<XXX> attribute** – name of an XDS attribute, followed by any discussion of the binding detail.
- **Optional?** - Indicates the required status of the XDS attribute, and is one of R, R2, or O (optional). This column is filled with the values specified in the XDS Profile as a convenience.
- **Source Type** – Will contain one of the following values:

<table>
<thead>
<tr>
<th>Source Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SA</td>
<td>Source document Attribute – value is copied directly from source document. The Source/Value column identifies where in the source document this attribute comes from. Specify the location in XPath when possible.</td>
</tr>
<tr>
<td>SAT</td>
<td>Source document Attribute with Transformation – value is copied from source document and transformed. The Source/Value column identifies where in the source document this attribute comes from. Specify the location in XPath when possible. Extended Discussion column must not be empty and the transform must be defined in the extended discussion.</td>
</tr>
<tr>
<td>FM</td>
<td>Fixed (constant) by Mapping - for all source documents. Source/Value column contains the value to be used in all documents.</td>
</tr>
<tr>
<td>FAD</td>
<td>Fixed by Affinity Domain – value configured into Affinity Domain, all documents will use this value.</td>
</tr>
<tr>
<td>CAD</td>
<td>Coded in Affinity Domain – a list of acceptable codes are to be configured into Affinity Domain. The value for this attribute shall be taken from this list.</td>
</tr>
<tr>
<td>CADT</td>
<td>Coded in Affinity Domain with Transform - a list of acceptable codes are to be configured into Affinity Domain. The value for this attribute shall be taken from this list.</td>
</tr>
<tr>
<td>n/a</td>
<td>Not Applicable – may be used with an optionality R2 or O attribute to indicate it is not to be used.</td>
</tr>
<tr>
<td>DS</td>
<td>Document Source – value comes from the Document Source actor. Use Source/Value column or Extended Discussion to give details.</td>
</tr>
<tr>
<td>O</td>
<td>Other – Extended Discussion must be 'yes' and details given in an Extended Discussion.</td>
</tr>
</tbody>
</table>
The following tables are intended to be summaries of the mapping and transforms. The accompanying sections labeled 'Extended Discussion' are to contain the details as necessary.

4.1 Medical Document Binding to XDS, XDM and XDR

This binding defines a transformation that generates metadata for the XDSDocumentEntry element of appropriate transactions from the XDS, XDM and XDR profiles given a medical document and information from other sources. The medical document refers to the document being stored in a repository that will be referenced in the registry. The other sources of information include the configuration of the Document Source actor, the Affinity Domain, the site or facility, local agreements, other documents in the registry/repository, and this Content Profile.

In many cases, the CDA document is created for the purposes of sharing within an affinity domain. In these cases the context of the CDA and the context of the affinity domain are the same, in which case the following mappings shall apply.

In other cases, the CDA document may have been created for internal use, and are subsequently being shared. In these cases the context of the CDA document would not necessarily coincide with that of the affinity domain, and the mappings below would not necessarily apply.

Please note the specifics given in the table below.

### 4.1.1 XDSDocumentEntry Metadata

<table>
<thead>
<tr>
<th>XDSDocumentEntry Attribute</th>
<th>Optional?</th>
<th>Source Type</th>
<th>Source/ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>availabilityStatus</td>
<td>R</td>
<td>DS</td>
<td>$inst &lt;= /ClinicalDocument/author /assignedAuthor /representedOrganization</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>concat($inst/name)</td>
</tr>
<tr>
<td>authorInstitution</td>
<td>R2</td>
<td>SAT</td>
<td>$person &lt;= /ClinicalDocument/author</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>The authorInstitution can be formatted using the following XPath expression, where $inst in the expression below represents the representedOrganization. concat($inst/name)</td>
</tr>
<tr>
<td>authorPerson</td>
<td>R2</td>
<td>SAT</td>
<td>$person &lt;= /ClinicalDocument/author</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>The author can be formatted using the following XPath expression, where $person in the expression below represents the author. concat($person/id/@extension,&quot;^&quot;, $person/assignedPerson/name/family,&quot;^&quot;, $person/assignedPerson/name/given[1],&quot;^&quot;, $person/assignedPerson/name/given[2],&quot;^&quot;, $person/assignedPerson/name/suffix,&quot;^&quot;, $person/assignedPerson/name/suffix),&quot;^&quot;, $person/assignedPerson/name/suffix,&quot;^&quot;, $person/assignedPerson/name/suffix)</td>
</tr>
<tr>
<td>Field</td>
<td>Requirement</td>
<td>Type</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-------------</td>
<td>------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>authorRole</td>
<td>R2</td>
<td>SAT</td>
<td>This metadata element should be based on a mapping of the participation function defined in the CDA document to the set of author roles configured for the affinity domain. If the context of the CDA coincides with that of the affinity domain, then the following x-path may be appropriate: /ClinicalDocument/author/participationFunction</td>
</tr>
<tr>
<td>authorSpecialty</td>
<td>R2</td>
<td>SAT</td>
<td>This metadata element should be based on a mapping of the code associated with the assignedAuthor to detailed defined classification system for healthcare providers such configured in the affinity domain. Possible classifications include those found in SNOMED-CT, or the HIPAA Healthcare Provider Taxonomy. If the context of the CDA coincides with that of the affinity domain, then the following x-path may be appropriate: /ClinicalDocument/author/assignedAuthor/code</td>
</tr>
<tr>
<td>classCode</td>
<td>R</td>
<td>CADT</td>
<td>Derived from a mapping of /ClinicalDocument/code/@code to an Affinity Domain specified coded value to use and coding system. Affinity Domains are encouraged to use the appropriate value for Type of Service, based on the LOINC Type of Service (see Page 53 of the LOINC User's Manual). Must be consistent with /ClinicalDocument/code/@code</td>
</tr>
<tr>
<td>classCodeDisplayName</td>
<td>R</td>
<td>CADT</td>
<td>DisplayName of the classCode derived. Derived from a mapping of /ClinicalDocument/code/@code to the appropriate Display Name based on the Type of Service. Must be Consitent with /ClinicalDocument/code/@code</td>
</tr>
<tr>
<td>confidentialityCode</td>
<td>R</td>
<td>CADT</td>
<td>Derived from a mapping of /ClinicalDocument/confidentialityCode/@code to an Affinity Domain specified coded value and coding system. When using the BPPC profile, the confidentialityCode may also be obtained from the &lt;authorization&gt; element.</td>
</tr>
<tr>
<td>comments</td>
<td>O</td>
<td>DS</td>
<td></td>
</tr>
<tr>
<td>creationTime</td>
<td>R</td>
<td>SAT</td>
<td>Times specified in clinical documents may be specified with a precision in fractional sections, and may contain a time zone offset. In the XDS Metadata, it can be precise to the second, and is always given in UTC, so the timezone offset if present must be added to the current time to</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Field</th>
<th>Access</th>
<th>Activity</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>entryUUID</td>
<td>R</td>
<td>DS</td>
<td>obtain the UTC time.</td>
</tr>
<tr>
<td>eventCodeList</td>
<td>O</td>
<td>CADT</td>
<td>These values express a collection of keywords that may be relevant to the consumer of the documents in the registry. They may come from anywhere in the CDA document, according to its purpose.</td>
</tr>
<tr>
<td>eventCodeDisplayNameList</td>
<td>R (if event Code is valued)</td>
<td>CADT</td>
<td>These are the display names for the collection of keywords described above.</td>
</tr>
<tr>
<td>formatCode</td>
<td>R</td>
<td>FM</td>
<td>The format code for each PCC Document content profile is provided within the document specifications.</td>
</tr>
<tr>
<td>healthcareFacilityTypeCode</td>
<td>R</td>
<td>CAD</td>
<td>A fixed value assigned to the Document Source and configured form a set of Affinity Domain defined values. Must be consistent with /clinicalDocument/code</td>
</tr>
<tr>
<td>healthcareFacilityTypeDisplayName</td>
<td>R</td>
<td>CAD</td>
<td>Must be consistent with /clinicalDocument/code</td>
</tr>
</tbody>
</table>
| intendedRecipient (for XDR, XDM)  | O      | SAT      | The intendedRecipient can be formatted using the following XPath expression, where $inst in the expression below represents the receivedOrganization and where $person in the expression below represents the intendedRecipient. $person <= /ClinicalDocument/intendedRecipient and/or $inst <= /ClinicalDocument/intendedRecipient/receivedOrganization  
| legalAuthenticator                | O      | SAT      | The legalAuthenticator can be formatted using the following XPath expression, where $person in the expression below represents the legalAuthenticator. concat($person/id/@extension,"^",$person/informationRecipient/name/family,"^",$person/informationRecipient/name/given[1],"^",$person/informationRecipient/name/given[2],"^",$person/informationRecipient/name/suffix,"^",$person/informationRecipient/name/prefix,"^",$person/id/@root,"&ISO","

### PCC Technical Framework V5.0

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Role</th>
<th>Format</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$person/id/@extension,&quot;^&quot;, $person/assignedPerson/name/family,&quot;^&quot;, $person/assignedPerson/name/given[1],&quot;^&quot;, $person/assignedPerson/name/given[2],&quot;^&quot;, $person/assignedPerson/name/suffix,&quot;^&quot;, $person/assignedPerson/name/prefix,&quot;^&quot;, &quot;^^^&amp;&quot;, $person/id/@root,&quot;&amp;ISO&quot;)</td>
<td></td>
<td></td>
<td>Local document versions need not always be published, and so no exact mapping can be determined from the content of the CDA document. The parentDocumentRelationship may be determined in some configurations from the relatedDocument element present in the CDA document. If the context of the CDA coincides with that of the affinity domain, then the following x-path may be appropriate: /ClinicalDocument/relatedDocument/@typeCode</td>
</tr>
<tr>
<td>parentDocumentRelationship</td>
<td>R</td>
<td>DS</td>
<td>Local document versions need not always be published, and so no exact mapping can be determined from the content of the CDA document. The parentDocumentRelationship may be determined in some configurations from the relatedDocument element present in the CDA document. If the context of the CDA coincides with that of the affinity domain, then the following x-path may be appropriate: $docID &lt;= /ClinicalDocument/relatedDocument/parentDocument/id</td>
</tr>
<tr>
<td>parentDocumentId</td>
<td>R</td>
<td>DS</td>
<td>The parentDocumentId can be formatted using the following XPath expression, where $docID in the expression below represents the identifier. concat($docID/@root,&quot;^&quot;, $docID/@extension)</td>
</tr>
<tr>
<td>patientId</td>
<td>R</td>
<td>DS</td>
<td>The XDS Affinity Domain patient ID can be mapped from the patientRole/id element using transactions from the ITI PIX or PDQ profiles. See sourcePatientId below. If the context of the CDA coincides with that of the affinity domain, then the following x-path may be appropriate: $patID &lt;= /ClinicalDocument/recordTarget/patientRole/id</td>
</tr>
<tr>
<td>practiceSettingCode</td>
<td>R</td>
<td>CAD</td>
<td>This elements should be based on a coarse classification system for the class of specialty practice. Recommend the use of the classification system for Practice Setting, such as that described by the Subject Matter Domain in LOINC.</td>
</tr>
<tr>
<td>practiceSettingCodeDisplayName</td>
<td>R</td>
<td>CAD</td>
<td>This element shall contain the display names associated with the codes described above.</td>
</tr>
<tr>
<td>serviceStartTime</td>
<td>R2</td>
<td>SAT</td>
<td>Times specified in clinical documents may be specified with a precision in fractional sections, and may contain a</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Field</th>
<th>Required</th>
<th>Access</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>serviceStopTime</td>
<td>R2</td>
<td>SAT</td>
<td>Times specified in clinical documents may be specified with a precision in fractional sections, and may contain a time zone offset. In the XDS Metadata, it can be precise to the second, and is always given in UTC, so the timezone offset if present must be added to the current time to obtain the UTC time.</td>
</tr>
<tr>
<td>sourcePatientId</td>
<td>R</td>
<td>SAT</td>
<td>$patID &lt;= /ClinicalDocument/recordTarget/patientRole/id</td>
</tr>
<tr>
<td>sourcePatientInfo</td>
<td>R</td>
<td>SAT</td>
<td>/ClinicalDocument/recordTarget/patientRole</td>
</tr>
<tr>
<td>title</td>
<td>O</td>
<td>SA</td>
<td>/ClinicalDocument/title</td>
</tr>
<tr>
<td>typeCode</td>
<td>R</td>
<td>CADT</td>
<td>/ClinicalDocument/code/@code</td>
</tr>
<tr>
<td>typeCodeDisplay Name</td>
<td>R</td>
<td>CADT</td>
<td>/ClinicalDocument/code/@displayName</td>
</tr>
<tr>
<td>uniqueId</td>
<td>R</td>
<td>SAT</td>
<td>$docID &lt;= /ClinicalDocument/id</td>
</tr>
</tbody>
</table>

### 4.1.1.1 XDSSubmissionSet Metadata

The submission set metadata is as defined for XDS, and is not necessarily affected by the content of the clinical document. Metadata values in an XDSSubmissionSet with names identical to those in the XDSDocumentEntry may be inherited from XDSDocumentEntry metadata, but this is left to affinity domain policy and/or application configuration.
4.1.1.2 Use of XDS Submission Set
This content format uses the XDS Submission Set to create a package of information to send from one provider to another. All documents referenced by the Medical Summary in this Package must be in the submission set.

4.1.1.3 Use of XDS Folders
No specific requirements identified.

4.1.1.4 Configuration
IHE Content Profiles using this binding require that Content Creators and Content Consumers be configurable with institution and other specific attributes or parameters. Implementers should be aware of these requirements to make such attributes easily configurable. There shall be a mechanism for the publishing and distribution of style sheets used to view clinical documents.

4.1.2 Extensions from other Domains

4.1.2.1 Scanned Documents (XDS-SD)
XDS-SD is a CDA R2 document and thus conforms to the XDS Metadata requirements in the PCC-TF, volume 2, Section 5 unless otherwise specified below.

4.1.2.1.1 XDSDocumentEntry
XDS-SD leverages the XDS DocumentEntry Metadata requirements in the PCC-TF, volume 2, Section 5.1.1.1.1.1 and in PCC_TF-2/Bindings unless otherwise specified below.

4.1.2.1.1.1 XDSDocumentEntry.formatCode
The XDSDocumentEntry.formatCode shall be urn:ihe:iti:xds-sd:pdf:2008 when the document is scanned pdf and urn:ihe:iti:xds-sd:text:2008 when the document is scanned text. The formatCode codeSystem shall be 1.3.6.1.4.1.19376.1.2.3.

4.1.2.1.1.2 XDSDocumentEntry.uniqueId
This value shall be the ClinicalDocument/id in the HL7 CDA R2 header. The root attribute is required, and the extension attribute is optional. In accordance with the XDS.a profile, total length is limited to 128 characters; for XDS.b the limit is 256 characters. Additionally see PCC-TF, volume 2, Section 5.1.1.1.1 or PCC_TF-2/Bindings for further content specification.

4.1.2.1.1.3 Relating instances of XDS-SD documents
In general, most instances of XDS-SD will not have parent documents. It is possible, however, in some specific use cases that instances of XDS-SD documents are related. For example, for a particular document it may be the case that both the PDF scanned content and somewhat equivalent plaintext need to be wrapped and submitted. Each document would correspond to separate XDSDocumentEntries linked via an XFRM Association that indicates one document is a transform of the other. These can be submitted in a
single submission set, or in separate ones. Other specific examples may exist and this profile does not preclude the notion of a parent document for these cases.

**4.1.2.1.2 XDSSubmissionSet**

No additional constraints. Particular to this profile, a legitimate use of submission sets would be to maintain a logical grouping of multiple XDS-SD documents. We encourage such usage. For more information, see PCC-TF-2 Section 5.1.1.1.2 or [PCC_TF-2/Bindings](#).

**4.1.2.1.3 XDSFolder**

No additional requirements. For more information, see PCC-TF-2 Section 5.1.1.1.3 or [PCC_TF-2/Bindings](#).

**4.1.2.2 Basic Patient Privacy Consents (BPPC)**

**4.1.2.3 Laboratory Reports (XD-LAB)**

XD-Lab is a CDA R2 document and thus conforms to the XDS Metadata requirements in the PCC-TF, volume 2, Section 5 unless otherwise specified below.

**4.1.2.3.1 XDSDocumentEntry**

XD-Lab leverages the XDS DocumentEntry Metadata requirements in the PCC-TF, volume 2, Section 5.1.1.1.1 and in [PCC_TF-2/Bindings](#) unless otherwise specified below.

**4.1.2.3.1.1 XDSDocumentEntry.eventCodeList**

XD-Lab documents further constrain the the XDSDocumentEntry.eventCodeList to the following.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Optional?</th>
<th>Source Type</th>
<th>Source/ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>eventCodeList</td>
<td>R2</td>
<td>SAT</td>
<td>ClinicalDocument / component / structuredBody / component / section / entry / act / entryRelationship / organizer (templateId=&quot;1.3.6.1.4.1.19376.1.3.1.1&quot;)/ component / observation(templateId=&quot;1.3.6.1.4.1.19376.1.3.1.1.1&quot;)/code AND ClinicalDocument / component / structuredBody / component / section / entry / act / subject / code If the document has Reportable Condition, then this code shall be among those listed in the eventCodeList. Additionally, if the document contains information about a</td>
</tr>
</tbody>
</table>
Non-Human Subject, then the code that indicates what this subject is shall be among those listed in the eventCodeList. Thus, this attribute has been enhanced from the XDS profile from O to R2.

4.1.2.3.1.2 XDSDocumentEntry.formatCode

The XDSDocumentEntry.formatCode shall be urn:ihe:lab:xd-lab:2008 The formatCode codeSystem shall be 1.3.6.1.4.1.19376.1.2.3.

4.1.2.3.2 XDSSubmissionSet

No additional constraints. For more information, see PCC-TF-2 Section 5.1.1.1.2 or PCC_TF-2/Bindings.

4.1.2.3.3 XDSFolder

No additional requirements. For more information, see PCC-TF-2 Section 5.1.1.1.3 or PCC_TF-2/Bindings.
5 Namespaces and Vocabularies

This section lists the namespaces and identifiers defined or referenced by the IHE PCC Technical Framework, and the vocabularies defined or referenced herein.

The following vocabularies are referenced in this document. An extensive list of registered vocabularies can be found at [http://hl7.amg-hq.net/oid/frames.cfm](http://hl7.amg-hq.net/oid/frames.cfm).

<table>
<thead>
<tr>
<th>codeSystem</th>
<th>codeSystemName</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1</td>
<td>IHE PCC Template Identifiers</td>
<td>This is the root OID for all IHE PCC Templates. A list of PCC templates can be found below in CDA Release 2.0 Content Modules.</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.2</td>
<td>IHEActCode</td>
<td>See IHEActCode Vocabulary below</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.3</td>
<td>IHE PCC RoleCode</td>
<td>See IHERoleCode Vocabulary below</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.4</td>
<td>CCD Root OID</td>
<td>Namespace OID used for IHE Extensions to CDA Release 2.0</td>
</tr>
<tr>
<td>2.16.840.1.113883.10.20.1</td>
<td>RouteOfAdministration</td>
<td>Root OID used for by ASTM/HL7 Continuity of Care Document</td>
</tr>
<tr>
<td>2.16.840.1.113883.5.112</td>
<td>SeverityObservation</td>
<td>See the HL7 RouteOfAdministration Vocabulary</td>
</tr>
<tr>
<td>2.16.840.1.113883.5.1063</td>
<td>LOINC</td>
<td>Logical Observation Identifier Names and Codes</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.96</td>
<td>SNOMED-CT</td>
<td>SNOMED Controlled Terminology</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.103</td>
<td>ICD-9CM (diagnosis codes)</td>
<td>International Classification of Diseases, Clinical Modifiers, Version 9</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.104</td>
<td>ICD-9CM (procedure codes)</td>
<td>International Classification of Diseases, Clinical Modifiers, Version 9</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.26</td>
<td>MEDCIN</td>
<td>A classification system from MEDICOMP Systems.</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.88</td>
<td>RxNorm</td>
<td>RxNorm</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.63</td>
<td>FDDC</td>
<td>First DataBank Drug Codes</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.257</td>
<td>Minimum Data Set for Long Term Care</td>
<td>The root OID for Minimum Data Set Answer Lists</td>
</tr>
</tbody>
</table>

5.1.1 IHE Format Codes

The table below lists the format codes, template identifiers and media types used by the IHE Profiles specified in the PCC Technical Framework, and also lists, for reference purposes the same values for other selected IHE Profiles from other committees.

Note that the code system for these codes is 1.3.6.1.4.1.19376.1.2.3 as assigned by the
ITI Domain for codes used for the purposes of cross-enterprise document sharing (XDS). For more information see XDS Coding System (1.3.6.1.4.1.19376.1.2.3).

<table>
<thead>
<tr>
<th>Profile</th>
<th>Format Code</th>
<th>Media Type</th>
<th>Template ID</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2006 Profiles</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medical Summaries (XDS-MS)</td>
<td>urn:ihe:pcc:xds-ms:2007</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.3 (Referral)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.4 (Discharge Summary)</td>
</tr>
<tr>
<td><strong>2007 Profiles</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exchange of Personal Health Records (XPHR)</td>
<td>urn:ihe:pcc:xphr:2007</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.5 (Extract)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.6 (Update)</td>
</tr>
<tr>
<td>Emergency Department Referral (EDR)</td>
<td>urn:ihe:pcc:edr:2007</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.10</td>
</tr>
<tr>
<td><strong>2008 Profiles</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Antepartum Summary (APS)</td>
<td>urn:ihe:pcc:aps:2007</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.11.2</td>
</tr>
<tr>
<td>Emergency Department Encounter Summary (EDES)</td>
<td>urn:ihe:pcc:edes:2007</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.1.1 (Triage Note)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.1.2 (Nursing Note)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.1.3 (Composite Triage and Nursing Note)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.1.4 (Physician Note)</td>
</tr>
<tr>
<td><strong>2009 Profiles</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Antepartum Record (APR) - History and Physical</td>
<td>urn:ihe:pcc:apr:handp:2008</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.16.1.1 (Antepartum History and Physical)</td>
</tr>
<tr>
<td>Antepartum Record (APR) - Laboratory</td>
<td>urn:ihe:pcc:apr:lab:2008</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.16.1.2 (Antepartum Laboratory)</td>
</tr>
<tr>
<td>Antepartum Record (APR) - Education</td>
<td>urn:ihe:pcc:apr:edu:2008</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.16.1.3 (Antepartum Education)</td>
</tr>
<tr>
<td>Immunization Registry Content (IRC)</td>
<td>urn:ihe:pcc:irc:2008</td>
<td>text/xml</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.18.1.2 (Immunization Detail)</td>
</tr>
<tr>
<td>Cancer Registry Content (CRC)</td>
<td>urn:ihe:pcc:crc:2008</td>
<td>text/xml</td>
<td></td>
</tr>
<tr>
<td>Care Management (CM)</td>
<td>urn:ihe:pcc:cm:2008</td>
<td>text/xml</td>
<td></td>
</tr>
</tbody>
</table>

**ITI Profiles**
5.1.2 IHEActCode Vocabulary

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>COMMENT</td>
<td>This is the act of commenting on another act.</td>
</tr>
<tr>
<td>PINSTRUCT</td>
<td>This is the act of providing instructions to a patient regarding the use of medication.</td>
</tr>
<tr>
<td>FINSTRUCT</td>
<td>This is the act of providing instructions to the supplier regarding the fulfillment of the medication order.</td>
</tr>
<tr>
<td>IMMUNIZ</td>
<td>The act of immunization of a patient using a particular substance or class of substances identified using a specified vocabulary. Use of this vocabulary term requires the use of either the SUBSTANCE or SUBSTCLASS qualifier described below, along with an identified substance or class of substances.</td>
</tr>
<tr>
<td>DRUG</td>
<td>The act of treating a patient with a particular substance or class of substances identified using a specified vocabulary. Use of this vocabulary term requires the use of either the SUBSTANCE or SUBSTCLASS qualifier described below, along with an identified substance or class of substances.</td>
</tr>
<tr>
<td>INTOL</td>
<td>An observation that a patient is somehow intollerant of (e.g., allergic to) a particular substance or class of substances using a specified vocabulary. Use of this vocabulary term requires the use of either the SUBSTANCE or SUBSTCLASS qualifier described below, along with an identified substance or class of substances.</td>
</tr>
<tr>
<td>SUBSTANCE</td>
<td>A qualifier that identifies the substance used to treat a patient in an immunization or drug treatment act. The substance is expected to be identified using a vocabulary such as RxNORM, SNOMED</td>
</tr>
</tbody>
</table>
CT or other similar vocabulary and should be specific enough to identify the ingredients of the substance used.

<table>
<thead>
<tr>
<th>SUBSTCLASS</th>
<th>Description</th>
</tr>
</thead>
</table>
| A qualifier that identifies the class of substance used to treat a patient in an immunization or drug treatment act. The class of substances is expected to be identified using a vocabulary such as NDF-RT, SNOMED CT or other similar vocabulary, and should be broad enough to classify substances by mechanism of action (e.g., Beta Blocker), intended effect (Diuretic, antibiotic) or ...

### 5.1.3 IHERoleCode Vocabulary

The IHERoleCode vocabulary is a small vocabulary of role codes that are not presently supported by the HL7 Role Code vocabulary. The root namespace (OID) for this vocabulary is 1.3.5.1.4.1.19376.1.5.3.3.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMPLOYER</td>
<td>The employer of a person.</td>
</tr>
<tr>
<td>SCHOOL</td>
<td>The school in which a person is enrolled.</td>
</tr>
<tr>
<td>AFFILIATED</td>
<td>An organization with which a person is affiliated (e.g., a volunteer organization).</td>
</tr>
<tr>
<td>PHARMACY</td>
<td>The pharmacy a person uses.</td>
</tr>
</tbody>
</table>
6  PCC Content Modules

6.1 Conventions

Various tables used in this section will further constrain the content. Within this volume, the follow conventions are used.

R

A "Required" data element is one that shall always be provided. If there is information available, the data element must be present. If there is no information available, or it cannot be transmitted, the data element must contain a value indicating the reason for omission of the data. (See PCC TF-2: 5.3.4.2 for a list of appropriate statements).

R2

A "Required if data present" data element is one that shall be provided when a value exists. If the information cannot be transmitted, the data element shall contain a value indicating the reason for omission of the data. If no such information is available to the creator or if such information is not available in a well identified manner (e.g. buried in a free form narrative that contains additional information relevant to other sections) or if the creator requires that information be absent, the R2 section shall be entirely absent. (See section PCC TF-2: 5.3.4.2 for a list of appropriate statements).

O

An optional data element is one that may be provided, irrespective of whether the information is available or not. If the implementation elects to support this optional section, then its support shall meet the requirement set forth for the "Required if data present" or R2.

C

A conditional data element is one that is required, required if known or optional depending upon other conditions. These will have further notes explaining when the data element is required, et cetera.

Note: The definitions of R, R2, and O differ slightly from other IHE profiles. This is due in part to the fact that local regulations and policies may in fact prohibit the transmission of certain information, and that a human decision to transmit the information may be required in many cases.

6.2 Folder Content Modules

This section contains modules that describe the content requirements of Folders used with XDS, XDM or XDR. When workflows are completed normally, the folders will contain documents with the optionality specified in the tables shown below. Under certain circumstances, the folders will not meet the optionality requirements described below, for example, when the patient leaves before treatment is completed.
6.2.1 EDES Folder Specification

6.2.2 APR Folder Specification
6.2.3 LDR Folder Specification

6.3 HL7 Version 3.0 Content Modules

This section contains content modules based upon the HL7 CDA Release 2.0 Standard, and related standards and/or implementation guides.

6.3.1 CDA Document Content Modules

6.3.1.1 Medical Documents Specification 1.3.6.1.4.1.19376.1.5.3.1.1.1

This section defines the base set of constraints used by almost all medical document profiles described the PCC Technical Framework.

6.3.1.1.1 Standards

<table>
<thead>
<tr>
<th>CDAR2</th>
<th>HL7 CDA Release 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>CDTHP</td>
<td>CDA for Common Document Types History and Physical Notes (DSTU)</td>
</tr>
<tr>
<td>XMLXSL</td>
<td>Associating Style Sheets with XML documents</td>
</tr>
</tbody>
</table>

6.3.1.1.2 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate <templateId> elements in the header of the document. This is shown in the sample document below.

```xml
<ClinicalDocument xmlns='urn:hl7-org:v3'>
  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.1'/>
  <id root=' ' extension=' '/>
  <code code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <title>Medical Documents</title>
  <effectiveTime value='20081004012005'/>
  <confidentialityCode code='N' displayName='Normal' codeSystem='2.16.840.1.113883.6.1' codeSystemName='Confidentiality' />
  <languageCode code='en-US'/>
  <component>
    <structuredBody>
      ...
    </structuredBody>
  </component>
</ClinicalDocument>
```

Figure 6.3-1 Sample Medical Documents Document
<!-- Verify the document type code -->

<assert test='cda:code[@code = "{{{LOINC}}}"]'>
   Error: The document type code of a Medical Documents must be {{{LOINC}}}.
</assert>

<assert test='cda:code[@codeSystem = "2.16.840.1.113883.6.1"]'>
   Error: The document type code must come from the LOINC code system (2.16.840.1.113883.6.1).
</assert>

6.3.1.1.3 Specification

The constraints for encoding of the CDA Header (Level 1) can be found in the CDA for Common Document Types History and Physical Implementation Guide, in the section 2. CDA Header -- General Constraints.

- IHE Medical Documents Shall follow all constraints found in that section with the exception of the constraint on realmcode found in CONF-HP-10:

- IHE Medical Documents which are implemented for the US Realm Shall follow ALL constraints found in that section, and Shall use both the IHE Medical Document templateId (1.3.6.1.4.1.19376.1.5.3.1.1.1) and the HL7 General Header Constraints templateId (2.16.840.1.113883.10.20.3).}

<table>
<thead>
<tr>
<th>Realm</th>
<th>Constraints</th>
<th>Template IDs Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Universal</td>
<td>CONF-HP-1 through CONF-HP-9</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.1</td>
</tr>
<tr>
<td></td>
<td>CONF-HP-11 through CONF-HP-40</td>
<td></td>
</tr>
<tr>
<td>US</td>
<td>CONF-HP-1 through CONF-HP-40</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.16.840.1.113883.10.20.3</td>
</tr>
</tbody>
</table>

6.3.1.1.4 Style Sheets

Document sources Should provide an XML style sheet to render the content of the Medical Summary document. The output of this style sheet Shall be an XHTML Basic (see http://www.w3.org/TR/xhtml-basic/) document that renders the clinical content of a Medical Summary Document as closely as possible as the sending provider viewed the completed document. When a style sheet is provided, at least one processing instruction Shall be included in the document that including a link to the URL for the XML style sheet. To ensure that the style sheet is available to all receivers, more than one stylesheet link May be included.

When a stylesheet is used within an XDS Affinity domain, the link to it Shall be provided using an HTTPS or HTTP URL.

<?xml-stylesheet href='https://foobar:8080/mystylesheet.xsl' type='text/xsl'?>

When using XDM or XDR to exchange documents, the stylesheet Shall also be exchanged on the media. The link to the stylesheet Shall be recorded as a relative URL.

<?xml-stylesheet href='../../stylesheets/mystylesheet.xsl' type='text/xsl'?>

Style sheets Should NOT rely on graphic or other media resources. If graphics other
media resources are used, these **SHALL** be accessible in the same way as the stylesheet. The **Content Creator** **NEED NOT** be the provider of the resources (stylesheet or graphics).

When a Content Creator provides a style sheet, Content Consumers **MUST** provide a mechanism to render the document with that style sheet. Content Consumers **MAY** view the document with their own style sheet.

To record the style sheet within a CDA Document that might be used in both an XDS and XDM environment, more than one style sheet processing instruction is required. In this case, all style sheet processing instructions included **MUST** include the alternate='yes' attribute.

```xml
<?xml-stylesheet href='https://foobar:8080/mystylesheet.xsl' type='text/xsl' alternate='yes'?>
<?xml-stylesheet href='../../stylesheets/mystylesheet.xsl' type='text/xsl' alternate='yes'?>
```

A Content Consumer that is attempting to render a document using the document supplied style sheet **MAY** use the first style sheet processing instruction for which it is able to obtain the style sheet content, and **SHALL NOT** report any errors if it is able to find at least one style sheet to render with.

### 6.3.1.1.5 Distinctions of None

Information that is sent **MUST** clearly identify distinctions between

- **None**
  - It is known with complete confidence that there are none. Used in the context of problem and medication lists, this indicates that the sender knows that there is no relevant information that can be sent.

- **None Known**
  - None are known at this time, but it is not known with complete confidence than none exist. Used in the context of allergy lists, where essentially, it is impossible to prove the negative that no allergies exist, it is only possible to assert that none have been found to date.

- **None Known Did Ask**
  - None are known at this time, and it is not known with complete confidence than none exist, but the information was requested. Also used in the context of allergy lists, where essentially, it is impossible to prove the negative that no allergies exist, it is only possible to assert that none have been found to date.

- **Unknown**
  - The information is not known, or is otherwise unavailable.

In the context of CDA, sections that are required to be present but have no information should use one of the above phrases where appropriate.

An appropriate machine readable entry shall be present for problems, medications and allergies to indicate the reason that no information. Codes for recording unknown or no information are provided in the section on the **Problem**, **Allergy** and **Medications** Entry.
6.3.1.2 Medical Summary Specification 1.3.6.1.4.1.19376.1.5.3.1.1.2

A medical summary contains a snapshot of the patient's medical information, including at the very least, a list of the patient's problems, medications and allergies. A Medical Summary is an abstract template that is expected to be further refined by additional document templates.

6.3.1.2.1 Parent Template

This document is an instance of the Medical Document template.

6.3.1.2.2 Standards

CDAR2 HL7 CDA Release 2.0

6.3.1.2.3 Specification

<table>
<thead>
<tr>
<th>Data Element Name</th>
<th>Opt</th>
<th>Template ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Problem Concern Entry</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.2</td>
</tr>
<tr>
<td>Allergy Concern Entry</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.3</td>
</tr>
<tr>
<td>Medications</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.7</td>
</tr>
</tbody>
</table>

Table 6.3.1.2.3-1

6.3.1.2.4 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate <templateId> elements in the header of the document. This is shown in the sample document below. A CDA Document may conform to more than one template. This content module inherits from the Medical Document content module, and so must conform to the requirements of that template as well, thus all <templateId> elements shown in the example below shall be included.

```
<ClinicalDocument xmlns='urn:hl7-org:v3'>
  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.1'/>  
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.2'/>
  <id root=' ' extension=' '/>
  <code code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <title>Medical Summary</title>
  <effectiveTime value='20081004012005'/>
  <confidentialityCode code='N' displayName='Normal' codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality'/>
  <languageCode code='en-US'/>
  :
  <component><structuredBody>
   :  
  </structuredBody></component>
</ClinicalDocument>
```

Figure 6.3-2 Sample Medical Summary Document
6.3.1.2.5 Document Specification

A medical summary is a type of medical document, and incorporates the constraints defined for Medical Documents, and requires the recording of Problems, Allergies and Medications.

6.3.1.3 Referral Summary Specification 1.3.6.1.4.1.19376.1.5.3.1.1.3

The use case is described fully in PCC TF-1:3.2.1 for the Ambulatory Specialist Referral. Briefly, it involves a "collaborative" transfer of care for the referral of a patient from a primary care provider (PCP) to a specialist. The important document data elements identified by physicians and nurses for this use case are listed in the table below under the column "Data Elements". These were then mapped to the categories given HL7 Care Record Summary Implementation Guide, and HL7 CDA Release 2.0. These mappings are provided in the next two columns.

A referral summary is a type of Medical Summary, and incorporates the constraints defined for a Medical Summary above. This section defines additional constraints for Medical Summary Content used in a Referral summary. These tables present the Categories, as defined in Section 3 of CRS. In no case are these IHE requirements less strict than those defined by CRS.

6.3.1.3.1 Format Code

The XDSDocumentEntry format code for this content is urn:ihe:pcc:xds-ms:2007

6.3.1.3.2 Parent Template

This document is an instance of the Medical Summary template.

6.3.1.3.3 Standards

<table>
<thead>
<tr>
<th>CDAR2</th>
<th>HL7 CDA Release 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRS</td>
<td>HL7 Care Record Summary</td>
</tr>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
</tbody>
</table>

6.3.1.3.4 Data Element Index

<table>
<thead>
<tr>
<th>Data Elements</th>
<th>HL7 Care Record Summary</th>
<th>CDA Release 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reason for Referral</td>
<td>Reason for Referral</td>
<td>REASON FOR REFERRAL</td>
</tr>
<tr>
<td>History Present Illness</td>
<td>History of Present Illness</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
<tr>
<td>Active Problems</td>
<td>Conditions</td>
<td>PROBLEM LIST</td>
</tr>
<tr>
<td>Current Meds</td>
<td>Medications</td>
<td>HISTORY OF MEDICATION USE</td>
</tr>
<tr>
<td>Allergies</td>
<td>Allergies and Adverse</td>
<td>HISTORY OF ALLERGIES</td>
</tr>
<tr>
<td>Data Element Name</td>
<td>Opt</td>
<td>Template ID</td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>-----</td>
<td>---------------------------------</td>
</tr>
<tr>
<td>Reason for Referral</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.1</td>
</tr>
<tr>
<td>History Present Illness</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.4</td>
</tr>
<tr>
<td>Active Problems</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.6</td>
</tr>
<tr>
<td>Current Meds</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.19</td>
</tr>
<tr>
<td>Allergies</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.13</td>
</tr>
<tr>
<td>History of Past Illness</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.8</td>
</tr>
<tr>
<td>List of Surgeries</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.11</td>
</tr>
<tr>
<td>Immunizations</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.23</td>
</tr>
<tr>
<td>Family History</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.14</td>
</tr>
<tr>
<td>Social History</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.16</td>
</tr>
</tbody>
</table>
### Pertinent Review of Systems

**Vital Signs**

- **1.3.6.1.4.1.19376.1.5.3.1.3.25**

**Physical Exam**

- **1.3.6.1.4.1.19376.1.5.3.1.3.24**

**Relevant Diagnostic Surgical Procedures / Clinical Reports and Relevant Diagnostic Test and Reports**

(Tests, Imaging, EKG's, etc.) including links.

- **1.3.6.1.4.1.19376.1.5.3.1.3.27**

**Plan of Care (new meds, labs, or x-rays ordered)**

- **1.3.6.1.4.1.19376.1.5.3.1.3.31**

**Advance Directives**

- **1.3.6.1.4.1.19376.1.5.3.1.3.34**

**Patient Administrative Identifiers**

Handled by the Medical Documents Content Profile by reference to constraints in HL7 CRS.

**Pertinent Insurance Information**

Refer to Appropriate Payers Section -- TBD

**Data needed for state and local referral forms, if different than above**

These are handed by including additional sections within the summary.

<table>
<thead>
<tr>
<th>Table 6.3.1.3.5-1</th>
</tr>
</thead>
</table>

### 6.3.1.3.6 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate `<templateId>` elements in the header of the document. This is shown in the sample document below. A CDA Document may conform to more than one template. This content module inherits from the **Medical Summary** content module, and so must conform to the requirements of that template as well, thus all `<templateId>` elements shown in the example below shall be included.

```xml
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  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.2'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.3'/>
  <id root=' ' extension=' '/>
  <code code=' ' display=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <title>Referral Summary</title>
  <effectiveTime value='20081004012005'/>
  <confidentialityCode code='N' display='Normal' codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality'/>
  <languageCode code='en-US'/>
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    <structuredBody>
      <component>
        <section>
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          <!-- Required Reason for Referral Section content -->
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      </component>
      <component>
        <section>
          <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.4'/>
          <!-- Required History Present Illness Section Content -->
        </section>
      </component>
    </structuredBody>
  </component>
</ClinicalDocument>
```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.8'/>
    <!-- Required if known History of Past Illness Section content -->
  </section>
</component>

<component>
  <section>
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</component>

<component>
  <section>
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</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18'/>
    <!-- Optional Pertinent Review of Systems Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.25'/>
    <!-- Required if known Vital Signs Section content -->
  </section>
</component>
Figure 6.33-3 Sample Referral Summary Document
6.3.1.4 Discharge Summary Specification 1.3.6.1.4.1.19376.1.5.3.1.1.4

This use case is described fully in the XDS-MS profile found in PCC TF-1:3.2.2. Briefly, it involves an episodic transfer of care in the form of a patient discharge from a hospital to home. The important data elements identified by physicians and nurses for this use case are listed in the table below under the column "Data Elements". These are mapped to the categories given HL7 Care Record Summary Implementation Guide, and HL7 CDA Release 2.0 in the next two columns.

A discharge summary is a type of medical summary, and incorporates the constraints defined for Medical Summaries.

This section defines additional constraints for Medical Summary Content used in a Discharge Summary. These tables present the data elements described above, along with their optionality, and references to the section and template where these sections or header data elements are further defined.

In no case are these IHE requirements less strict than those defined by the HL7 Care Record Summary.

6.3.1.4.1 Format Code

The XDSDocumentEntry format code for this content is urn:ihe:pcc:xds-ms:2007

6.3.1.4.2 Parent Template

This document is an instance of the Medical Summary template.

6.3.1.4.3 Standards

<table>
<thead>
<tr>
<th>CDAR2</th>
<th>HL7 CDA Release 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRS</td>
<td>HL7 Care Record Summary</td>
</tr>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
</tbody>
</table>

6.3.1.4.4 Data Element Index

<table>
<thead>
<tr>
<th>Data Elements</th>
<th>HL7 Care Record Summary</th>
<th>CDA Release 2.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date of Admission</td>
<td>Header</td>
<td>encompassingEncounter/effectiveTime</td>
</tr>
<tr>
<td>Date of Discharge</td>
<td>Header</td>
<td>encompassingEncounter/effectiveTime</td>
</tr>
<tr>
<td>Participating Providers and Roles</td>
<td>Header</td>
<td>documentationOf/serviceEvent/performer</td>
</tr>
<tr>
<td>Discharge Disposition (who, how, where)</td>
<td>Care Plan</td>
<td>DISCHARGE DISPOSITION</td>
</tr>
<tr>
<td>Admitting Diagnosis</td>
<td>Conditions</td>
<td>HOSPITAL ADMISSION DX</td>
</tr>
<tr>
<td>History of Present Illness</td>
<td>History of Present Illness</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
</tbody>
</table>
## 6.3.1.4.5 Specification

<table>
<thead>
<tr>
<th>Data Element Name</th>
<th>Opt</th>
<th>Template ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active Problems</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.6</td>
</tr>
<tr>
<td>Resolved Problems</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.8</td>
</tr>
<tr>
<td>Discharge Diagnosis</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.7</td>
</tr>
<tr>
<td>Admitting Diagnosis</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.3</td>
</tr>
<tr>
<td>Selected Meds Administered</td>
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</tr>
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<td>Discharge Meds</td>
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</tr>
<tr>
<td>Admission Medications</td>
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</tr>
<tr>
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<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.13</td>
</tr>
<tr>
<td>Hospital Course</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.5</td>
</tr>
<tr>
<td>Advance Directives</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.34</td>
</tr>
<tr>
<td>History of Present Illness</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.4</td>
</tr>
<tr>
<td>Functional Status</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.17</td>
</tr>
<tr>
<td>Section</td>
<td>Status</td>
<td>Template Id</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>--------</td>
<td>------------------------------</td>
</tr>
<tr>
<td>Review of Systems</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.18</td>
</tr>
<tr>
<td>Physical Examination</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.24</td>
</tr>
<tr>
<td>Vital Signs</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.25</td>
</tr>
<tr>
<td>Discharge Procedures Tests, Reports</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.29</td>
</tr>
<tr>
<td>Plan of Care</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.31</td>
</tr>
<tr>
<td>Discharge Diet</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.33</td>
</tr>
</tbody>
</table>

**Table 6.3.1.4.5-1**

**6.3.1.4.6 Conformance**

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate `<templateId>` elements in the header of the document. This is shown in the sample document below. A CDA Document may conform to more than one template. This content module inherits from the [Medical Summary](#) content module, and so must conform to the requirements of that template as well, thus all `<templateId>` elements shown in the example below shall be included.
<ClinicalDocument xmlns='urn:hl7-org:v3'>
  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.2'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.4'/>
  <id root=' ' extension=' '/>
  <code code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <title>Discharge Summary</title>
  <effectiveTime value='20081004012005'/>
  <confidentialityCode code='N' displayName='Normal' codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality' />
  <languageCode code='en-US'/>

  <component>
    <structuredBody>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.6'/>
        <!-- Required Active Problems Section content -->
      </section>
    </component>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.8'/>
      <!-- Required Resolved Problems Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.7'/>
      <!-- Required Discharge Diagnosis Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.3'/>
      <!-- Required Admitting Diagnosis Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.21'/>
      <!-- Required if known Selected Meds Administered Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.22'/>
      <!-- Required Discharge Meds Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.20'/>
      <!-- Required if known Admission Medications Section content -->
    </section>

    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.13'/>
      <!-- Required Allergies Section content -->
    </section>
  </structuredBody>
</ClinicalDocument>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.5'/>
    <!-- Required Hospital Course Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.34'/>
    <!-- Optional Advance Directives Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.17'/>
    <!-- Required if known History of Present Illness Section content -->
  </section>
</component>

<component>
  <section>
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    <!-- Optional Functional Status Section content -->
  </section>
</component>

<component>
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    <!-- Optional Review of Systems Section content -->
  </section>
</component>

<component>
  <section>
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    <!-- Optional Physical Examination Section content -->
  </section>
</component>

<component>
  <section>
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    <!-- Optional Discharge Procedures Tests, Reports Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31'/>
    <!-- Required Plan of Care Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.33'/>
    <!-- Optional Discharge Diet Section content -->
  </section>
</component>
6.3.1.5 PHR Extract Specification 1.3.6.1.4.1.19376.1.5.3.1.1.5

The PHR Extract module describes the document content that summarizes information contained within a Personal Health Record. While a PHR can contain a great deal more information (including clinical documents, lab reported, images, trend data, monitoring data) et cetera, this content module only deals with the format of the summary information from the PHR.

An PHR Extract Module is a type of medical summary, and incorporates the constraints defined for Medical Summaries. While mappings have been provided to various standards, this content module conforms to the ASTM/HL7 Continuity of Care Document as well as this guide.

The following table describes the data elements that may be present in a PHR Extract. The first column of this table is drawn from the Common Data Elements in the PHR found in Appendix B of the AHIMA Report: The Role of the Personal Health Record in the EHR. Indented items in this column of the table provide more detail for the item they appear underneath.

These data elements were then mapped to the ASTM CCR, HL7 CDA, CRS and CCD and the implicit data elements referenced by the HL7 PHR Conformance Criteria.

A further requirement of transfers of information between PHR and EHR systems is that authorship of the information stored within the PHR shall be tracable through the various import/export cycles. PHR Manager Actors must be secure nodes, which requires logging of any updates to or accesses of PHR information. The DSG profile should be used to ensure that information coming into, or exiting these systems is verifiably authored.

### 6.3.1.5.1 Format Code

The XDSDocumentEntry format code for this content is urn:ihe:pcc:xphr:2007

### 6.3.1.5.2 Parent Template

This document is an instance of the Medical Summary template.

### 6.3.1.5.3 LOINC Code

The LOINC code for this document is 34133-9 Summary of Episode Note

### 6.3.1.5.4 Standards

<table>
<thead>
<tr>
<th>AHIMA-PHR</th>
<th>AHIMA PHR Common Data Elements</th>
</tr>
</thead>
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<tr>
<td>CDAR2</td>
<td>HL7 CDA Release 2.0</td>
</tr>
<tr>
<td>CRS</td>
<td>HL7 Care Record Summary</td>
</tr>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
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<td>HL7-PHR</td>
<td>HL7 PHR Functional Model (Draft)</td>
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<tr>
<td>---------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>LOINC</td>
<td>Logical Observation Identifier Names and Codes</td>
</tr>
</tbody>
</table>
6.3.1.5.5 Data Element Index

<table>
<thead>
<tr>
<th>AHIMA Common Data Elements</th>
<th>ASTM Continuity of Care Record</th>
<th>HL7 Clinical Document Architecture, Care Record Summary or Continuity of Care Document</th>
<th>HL7 PHR Conformance Criteria</th>
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</thead>
<tbody>
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<td>patientRole</td>
<td>Demographic Information</td>
</tr>
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<td>patient/name</td>
<td>Demographic Information</td>
</tr>
<tr>
<td>Address</td>
<td>Patient</td>
<td>patientRole/addr</td>
<td>Contact Information</td>
</tr>
<tr>
<td>Contact Information</td>
<td>Patient</td>
<td>patientRole/telecom</td>
<td>Contact Information</td>
</tr>
<tr>
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<td>Demographic Information</td>
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<td>(Religious Affiliation[1])</td>
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<td>patient/religiousAffiliationCode</td>
<td>Spiritual Affiliation / Considerations</td>
</tr>
<tr>
<td>Languages Spoken</td>
<td>Patient</td>
<td>patient/languageCommunication</td>
<td></td>
</tr>
<tr>
<td>Employer and School Contacts</td>
<td>Social History</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hazardous Working Conditions</td>
<td>Social History</td>
<td>HISTORY OF OCCUPATIONAL EXPOSURE</td>
<td></td>
</tr>
<tr>
<td>Emergency Contacts</td>
<td>Social History</td>
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<td>Health Insurance or Pharmacy Insurance</td>
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<td>performer</td>
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<td>ADVANCE DIRECTIVES</td>
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<td>General Medical Information Height, Weight</td>
<td>Vital Signs</td>
<td>VITAL SIGNS</td>
<td></td>
</tr>
<tr>
<td>Blood Type</td>
<td>Results</td>
<td>RELEVANT DIAGNOSTIC TESTS AND/OR LABORATORY DATA</td>
<td></td>
</tr>
<tr>
<td>Last Physical or Checkup</td>
<td>Encounters</td>
<td>HISTORY OF OUTPATIENT VISITS</td>
<td>Clinical Encounters and Procedures List</td>
</tr>
<tr>
<td>Allergies and Drug Sensitivities</td>
<td>Alerts</td>
<td>HISTORY OF ALLERGIES</td>
<td>Allergy and Reaction List</td>
</tr>
<tr>
<td>Conditions</td>
<td>Problems</td>
<td>HISTORY OF PAST ILLNESS - or - PROBLEM LIST</td>
<td>Problem List</td>
</tr>
<tr>
<td>Surgeries</td>
<td>Procedures</td>
<td>HISTORY OF SURGICAL PROCEDURES</td>
<td>Clinical Encounters and Procedures List</td>
</tr>
<tr>
<td>Medications – Prescription and Non-Prescription</td>
<td>Medications</td>
<td>HISTORY OF MEDICATION USE</td>
<td>Medication List</td>
</tr>
<tr>
<td>Immunizations</td>
<td>Immunizations</td>
<td>HISTORY OF IMMUNIZATIONS</td>
<td>Immunizations List</td>
</tr>
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<td>Doctor Visits</td>
<td>Encounters</td>
<td>HISTORY OF OUTPATIENT VISITS</td>
<td>Clinical Encounters and Procedures List</td>
</tr>
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<td>Encounters</td>
<td>HISTORY OF HOSPITALIZATIONS</td>
<td>Clinical Encounters and Procedures List</td>
</tr>
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<td>Other Healthcare Visits</td>
<td>Encounters</td>
<td>HISTORY OF OUTPATIENT VISITS</td>
<td>Clinical Encounters and Procedures List</td>
</tr>
<tr>
<td>Clinical Tests</td>
<td>Results</td>
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<td>Laboratory and Test Results</td>
</tr>
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<td>HISTORY OF PREGNANCIES</td>
<td></td>
</tr>
<tr>
<td>Medical Devices</td>
<td>Medical Devices</td>
<td>HISTORY OF MEDICAL DEVICE USE</td>
<td></td>
</tr>
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<td>Family Member History</td>
<td>Family History</td>
<td>HISTORY OF FAMILY MEMBER DISEASES</td>
<td>Family History</td>
</tr>
<tr>
<td>Foreign Travel</td>
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<td>HISTORY OF TRAVEL</td>
<td></td>
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<td>Therapy</td>
<td>Plan of Care</td>
<td>TREATMENT PLAN</td>
<td>Care Plans, Goals and Disease Management</td>
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<td>Vital signs</td>
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# 6.3.1.5.6 Specification

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<tr>
<td><strong>Legal Documents and Medical Directives</strong></td>
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<td>1.3.6.1.4.1.19376.1.5.3.1.3.13</td>
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<td>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3</td>
</tr>
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<td>O</td>
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</tr>
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</tr>
<tr>
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<td>R2</td>
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<tr>
<td>-----------------------------</td>
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</tr>
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<td>Foreign Travel</td>
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</tr>
<tr>
<td>Plan of Care</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.31</td>
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<td>Coded Vital signs</td>
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<td>Functional Status</td>
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</tbody>
</table>

**Table 6.3.1.5.6-1**

### 6.3.1.5.7 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate `<templateId>` elements in the header of the document. This is shown in the sample document below. A CDA Document may conform to more than one template. This content module inherits from the Medical Summary content module, and so must conform to the requirements of that template as well, thus all `<templateId>` elements shown in the example below shall be included.
<ClinicalDocument xmlns='urn:hl7-org:v3'>
	<typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
	<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.2'/>
	<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5'/>
	<id root=' ' extension=' '/>
	<code code='34133-9' displayName='Summary of Episode Note' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
	<title>PHR Extract</title>
	<effectiveTime value='20081004012005'/>
	<confidentialityCode code='N' displayName='Normal' codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality'/>
	<languageCode code='en-US'/>

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		<section>
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			<!-- Optional Hazardous Working Conditions Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.34'/>
			<!-- Required if known Legal Documents and Medical Directives Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.13'/>
			<!-- Required Allergies and Drug Sensitivities Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.8'/>
			<!-- Required Conditions Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.6'/>
			<!-- Required Conditions (cont) Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.12'/>
			<!-- Required if known Surgeries Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.19'/>
			<!-- Required Medications - Prescription and Non-Prescription Section content -->
		</section>
		<section>
			<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.23'/>
			<!-- Required if known Immunizations Section content -->
		</section>
	</component>
</ClinicalDocument>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3'/>
    <!-- Optional Doctor Visits / Last Physical or Checkup Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3'/>
    <!-- Optional Hospitalizations Section content -->
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</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3'/>
    <!-- Optional Other Healthcare Visits Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.4'/>
    <!-- Optional Clinical Tests / Blood Type Section content -->
  </section>
</component>

<component>
  <section>
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  </section>
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<component>
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  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.6'/>
    <!-- Optional Foreign Travel Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.6'/>
    <!-- Optional Plan of Care Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.2'/>
    <!-- Optional Coded Vital signs Section content -->
  </section>
</component>
6.3.1.5.8 Additional Constraints

The assignedAuthoring device shall be populated with information about the EHR and/or PHR which assisted in creation of the document.

All sections and entries within the document shall contain an <id> element.

6.3.1.6 PHR Update Specification 1.3.6.1.4.1.19376.1.5.3.1.1.6

The PHR Update Content Module is similar to the PHR Extract content module, except that it has a number of different constraints. First of all, it is not required to contain all of the information that the PHR Extract content module does. The reason for this is because the purpose of this module is to reflect the changes that should be made to a PHR based on a previously existing PHR Extract content module. So, while it makes use of the same data element index, almost all of the data elements are optional. The purpose of this module is to make it easier for an EHR to create content that can be used to update a PHR.

6.3.1.6.1 Format Code

The XDSDocumentEntry format code for this content is urn:ihe:pcc:xphr:2007

6.3.1.6.2 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate <templateId> elements in the header of the document. This is shown in the sample document below.
6.3.1.6.3 Requirements

The requirements of this module are that it support recording updates to the original PHR Extract. The PHR Extract is made up of a header, and several sections, each of which may contain one or more entries. Suggestions to add, remove or update a section or entry are described in more detail below.

6.3.1.6.4 Adding a New Section or Appending to an Existing Section

A PHR Reviewer Actor may suggest additional material for an existing or new section by simply adding that section to the PHR Update document.

6.3.1.6.5 Replacing a Section

A PHR Reviewer Actor may suggest a revision to a section in the PHR Extract by replacing that section. To replace a section, the PHR Reviewer Actor creates a section in the PHR Update document that is of the same type as the section to be replaced in the PHR Extract document, and adds a <ppc:replacementOf> element to that section to indicate the section that it replaces.

The replacementOf element is an extension to the CDA Release 2.0 standard, and is further described below in Appendix C Extensions to CDA Release 2.0.

6.3.1.6.6 Adding an Entry

A PHR Reviewer Actor may suggest a new entry be added to a section by simply including that entry in a like section in the PHR Update document.

6.3.1.6.7 Replacing or Removing an Entry

The PHR Review Actor can replace an existing entry by adding an entry of the same type with new or modified information, and including in that entry a <reference> element that has an <externalAct> element. The <id> element of the <externalAct> shall be that of the act that is being replaced.

6.3.1.6.8 Removing an Entry

The PHR Reviewer Actor can suggest that an entry be removed by replacing it with an act who <statusCode> element has been set to nullified.

6.3.1.6.9 Constraints

The LOINC document type code is the same as for the PHR Extract content module. The PHR Update Content module must record the PHR Extract which it is updating.
6.3.1.7 Emergency Department Referral Specification
1.3.6.1.4.1.19376.1.5.3.1.1.10

An ED Referral is a type of Referral Summary, and incorporates the constraints defined for Referral Summaries.

This use case is described fully in the EDR Profile in PCC TF-1. Briefly, it involves a collaborative transfer of care for the referral of a patient from a care provider to an emergency department. Using this use case the contents of documents used in collaborative transfers of care were discussed with physicians and nurses in detail to identify major sections. The sections identified by physicians during the use case exercise as important are listed in the table below.

Using this information from the use case, the following mappings were made to existing standards.

6.3.1.7.1 Format Code
The XDSDocumentEntry format code for this content is urn:ihe:pcc:edr:2007

6.3.1.7.2 Parent Template
This document is an instance of the Medical Summary template.

6.3.1.7.3 Data Element Index

<table>
<thead>
<tr>
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<th>HL7 Care Record Summary</th>
<th>CDA Release 2.0</th>
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</thead>
<tbody>
<tr>
<td>Reason for Referral</td>
<td>Reason for Referral</td>
<td>REASON FOR REFERRAL</td>
</tr>
<tr>
<td>History Present Illness</td>
<td>History of Present Illness</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
<tr>
<td>Active Problems</td>
<td>Conditions</td>
<td>PROBLEM LIST</td>
</tr>
<tr>
<td>Current Meds</td>
<td>Medications</td>
<td>HISTORY OF MEDICATION USE</td>
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<td>Allergies</td>
<td>Allergies and Adverse Reactions</td>
<td>HISTORY OF ALLERGIES</td>
</tr>
<tr>
<td>Resolved Problems</td>
<td>Conditions</td>
<td>HISTORY OF PAST ILLNESS</td>
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<td>List of Surgeries</td>
<td>Past Surgical History</td>
<td>HISTORY OF PRIOR SURGERIES</td>
</tr>
<tr>
<td>Immunizations</td>
<td>Immunizations</td>
<td>HISTORY OF IMMUNIZATIONS</td>
</tr>
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<td>Family History</td>
<td>Family History</td>
<td>HISTORY OF FAMILY ILLNESS</td>
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<td>Physical Exam</td>
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#### Physical Exam

<table>
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<tr>
<td>Relevant Surgical Procedures / Clinical Reports (including links)</td>
<td>Studies and Reports</td>
<td>Relevant Diagnostic Tests and/or Laboratory Data</td>
</tr>
<tr>
<td>Relevant Diagnostic Test and Reports (Lab, Imaging, EKG’s, etc.) including links.</td>
<td>Studies and Reports</td>
<td>Relevant Diagnostic Tests and/or Laboratory Data</td>
</tr>
<tr>
<td>Care Plan (new meds labs, or x-rays ordered)</td>
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<td>Treatment Plan</td>
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</tr>
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<td>Mode of Transport to the Emergency Department</td>
<td>Care Plan</td>
<td>Mode of Transport</td>
</tr>
<tr>
<td>Estimated Time of Arrival to the ED</td>
<td>Care Plan</td>
<td>Mode of Transport</td>
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<td>Advance Directives</td>
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<td>Advance Directives</td>
</tr>
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<td>Patient Administrative Identifiers</td>
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</tr>
<tr>
<td>Data needed for state and local referral forms, if different than above</td>
<td>Optional Sections</td>
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### 6.3.1.7.4 Specification

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<td>History Present Illness</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.4</td>
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<tr>
<td>Active Problems</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.6</td>
</tr>
<tr>
<td>Current Meds</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.19</td>
</tr>
<tr>
<td>Allergies</td>
<td>R</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.13</td>
</tr>
<tr>
<td>Resolved Problems</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.8</td>
</tr>
<tr>
<td>List of Surgeries</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.11</td>
</tr>
<tr>
<td>Immunizations</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.23</td>
</tr>
<tr>
<td>Family History</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.14</td>
</tr>
<tr>
<td>Social History</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.16</td>
</tr>
<tr>
<td>Pertinent Review of Systems</td>
<td>O</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.18</td>
</tr>
<tr>
<td>Vital Signs</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.25</td>
</tr>
<tr>
<td>Physical Exam</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.24</td>
</tr>
<tr>
<td>Relevant Diagnostic Results and/or Clinical Reports</td>
<td>R2</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.27</td>
</tr>
</tbody>
</table>

Includes Diagnostic Surgical Procedures, Clinical Reports and Diagnostic Tests and Results (Lab, Imaging, EKG’s, etc.) including links to relevant documents.

Care Plan | R2 | 1.3.6.1.4.1.19376.1.5.3.1.3.31 |
### (new meds, labs, or x-rays ordered)

| **Mode of Transport to the Emergency Department (includes ETA)** | R | 1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2 |
| **Proposed ED Disposition** | R2 | 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.10 |
| **Advance Directives** | R | 1.3.6.1.4.1.19376.1.5.3.1.3.34 |
| **Patient Administrative Identifiers** | R |  |
| **Pertinent Insurance Information** | R2 |  |
| **Data needed for state and local referral forms, if different than above** | R2 |  |

**Note:** *Highlighted* items in the table above are different from what appears in the XDS-MS profile. All other data elements have identical definitions.

### 6.3.1.7.5 Conformance

CDA Release 2.0 documents that conform to the requirements of this content module shall indicate their conformance by the inclusion of the appropriate `<templateId>` elements in the header of the document. This is shown in the sample document below. A CDA Document may conform to more than one template. This content module inherits from the Medical Summary content module, and so must conform to the requirements of that template as well, thus all `<templateId>` elements shown in the example below shall be included.
<ClinicalDocument xmlns='urn:hl7-org:v3'>
<typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.3'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10'/>
</ClinicalDocument>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.14'/>
    <!-- Required if known Family History Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.16'/>
    <!-- Required if known Social History Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18'/>
    <!-- Optional Pertinent Review of Systems Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.25'/>
    <!-- Required if known Vital Signs Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.24'/>
    <!-- Required if known Physical Exam Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.27'/>
    <!-- Required if known Relevant Diagnostic Results and/or Clinical Reports Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31'/>
    <!-- Required if known Care Plan Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.10.3.2'/>
    <!-- Required Mode of Transport to the Emergency Department (includes ETA) Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.10'/>
    <!-- Required if known Proposed ED Disposition Section content -->
  </section>
  </component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.34'/>
    <!-- Required Advance Directives Section content -->
  </section>
  </component>

  <component>
    <section>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.19'/>
      <!-- Optional Additional History Section content -->
    </section>
  </component>
6.3.1.8 History and Physical Specification

6.3.1.9 Antepartum Summary Specification

6.3.2 CDA Header Content Modules

6.3.2.1 Language Communication 1.3.6.1.4.1.19376.1.5.3.1.2.1

Languages spoken shall be recorded using the languageCommunication infrastructure class associated with the patient. The <languageCommunication> element describes the primary and secondary languages of communication for a person. When used, these shall be described using the languageCommunication element as follows.

6.3.2.1.1 Specification

```xml
<languageCommunication>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.1'/>
  <languageCode code='en-US'/>
  <modeCode code='' codeSystem='2.16.840.1.113883.5.60' codeSystemName='LanguageAbilityMode'/>
  <preferenceInd value='true|false'/>
</languageCommunication>
```

6.3.2.1.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.1'/>

The <templateId> element identifies this <languageCommunication> element for validation of the content. The templateId must have root='1.3.6.1.4.1.19376.1.5.3.1.2.1'.

6.3.2.1.3 <languageCode code=' '/>

This element describes the language code. It uses the same vocabulary described for the ClinicalDocument/languageCode element described in more detail in HL7 CRS: 2.1.1. This element is required.

6.3.2.1.4 <modeCode code='' codeSystem='2.16.840.1.113883.5.60' codeSystemName='LanguageAbilityMode'/>

This element describes the mode of use, and is only necessary when there are differences between expressive and receptive abilities. This element is optional. When not present, the assumption is that any further detail provided within the languageCommunication element refers to all common modes of communication. The coding system used shall be the HL7 LanguageAbilityMode vocabulary when this element is communicated.
6.3.2.1.5 <proficiencyLevelCode code=''
   codeSystem='2.16.840.1.113883.5.61'
   codeSystemName='LanguageProficiencyCode' />

This element describes the proficiency of the patient (with respect to the mode if specified). This element is optional. The coding system used shall be the HL7 LanguageProficiencyCode vocabulary when this element is communicated.

6.3.2.1.6 <preferenceInd value=''/>

This element shall be present on all languageCommunication elements when more than one is provided. It shall be valued "true" if this language is the patient’s preferred language for communication, or "false" if this is not the patient’s preferred language. More than one language may be preferred, and at least one must be preferred.

6.3.2.2 Employer and School Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.2

Employer and school informational contacts shall be recorded as participants in the CDA Header as demonstrated in the figure below. These contacts shall conform to the General Constraints found in HL7 CRS: 2.1.1 with respect to the requirements for name, address, telephone numbers and other contact information.

The figure below shows how the information for this element is coded, and further constraints are provided in the following sections.

6.3.2.2.1 Specification

<participant typeCode='IND'>
   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.2'/>
   <time>
      <low value=''/>
      <high value=''/>
   </time>
   <associatedEntity classCode='CON'>
      <id root='' extension=''/>
      <code code='EMPLOYER|SCHOOL|AFFILIATED' codeSystem='1.3.5.1.4.1.19376.1.5.3.3'
            codeSystemName='IHERoleCode'/>
      <associatedPerson><name>…</name></associatedPerson>
      <scopingOrganization>
         <name>…</name>
         <telecom value='' use=''/>
         <addr>…</addr>
      </scopingOrganization>
   </associatedEntity>
</participant>

6.3.2.2.2 <participant typeCode='IND'>

The typeCode of the participant shall be IND.

6.3.2.2.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.2'>

The <templateId> element identifies this <participant> as a school or employer contact for validation of the content. The templateId must have root='1.3.6.1.4.1.19376.1.5.3.1.2.2'.
6.3.2.4 `<time>`<low value=' '/><high value=' '/></time>

The time element indicates the start and stop time range for this contact. These dates shall correspond to the start and stop dates for employment, enrollment, or other affiliation with the organization described.

6.3.2.5 `<associatedEntity classCode='CON'>`

The `<associatedEntity>` element provides the contact information (classCode='CON') for the school, employer or affiliated organization.

6.3.2.6 `<code code='EMPLOYER|SCHOOL|AFFILIATED' codeSystem='1.3.5.1.4.1.19376.1.5.3.3' codeSystemName='IHERoleCode'>`

The code value shall indicate whether the participant is the employer, school or other affiliated (e.g., volunteer) organization. See also the [IHE Role Code Vocabulary](1.3.5.1.4.1.19376.1.5.3.3).

6.3.2.7 `<associatedPerson>`<name>…</name></associatedPerson>

This element should be present. When present is shall provide the name of a contact person within the organization.

6.3.2.8 `<scopingOrganization>`<name>…</name><telecom value= use=/><addr>…</addr></scopingOrganization>

This element shall be present, and shall provide the name, address and telephone number of the organization.

6.3.2.3 Healthcare Providers and Pharmacies 1.3.6.1.4.1.19376.1.5.3.1.2.3

Healthcare providers (including pharmacies) shall be recorded as described in CCD: 3.17. The identifier that the patient is known by to these providers may be included using the Patient Identifier extension described in Extensions to HL7 CDA Release 2.0. See the example shown in for use of this extension element.
6.3.2.3.1 Specification

The `<documentationOf>` element records the service events that were performed. This element shall be present.

6.3.2.3.2 `<documentationOf>`

The `<documentationOf>` element records the service events that were performed. This element shall be present.

6.3.2.3.3 `<serviceEvent classCode="PCPR">`

The `<serviceEvent>` element describes the activity being documented. This element shall be present, and shall have a classCode attribute of 'PCPR'.

6.3.2.3.4 `<effectiveTime><low value=""/><high value=""/></effectiveTime>`

The `<effectiveTime>` element records the time over which care provision activities are recorded in the document. There shall be a `<low>` element which records the starting date of care provision, and a `<high>` element which records the ending date of care provision. The ending date may extend into the future in the document describes care that is intended to be provided, but that has not actually occurred.

6.3.2.3.5 `<performer typeCode="PRF">`

The `<performer>` elements in the `<serviceEvent>` identify the providers of care. At least one `<performer>` element should be present. When a provider gives care over two distinct time intervals (e.g., as in the case of a specialist who treats the patient for short periods of time in different years), the provider may be recorded multiple times as a performer.
6.3.2.3.6 <functionCode code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>

The function of the provider in the care of the patient should be present, and will be described in the <functionCode> element. This may be used for example, to identify the primary care provider.

6.3.2.3.7 <time><low value=' '/><high value=' '/></time>

The <time> element is used to show the time period over which the provider gave care to the patient. The <low> and <high> elements must be present, and indicate the time over which care was (or is to be) provided.

6.3.2.3.8 <assignedEntity classCode='ASSIGNED'>

The <assignedEntity> element contains elements that identify the individual provider, and shall be present.

6.3.2.3.9 <id root=' ' extension=' '/>

The <id> element may be present and identifies the provider.

6.3.2.3.10<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>

The <code> element describes the type of provider and can be used to distinguish pharmacies from other providers.

6.3.2.3.11<addr></addr>

The <addr> element gives the address of the provider.

6.3.2.3.12<telecom value=' ' use=' '/>

The <telecom> element gives the telephone number of the provider.

6.3.2.3.13<assignedPerson><name></name></assignedPerson>

The providers name should be present. If not present, then the <scopingOrganization> shall be present (see below).

6.3.2.3.14<representedOrganization><name></name></representedOrganization>

This element should be present, and shall provide the name of the organization.

6.3.2.3.15<sdtc:patient><sdtc:id root=' ' extension=' '/></sdtc:patient>

The <sdtc:patient> element may be present to represent the patient's medical record number with the given provider. The root attribute of <sdtc:id> element shall be present and identifies the namespace used for the identifier. The extension attribute shall be present and is the patient's medical record or account number with the provider. This element is an HL7 extension to CDA Release 2.0.
6.3.2.4 Patient Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.4

Patient contacts are recorded as described in HL7 CCD: 3.3

6.3.2.4.1 Specification

The guardians of a patient shall be recorded in the <guardian> element beneath the <patient> element.

Other contacts are recorded as <participant> elements appearing in the document header. The classCode attribute shall be set to 'IND'.

The <templateId> element identifies this person as a patient contact and must be recorded exactly as shown above.

The <time> element may be present and indicates the time of the participation.

The <associatedEntity> element identifies the type of contact. The classCode attribute shall be present, and contains a value from the set AGNT, CAREGIVER, ECON, NOK, or PRS to identify contacts that are agents of the patient, care givers, emergency contacts, next of kin, or other relations respectively.
6.3.2.4.7 <code code='' displayName='' codeSystem='2.16.840.1.113883.5.111' codeSystemName='RoleCode'/>

The relationship between the patient and the guardian or other contact should be recorded in the <code> element. The code attribute is required and comes from the HL7 PersonalRelationshipRoleType vocabulary. The codeSystem attribute is required and shall be represented exactly as shown above.

6.3.2.4.8 <addr>

The address of the guardian or other contact should be present, and shall be represented as any other address would be in CDA.

6.3.2.4.9 <telecom>

The phone number of the guardian or other contact should be present, and shall be represented as any other phone number would be in CDA.

6.3.2.4.10 <guardianPerson><name/> or <assignedPerson><name/>

The name of the guardian or other contact shall be present, and shall be represented as any other name would be in CDA.

6.3.3 CDA Section Content Modules

This list defines the sections that may appear in a medical document. It is intended to be a comprehensive list of all document sections that are used by any content profile defined in the Patient Care Coordination Technical Framework. All sections shall have a narrative component that may be freely formatted into normal text, lists, tables, or other appropriate human-readable presentations. Additional subsections or entry content modules may be required.

6.3.3.1 Reasons for Care

The sections described below describe various reasons why healthcare is being provided to the patient.
6.3.3.1.1 Reason for Referral Section 1.3.6.1.4.1.19376.1.5.3.1.3.1

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The reason for referral section shall contain a narrative description of the reason that the patient is being referred.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42349-1</td>
<td>R</td>
<td>REASON FOR REFERRAL</td>
</tr>
</tbody>
</table>

Figure 6.3-8Sample Reason for Referral Section

6.3.3.1.2 Coded Reason for Referral Section 1.3.6.1.4.1.19376.1.5.3.1.3.2

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>Reason for Referral (1.3.6.1.4.1.19376.1.5.3.1.3.1)</td>
</tr>
<tr>
<td>General Description</td>
<td>This section shall include at least one entry describing the reason for referral as described in the Entry Content Module.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42349-1</td>
<td>R</td>
<td>REASON FOR REFERRAL</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.13</td>
<td>R</td>
<td>Simple Observations</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5</td>
<td>R</td>
<td>Conditions Entry</td>
</tr>
</tbody>
</table>

6.3.3.1.2.1 Parent Template

The parent of this template is Reason for Referral.
Figure 6.33-11 Sample Coded Reason for Referral Section

6.3.3.1.3 Chief Complaint Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>This contains a narrative description of the patient's chief complaint.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>CHIEF COMPLAINT</td>
</tr>
<tr>
<td>10154-3</td>
<td>R</td>
</tr>
</tbody>
</table>

Figure 6.3-12 Sample Chief Complaint Section

6.3.3.1.4 Hospital Admission Diagnosis Section 1.3.6.1.4.1.19376.1.5.3.1.3.3

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.3</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The hospital admitting diagnosis section shall contain a narrative description of the primary reason for admission to a hospital facility. It shall include entries for observations as described in the Entry Content Modules.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>HOSPITAL ADMISSION DX</td>
</tr>
<tr>
<td>46241-6</td>
<td>R</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td></td>
</tr>
</tbody>
</table>
### Figure 6.33-13 Sample Hospital Admission Diagnosis Section

2010

```xml
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.3'/>
    <id root=' ' extension=' '/>
    <code code='46241-6' displayName='HOSPITAL ADMISSION DX' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
    <entry>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2'/>
    </entry>
  </section>
</component>
```

2015

Figure 6.33-13 Sample Hospital Admission Diagnosis Section

2020 6.3.3.2 Other Condition Histories

The sections defined below provide historical information about the patient's conditions.

#### 6.3.3.2.1 History of Present Illness Section 1.3.6.1.4.1.19376.1.5.3.1.3.4

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The history of present illness section shall contain a narrative description of the sequence of events preceding the patient’s current complaints.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10164-2</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
</tbody>
</table>

2030

<component>

2035 6.3.3.2.2 Hospital Course Section 1.3.6.1.4.1.19376.1.5.3.1.3.5

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The hospital course section shall contain a narrative description of the sequence of events from admission to discharge in a hospital facility.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10164-2</td>
<td>HISTORY OF PRESENT ILLNESS</td>
</tr>
</tbody>
</table>

2040

Figure 6.33-14 Sample History of Present Illness Section

2045

Figure 6.33-14 Sample History of Present Illness Section
6.3.3.2.3 Active Problems Section 1.3.6.1.4.1.19376.1.5.3.1.3.6

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.6</td>
<td>The active problem section shall contain a narrative description of the conditions currently being monitored for the patient. It shall include entries for patient conditions as described in the Entry Content Module.</td>
</tr>
</tbody>
</table>

### Entries

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11450-4</td>
<td>R</td>
<td>PROBLEM LIST</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.2</td>
<td>R</td>
<td>Problem Concern Entry</td>
</tr>
</tbody>
</table>

#### 6.3.3.2.3.1 Parent Template

The parent of this template is CCD 3.5.
6.3.3.2.4 Discharge Diagnosis Section 1.3.6.1.4.1.19376.1.5.3.1.3.7

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.7</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The discharge diagnosis section shall contain a narrative description of the conditions that need to be monitored after discharge from the hospital and those that were resolved during the hospital course. It shall include entries for patient conditions as described in the Entry Content Module.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>11535-2</td>
<td>R</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.2</td>
<td>R</td>
</tr>
</tbody>
</table>

```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.7'/>
    <id root=' ' extension=' '/>
    <code code='11535-2' displayName='HOSPITAL DISCHARGE DX' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>
```

Figure 6.3-17 Sample Discharge Diagnosis Section

6.3.3.2.5 History of Past Illness Section 1.3.6.1.4.1.19376.1.5.3.1.3.8

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The History of Past Illness section shall contain a narrative description of the conditions the patient suffered in the past. It shall include entries for problems as described in the Entry Content Modules.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>11348-0</td>
<td>R</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.2</td>
<td>R</td>
</tr>
</tbody>
</table>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.8'/>
    <id root=' ' extension=' '/>
    <code code='11348-0' displayName='HISTORY OF PAST ILLNESS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
    <entry>
      <!-- Required Problem Concern Entry element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2' />
    </entry>
  </section>
</component>

Figure 6.3-18 Sample History of Past Illness Section

6.3.3.2.6 Encounter Histories Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>2.16.840.1.113883.10.20.1.3 (2.16.840.1.113883.10.20.1.3)</td>
</tr>
<tr>
<td>General Description</td>
<td>The encounter history section contains coded entries describing the patient history of encounters.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>46240-8</td>
<td>R</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.14</td>
<td>R</td>
<td>Encounters</td>
</tr>
</tbody>
</table>

6.3.3.2.6.1 Parent Template

The parent of this template is 2.16.840.1.113883.10.20.1.3.

Figure 6.33-19 Sample Encounter Histories Section
6.3.3.2.7 History of Outpatient Visits Section 1.3.6.1.4.1.19376.1.5.3.1.3.9

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The outpatients visit section shall contain a narrative description of the completed visits to ambulatory facilities.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>11346-4</td>
<td>R</td>
</tr>
</tbody>
</table>

Figure 6.3-20 Sample History of Outpatient Visits Section

6.3.3.2.8 History of Inpatient Visits Section 1.3.6.1.4.1.19376.1.5.3.1.3.10

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.10</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The inpatient admissions section shall contain a narrative description of the admissions and discharges to inpatient facilities.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>11336-5</td>
<td>R</td>
</tr>
</tbody>
</table>

Figure 6.33-21 Sample History of Inpatient Visits Section

6.3.3.2.9 List of Surgeries Section 1.3.6.1.4.1.19376.1.5.3.1.3.11

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.11</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.14 (2.16.840.1.113883.10.20.1.12)</td>
</tr>
<tr>
<td>General Description</td>
<td>The list of surgeries section shall contain a narrative description of the diagnostic and therapeutic operative procedures and associated anesthetic techniques the patient received in the past.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>47519-4</td>
<td>R</td>
</tr>
</tbody>
</table>
### 6.3.3.2.9.1 Parent Template

The parent of this template is [CCD 3.14](#).

```xml
<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.12'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.11'/>
    <id root=' ' extension=' '/>
    <code code='47519-4' displayName='HISTORY OF PROCEDURES' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>
```

**Figure 6.33-22 Sample List of Surgeries Section**

### 6.3.3.2.10 Coded List of Surgeries Section 1.3.6.1.4.1.19376.1.5.3.1.3.12

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.12</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parent Template</strong></td>
<td>List of Surgeries (1.3.6.1.4.1.19376.1.5.3.1.3.11)</td>
</tr>
<tr>
<td><strong>General Description</strong></td>
<td>The list of surgeries section shall include entries for procedures and references to procedure reports when known as described in the Entry Content Modules.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td>Opt</td>
</tr>
<tr>
<td><strong>Description</strong></td>
<td></td>
</tr>
<tr>
<td>47519-4</td>
<td>R</td>
</tr>
<tr>
<td>HISTORY OF PROCEDURES</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.19</td>
<td>R</td>
<td>Procedure Entry</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.4</td>
<td>R2</td>
<td>References Entry</td>
</tr>
</tbody>
</table>

### 6.3.3.2.10.1 Parent Template

The parent of this template is [List of Surgeries](#).
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.11'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.12'/>
    <id root=' ' extension=' '/>
    <code code='47519-4' displayName='HISTORY OF PROCEDURES' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
    <entry/>
    <entry/>
  </section>
</component>

Figure 6.3-23 Sample Coded List of Surgeries Section

6.3.3.2.11 Allergies and Other Adverse Reactions Section
1.3.6.1.4.1.19376.1.5.3.1.3.13

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.13</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.8 (2.16.840.1.113883.10.20.1.2)</td>
</tr>
<tr>
<td>General Description</td>
<td>The adverse and other adverse reactions section shall contain a narrative description of the substance intolerances and the associated adverse reactions suffered by the patient. It shall include entries for intolerances and adverse reactions as described in the Entry Content Modules.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>48765-2 R Allergies, adverse reactions, alerts</td>
</tr>
<tr>
<td>Entries</td>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.5.3 R Allergies and Intolerances Concern</td>
</tr>
</tbody>
</table>

6.3.3.2.11.1 Parent Template

The parent of this template is CCD 3.8. This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.2
<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.2'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.13'/>
    <id root=' ' extension=' '/>
    <code code='48765-2' displayName='Allergies, adverse reactions, alerts'
        codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </entry>
</section>

Figure 6.33-24 Sample Allergies and Other Adverse Reactions Section

### 6.3.3.2.12 Family Medical History Section 1.3.6.1.4.1.19376.1.5.3.1.3.14

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.14</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.6 (2.16.840.1.113883.10.20.1.4)</td>
</tr>
<tr>
<td>General Description</td>
<td>The family history section shall contain a narrative description of the genetic family members, to the extent that they are known, the diseases they suffered from, their ages at death, and other relevant genetic information.</td>
</tr>
</tbody>
</table>

### 6.3.3.2.12.1 Parent Template

The parent of this template is **CCD 3.6**.

<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.4'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.14'/>
    <id root=' ' extension=' '/>
    <code code='10157-6' displayName='HISTORY OF FAMILY MEMBER DISEASES'
        codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </entry>
</section>

Figure 6.33-25 Sample Family Medical History Section

### 6.3.3.2.13 Coded Family Medical History Section 1.3.6.1.4.1.19376.1.5.3.1.3.15

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.15</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>Family Medical History (1.3.6.1.4.1.19376.1.5.3.1.3.14)</td>
</tr>
<tr>
<td>General Description</td>
<td>The family history section shall include entries for family history as described in the Entry Content Modules.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>---------------</td>
<td>-----</td>
</tr>
<tr>
<td>10157-6</td>
<td>R</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.15</td>
<td>R</td>
<td>Family History Organizer</td>
</tr>
</tbody>
</table>

### 6.3.3.2.13.1 Parent Template

The parent of this template is [Family Medical History](#).

```xml
<component>
  <section>
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    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.15'/>
    <id root=' ' extension=' '/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.16'/>
    <text>
      Text as described above
    </text>
    <entry>
      <!-- Required Family History Organizer element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.15'/>
    </entry>
  </section>
</component>
```

**Figure 6.3-26 Sample Coded Family Medical History Section**

### 6.3.3.2.14 Social History Section 1.3.6.1.4.1.19376.1.5.3.1.3.16

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.16</td>
<td>The social history section shall contain a narrative description of the person’s beliefs, home life, community life, work life, hobbies, and risky habits.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>29762-2</td>
<td>R</td>
<td>SOCIAL HISTORY</td>
</tr>
</tbody>
</table>

### 6.3.3.2.14.1 Parent Template

The parent of this template is [CCD 3.7](#).
6.3.3.2.15 Functional Status Section 1.3.6.1.4.1.19376.1.5.3.1.3.17

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.17</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.4 (2.16.840.1.113883.10.20.1.5)</td>
</tr>
<tr>
<td>General Description</td>
<td>The functional status section shall contain a narrative description of capability of the patient to perform acts of daily living.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>47420-5</td>
</tr>
</tbody>
</table>

6.3.3.2.16 Review of Systems Section 1.3.6.1.4.1.19376.1.5.3.1.3.18

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.18</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The review of systems section shall contain a narrative description of the responses the patient gave to a set of routine questions on the functions of each anatomic body system.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>10187-3</td>
</tr>
</tbody>
</table>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18'/>
    <id root=' ' extension=' '/>
    <code code='10187-3' displayName='REVIEW OF SYSTEMS' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>

Figure 6.3-29 Sample Review of Systems Section

6.3.3.2.17 Hazardous Working Conditions Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.1

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>Hazardous working conditions contains a narrative description of the patient's hazardous risks.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>10161-8</td>
</tr>
<tr>
<td>Description</td>
<td>R HISTORY OF OCCUPATIONAL EXPOSURE</td>
</tr>
</tbody>
</table>

Figure 6.33-30 Sample Hazardous Working Conditions Section

6.3.3.2.18 Pregnancy History Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.4

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The pregnancy history section contains coded entries describing the patient history of pregnancies.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>10162-6</td>
</tr>
<tr>
<td>Description</td>
<td>R HISTORY OF PREGNANCIES</td>
</tr>
</tbody>
</table>

Entries

<table>
<thead>
<tr>
<th>Structure ID</th>
<th>R Pregnancy Observation</th>
</tr>
</thead>
</table>
6.3.3.2.19 Medical Devices Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.5

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>2.16.840.1.11383.10.20.1.7 (2.16.840.1.11383.10.20.1.7)</td>
</tr>
<tr>
<td>General Description</td>
<td>The medical devices section contains narrative text describing the patient history of medical device use.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>46264-8</td>
</tr>
<tr>
<td>Opt</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>HISTORY OF MEDICAL DEVICE USE</td>
</tr>
</tbody>
</table>

6.3.3.2.19.1 Parent Template

The parent of this template is 2.16.840.1.11383.10.20.1.7.

6.3.3.2.20 Foreign Travel Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.6

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.6</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The foreign travel section contains only narrative text describing the patient's travel history.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>10182-4</td>
</tr>
<tr>
<td>Opt</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>HISTORY OF TRAVEL</td>
</tr>
</tbody>
</table>

Figure 6.33-31 Sample Pregnancy History Section

Figure 6.33-32 Sample Medical Devices Section

Figure 6.33-32 Sample Medical Devices Section
6.3.3.3 Medications

This section contains section content modules that describe activities surrounding the use of medication.

### 6.3.3.3.1 Medications Section 1.3.6.1.4.1.19376.1.5.3.1.3.19

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.19</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.9 (2.16.840.1.113883.10.20.1.8)</td>
</tr>
<tr>
<td>General Description</td>
<td>The medications section shall contain a description of the relevant medications for the patient, e.g. an ambulatory prescription list. It shall include entries for medications as described in the Entry Content Module.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>Opt</td>
</tr>
<tr>
<td>10160-0</td>
<td>R</td>
</tr>
</tbody>
</table>

### 6.3.3.3.1.1 Parent Template

The parent of this template is [CCD 3.9](#).
Note: This LOINC code is typically used to record the current medication list found in an EHR.

6.3.3.3.2 Admission Medication History Section

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.20</th>
</tr>
</thead>
</table>

**General Description**
The admission medication history section shall contain a narrative description of the relevant medications administered to a patient prior to admission to a facility. It shall include entries for medication administration as described in the Entry Content Module.

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42346-7</td>
<td>R</td>
<td>MEDICATIONS ON ADMISSION</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.7</td>
<td>R</td>
<td>Medications</td>
</tr>
</tbody>
</table>

Figure 6.3-35 Sample Admission Medication History Section
6.3.3.3.3 Medications Administered Section 1.3.6.1.4.1.19376.1.5.3.1.3.21

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.21</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The medications administered section shall contain a narrative description of the relevant medications administered to a patient during the course of an encounter. It shall include entries for medication administration as described in the Entry Content Module.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>18610-6</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>MEDICATION ADMINISTERED</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.7</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>Medications</td>
</tr>
</tbody>
</table>

Figure 6.3-36 Sample Medications Administered Section

6.3.3.3.4 Hospital Discharge Medications Section 1.3.6.1.4.1.19376.1.5.3.1.3.22

<table>
<thead>
<tr>
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<th>1.3.6.1.4.1.19376.1.5.3.1.3.22</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The hospital discharge medications section shall contain a narrative description of the medications requested (ordered) to be administered to the patient after discharge from the hospital. It shall include entries for medication requests as described in the Entry Content Module.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>10183-2</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>HOSPITAL DISCHARGE MEDICATIONS</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.7</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>Medications</td>
</tr>
</tbody>
</table>
Figure 6.3-37 Sample Hospital Discharge Medications Section

Note: All medications in this section must have substanceAdministration/@moodCode = "INT"

6.3.3.3.5 Immunizations Section 1.3.6.1.4.1.19376.1.5.3.1.3.23

<table>
<thead>
<tr>
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<td>Parent Template</td>
<td>CCD 3.11 (2.16.840.1.113883.10.20.1.6)</td>
</tr>
<tr>
<td>General Description</td>
<td>The immunizations section shall contain a narrative description of the immunizations administered to the patient in the past. It shall include entries for medication administration as described in the Entry Content Modules.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>11369-6</td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>HISTORY OF IMMUNIZATIONS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.12</td>
<td>R</td>
</tr>
</tbody>
</table>

6.3.3.3.5.1 Parent Template

The parent of this template is CCD 3.11.

Figure 6.33-38 Sample Immunizations Section
### 6.3.3.4 Physical Exams

#### 6.3.3.4.1 Physical Exam Section 1.3.6.1.4.1.19376.1.5.3.1.3.24

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.24</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>General Description</strong></td>
<td>The physical exam section shall contain a narrative description of the patient’s physical findings.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>29545-1</td>
<td>R</td>
</tr>
</tbody>
</table>

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    <id root=' ' extension=' '/>
    <code code='29545-1' displayName='PHYSICAL EXAMINATION' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>
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**Figure 6.33-39 Sample Physical Exam Section**

#### 6.3.3.4.2 Physical Exam Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.15

<table>
<thead>
<tr>
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<tbody>
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</tr>
<tr>
<td><strong>General Description</strong></td>
<td>The physical exam section shall contain only the required and optional subsections performed.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>29545-1</td>
<td>R</td>
</tr>
<tr>
<td><strong>Entries</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td><strong>Subsections</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.25</td>
<td>O</td>
</tr>
<tr>
<td>Vital signs may be a subsection of the physical exam or they may stand alone</td>
<td></td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.16</td>
<td>O</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.48</td>
<td>O</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.17</td>
<td>O</td>
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<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.18</td>
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<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.21</td>
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<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.22</td>
<td>O</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.23</td>
<td>O</td>
</tr>
<tr>
<td>OID</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.24</td>
<td>Neck</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.25</td>
<td>Endocrine System</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.26</td>
<td>Thorax and Lungs</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.27</td>
<td>Chest Wall</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.28</td>
<td>Breasts</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.29</td>
<td>Heart</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.30</td>
<td>Respiratory System</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.31</td>
<td>Abdomen</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.32</td>
<td>Lymphatic System</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.33</td>
<td>Vessels</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.34</td>
<td>Musculoskeletal System</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.35</td>
<td>Neurologic System</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.36</td>
<td>Genitalia</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.9.37</td>
<td>Rectum</td>
</tr>
</tbody>
</table>

2560  **6.3.3.4.2.1 Parent Template**

The parent of this template is [1.3.6.1.4.1.19376.1.5.3.1.3.24](#).
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.24'/>
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    <text>
      Text as described above
    </text>
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</component>

<component>
  <section>
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    <!-- Optional Vital Signs Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.16'/>
    <!-- Optional General Appearance Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.48'/>
    <!-- Optional Visible Implanted Medical Devices Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.17'/>
    <!-- Optional Integumentary System Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.18'/>
    <!-- Optional Head Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.20'/>
    <!-- Optional Ears, Nose, Mouth and Throat Section content -->
  </section>
</component>

<component>
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    <!-- Optional Ears Section content -->
  </section>
</component>

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.22'/>
    <!-- Optional Nose Section content -->
  </section>
</component>
<section>
  <!-- Optional Mouth, Throat, and Teeth Section content -->
</section>

<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.23'/>

<component>
  <section>
    <!-- Optional Neck Section content -->
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</component>

<component>
  <section>
    <!-- Optional Endocrine System Section content -->
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  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.25'/>
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<component>
  <section>
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  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.26'/>
</component>

<component>
  <section>
    <!-- Optional Chest Wall Section content -->
  </section>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.27'/>
</component>

<component>
  <section>
    <!-- Optional Breasts Section content -->
  </section>
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</component>

<component>
  <section>
    <!-- Optional Heart Section content -->
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  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.29'/>
</component>

<component>
  <section>
    <!-- Optional Respiratory System Section content -->
  </section>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.30'/>
</component>

<component>
  <section>
    <!-- Optional Abdomen Section content -->
  </section>
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</component>

<component>
  <section>
    <!-- Optional Lymphatic System Section content -->
  </section>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.32'/>
</component>
Figure 6.33-40 Sample Physical Exam Section

6.3.3.4.3 Hospital Discharge Physical Exam Section

| Template ID | 1.3.6.1.4.1.19376.1.5.3.1.3.26 |
| General Description | The hospital discharge physical exam section shall contain a narrative description of the patient’s physical findings at discharge from a hospital facility. |
| LOINC Code | Opt | Description |
| 10184-0 | R | HOSPITAL DISCHARGE PHYSICAL |
Figure 6.33-91 Sample Hospital Discharge Physical Exam Section

6.3.3.4.4 Vital Signs Section 1.3.6.1.4.1.19376.1.5.3.1.3.25

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.25</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.12</td>
</tr>
<tr>
<td>General Description</td>
<td>The vital signs section shall contain a narrative description of the measurement results of a patient’s vital signs.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>R</td>
</tr>
<tr>
<td>8716-3</td>
<td>VITAL SIGNS</td>
</tr>
</tbody>
</table>

6.3.3.4.4.1 Parent Template

The parent of this template is CCD 3.12.

Figure 6.33-102 Sample Vital Signs Section

6.3.3.4.5 Coded Vital Signs Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.2

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>Vital Signs (1.3.6.1.4.1.19376.1.5.3.1.3.25)</td>
</tr>
<tr>
<td>General Description</td>
<td>The vital signs section contains coded measurement results of a patient’s vital signs.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>R</td>
</tr>
<tr>
<td>8716-3</td>
<td>VITAL SIGNS</td>
</tr>
<tr>
<td>Entries</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td>R</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.13.1</td>
<td>Vital Signs Organizer</td>
</tr>
</tbody>
</table>
6.3.3.4.5.1 Parent Template
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      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
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    </text>
  </section>
</component>

Figure 6.33-113 Sample Coded Vital Signs Section

6.3.3.4.6 General Appearance Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.16

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The general appearance section shall contain a description of the overall, visibly-apparent condition of the patient.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>10210-3</td>
<td>R</td>
</tr>
</tbody>
</table>

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      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>

Figure 6.33-124 Sample General Appearance Section

6.3.3.4.7 Visible Implanted Medical Devices Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.48

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.9.48</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The visible implanted medical devices section shall contain a description of the medical devices apparent on physical exam that have been inserted into the patient, whether internal or partially external.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>TBD</td>
<td>R</td>
</tr>
</tbody>
</table>

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6.3.3.4.8 Integumentary System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.17

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The integumentary system section shall contain a description of any type of integumentary system exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>29302-7</td>
<td>R</td>
</tr>
</tbody>
</table>

6.3.3.4.9 Head Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.18

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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The head section shall contain a description of any type of head exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>10199-8</td>
<td>R</td>
</tr>
</tbody>
</table>
6.3.3.4.10 Eyes Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.19

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The eyes section shall contain a description of any type of eye exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10197-2</td>
<td>R</td>
<td>EYE</td>
</tr>
</tbody>
</table>

Figure 6.33-48 Sample Eyes Section

6.3.3.4.11 Ears, Nose, Mouth and Throat Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.20

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.9.20</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The ears, nose, mouth, and throat section shall contain a description of any type of ears, nose, mouth, or throat exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11393-6</td>
<td>R</td>
<td>EARS and NOSE and MOUTH and THROAT</td>
</tr>
</tbody>
</table>

Figure 6.33-49 Sample Ears, Nose, Mouth and Throat Section

6.3.3.4.12 Ears Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.21

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The ears section shall contain a description of any type of ear exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
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<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10195-6</td>
<td>R</td>
<td>EAR</td>
</tr>
</tbody>
</table>
Figure 6.33-160 Sample Ears Section

6.3.3.4.13 Nose Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.22

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The nose section shall contain a description of any type of nose exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10203-8</td>
<td>R</td>
<td>NOSE</td>
</tr>
</tbody>
</table>

Figure 6.33-171 Sample Nose Section

6.3.3.4.14 Mouth, Throat and Teeth Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.23

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The mouth, throat, and teeth section shall contain a description of any type of mouth, throat, or teeth exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10201-2</td>
<td>R</td>
<td>MOUTH and THROAT and TEETH</td>
</tr>
</tbody>
</table>

Figure 6.33-182 Sample Mouth, Throat and Teeth Section
### Neck Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.24

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td><strong>General Description</strong></td>
<td>The neck section shall contain a description of any type of neck exam.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>11411-6</td>
<td>R</td>
</tr>
</tbody>
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    </text>
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#### Figure 6.33-193Sample Neck Section

### Endocrine System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.25

<table>
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<tr>
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</tr>
</thead>
<tbody>
<tr>
<td><strong>General Description</strong></td>
<td>The endocrine system section shall contain a description of any type of endocrine system exam.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>29307-6</td>
<td>R</td>
</tr>
</tbody>
</table>

```
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    <text>
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    </text>
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</component>
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#### Figure 6.33-204Sample Endocrine System Section

### Thorax and Lungs Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.26

<table>
<thead>
<tr>
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<th>1.3.6.1.4.1.19376.1.5.3.1.1.9.26</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>General Description</strong></td>
<td>The thorax and lungs section shall contain a description of any type of thoracic or lung exams.</td>
</tr>
<tr>
<td><strong>LOINC Code</strong></td>
<td><strong>Opt</strong></td>
</tr>
<tr>
<td>10207-9</td>
<td>R</td>
</tr>
</tbody>
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<component>
  <section>
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    <id root=' ' extension=' '/>
    <code code='10207-9' displayName='THORAX+LUNGS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
      <text>
      Text as described above
      </text>
  </section>
</component>

Figure 6.33-215 Sample Thorax and Lungs Section

6.3.3.4.18 Chest Wall Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.27

<table>
<thead>
<tr>
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<th>1.3.6.1.4.1.19376.1.5.3.1.1.9.27</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The chest wall section shall contain a description of any type of chest wall exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11392-8</td>
<td>R</td>
<td>CHEST WALL</td>
</tr>
</tbody>
</table>

<component>
  <section>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.27'/>
    <id root=' ' extension=' '/>
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      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
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      Text as described above
      </text>
  </section>
</component>

Figure 6.33-226 Sample Chest Wall Section

6.3.3.4.19 Breast Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.28

<table>
<thead>
<tr>
<th>Template ID</th>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The breast section shall contain a description of any type of breast exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10193-1</td>
<td>R</td>
<td>BREASTS</td>
</tr>
</tbody>
</table>

<component>
  <section>
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      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
      <text>
      Text as described above
      </text>
  </section>
</component>

Figure 6.33-237 Sample Breast Section
6.3.3.4.20 Heart Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.29

<table>
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<tr>
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</tr>
</thead>
<tbody>
<tr>
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<td>The heart section shall contain a description of any type of heart exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>10200-4</td>
<td>R</td>
</tr>
<tr>
<td></td>
<td>HEART</td>
</tr>
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    </text>
  </section>
</component>
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Figure 6.3-58 Sample Heart Section

6.3.3.4.21 Respiratory System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.30

<table>
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</tr>
</thead>
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<tr>
<td>General Description</td>
<td>The respiratory system section shall contain a description of any type of respiratory exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>11412-4</td>
<td>R</td>
</tr>
<tr>
<td></td>
<td>RESPIRATORY SYSTEM</td>
</tr>
</tbody>
</table>

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    </text>
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</component>
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Figure 6.3.3-59 Sample Respiratory System Section

6.3.3.4.22 Abdomen Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.31

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The abdomen system section shall contain a description of any type of abdominal exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>10191-5</td>
<td>R</td>
</tr>
<tr>
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<td>ABDOMEN</td>
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    </text>
  </section>
</component>
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Figure 6.3.3.3-59 Sample Abdomen Section
Figure 6.33-240 Sample Abdomen Section

6.3.3.4.23 Lymphatic System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.32

<table>
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</tr>
</thead>
<tbody>
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<td>General Description</td>
<td>The lymphatic system section shall contain a description of any type of lymphatic exam.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
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<tr>
<td>11447-0</td>
<td></td>
<td>HEMATOLOGIC+LYMPHATIC+IMMUNOLOGIC SYSTEM</td>
</tr>
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</table>

Figure 6.33-251 Sample Lymphatic System Section

6.3.3.4.24 Vessels Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.33

<table>
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<td>The vessels section shall contain a description of any type of vessels exam.</td>
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<table>
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<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
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<tbody>
<tr>
<td>10208-7</td>
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</tr>
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</table>

Figure 6.33-262 Sample Vessels Section
### Musculoskeletal System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.34

<table>
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</thead>
<tbody>
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<td>The musculoskeletal system section shall contain a description of any type of musculoskeletal exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>11410-8</td>
</tr>
<tr>
<td></td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>MUSCULOSKELETAL SYSTEM</td>
</tr>
</tbody>
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</section>
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**Figure 6.33-27** Sample Musculoskeletal System Section

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### Neurologic System Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.35

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<td>The neurologic system section shall contain a description of any type of neurologic exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>10202-0</td>
</tr>
<tr>
<td></td>
<td>R</td>
</tr>
<tr>
<td>Description</td>
<td>NEUROLOGIC SYSTEM</td>
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</section>
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**Figure 6.33-28** Sample Neurologic System Section

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### Genitalia Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.36

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</thead>
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<td>The genitalia section shall contain a description of any type of genital exam.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>11400-9</td>
</tr>
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<td>R</td>
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<td>Description</td>
<td>GENITALIA</td>
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</section>
</component>
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**Figure 6.33-29** Sample Genitalia System Section
Figure 6.33-295 Sample Genitalia Section

6.3.3.4.28 Rectum Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.37

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</tr>
</thead>
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<td>The rectum section shall contain a description of any type of rectal exam.</td>
</tr>
</tbody>
</table>

<table>
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<tr>
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<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10205-3</td>
<td>R</td>
<td>RECTUM</td>
</tr>
</tbody>
</table>

Figure 6.33-306 Sample Rectum Section

6.3.3.4.29 Extremities Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The Extremities section SHALL contain a description of any type of exam on the patient's extremities.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>10196-4</td>
<td>R</td>
<td>EXTREMITIES</td>
</tr>
</tbody>
</table>

Figure 6.33-317 Sample Extremities Section
6.3.3.5 Relevant Studies

6.3.3.5.1 Results Section 1.3.6.1.4.1.19376.1.5.3.1.3.27

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The results section shall contain a narrative description of the patient’s relevant studies.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>30954-2</td>
<td>R</td>
<td>STUDIES SUMMARY</td>
</tr>
</tbody>
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<component>
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    <text>
      Text as described above
    </text>
  </section>
</component>
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Figure 6.33-68Sample Results Section

6.3.3.5.2 Coded Results Section 1.3.6.1.4.1.19376.1.5.3.1.3.28

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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The results section shall contain a narrative description of the relevant diagnostic procedures the patient received in the past. It shall include entries for procedures and references to procedure reports when known as described in the Entry Content Modules.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>30954-2</td>
<td>R</td>
<td>STUDIES SUMMARY</td>
</tr>
</tbody>
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<th>Description</th>
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</thead>
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<td>Procedure Entry</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.4</td>
<td>R2</td>
<td>References Entry</td>
</tr>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.13</td>
<td>O</td>
<td>Simple Observation</td>
</tr>
</tbody>
</table>
**6.3.3.5.3 Hospital Studies Summary Section**

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The hospital studies summary section shall contain a narrative description of the relevant diagnostic procedures the patient received during the hospital admission.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11493-4</td>
<td>R</td>
<td>HOSPITAL DISCHARGE STUDIES SUMMARY</td>
</tr>
</tbody>
</table>

**6.3.3.5.3.1 Coded Hospital Studies Summary Section**

<table>
<thead>
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<th>1.3.6.1.4.1.19376.1.5.3.1.3.30</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>Hospital Studies Summary (1.3.6.1.4.1.19376.1.5.3.1.3.29)</td>
</tr>
<tr>
<td>General Description</td>
<td>The hospital studies summary section shall include entries for diagnostic procedures.</td>
</tr>
</tbody>
</table>
6.3.3.5.3.1.1 Parent Template

The parent of this template is [Hospital Studies Summary](#).

```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.29'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.30'/>
    <id root='' extension=' '/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.29'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.30'/>
    <id root='' extension=' '/>
    <code code='11493-4' displayName='HOSPITAL DISCHARGE STUDIES SUMMARY'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
    <entry>
      <!-- Required Procedure Entry element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.16'/>
    </entry>
    <entry>
      <!-- Required if known References Entry element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4'/>
    </entry>
  </section>
</component>
```

**Figure 6.33-331 Sample Coded Hospital Studies Summary Section**

### 6.3.3.6 Plans of Care

This section provides content modules for sections that describe the plan of care intended for the patient.

#### 6.3.3.6.1 Care Plan Section 1.3.6.1.4.1.19376.1.5.3.1.3.31

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.31</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td><a href="2.16.840.1.113883.10.20.1.10">CCD 3.16</a></td>
</tr>
<tr>
<td>General Description</td>
<td>The care plan section shall contain a narrative description of the expectations for care including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>18776-5</td>
<td>R</td>
<td>TREATMENT PLAN</td>
</tr>
</tbody>
</table>
6.3.3.6.1.1 Parent Template

The parent of this template is CCD 3.16.

```xml
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5'/>
    <id root=' ' extension=' '/>
    <code code='51847-2' displayName='ASSESSMENT AND PLAN' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>
```

**Figure 6.33-342 Sample Care Plan Section**

6.3.3.6.2 Assessment and Plan Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5</td>
<td>The assessment and plan section shall contain a narrative description of the assessment of the patient condition and expectations for care including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>51847-2</td>
<td>R</td>
<td>ASSESSMENT AND PLAN</td>
</tr>
</tbody>
</table>

```xml
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5'/>
    <id root=' ' extension=' '/>
    <code code='18776-5' displayName='TREATMENT PLAN' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>
```

**Figure 6.33-353 Sample Assessment and Plan Section**

6.3.3.6.3 Discharge Disposition Section 1.3.6.1.4.1.19376.1.5.3.1.3.32

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.3.32</td>
<td>The plan of care section shall contain a narrative description of the expectations for care including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient, specifically used in a discharge from a facility such as a hospital or nursing home.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>18776-5</td>
<td>R</td>
<td>TREATMENT PLAN</td>
</tr>
</tbody>
</table>
6.3.3.6.4 Discharge Diet Section 1.3.6.1.4.1.19376.1.5.3.1.3.33

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.33</th>
</tr>
</thead>
</table>

**General Description**
The discharge diet section shall contain a narrative description of the expectations for diet including proposals, goals, and order requests for monitoring, tracking, or improving the dietary control of the patient, specifically used in a discharge from a facility such as an emergency department, hospital, or nursing home.

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42344-2</td>
<td>R</td>
<td>DISCHARGE DIET</td>
</tr>
</tbody>
</table>

6.3.3.6.5 Advance Directives Section 1.3.6.1.4.1.19376.1.5.3.1.3.34

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.34</th>
</tr>
</thead>
</table>

**Parent Template**
CCD 3.2 (2.16.840.1.113883.10.20.1.1)

**General Description**
The advance directive section shall contain a narrative description of the list of documents that define the patient’s expectations and requests for care along with the locations of the documents.

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42348-3</td>
<td>R</td>
<td>ADVANCE DIRECTIVES</td>
</tr>
</tbody>
</table>

6.3.3.6.5.1 Parent Template
The parent of this template is CCD 3.2. This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.1
3340
<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.1'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.34'/>
    <id root=' ' extension=' '/>
    <code code='42348-3' displayName='ADVANCE DIRECTIVES'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>

Figure 6.33-386 Sample Advance Directives Section

6.3.3.6.6 Coded Advance Directives Section 1.3.6.1.4.1.19376.1.5.3.1.3.35

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.3.35</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>Advance Directives (1.3.6.1.4.1.19376.1.5.3.1.3.34)</td>
</tr>
<tr>
<td>General Description</td>
<td>The advance directive section shall include entries for references to consent and advance directive documents when known as described in the Entry Content Modules.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>42348-3</td>
<td>R</td>
<td>ADVANCE DIRECTIVES</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.13.7</td>
<td>R2</td>
<td>Advance Directive Observation</td>
</tr>
</tbody>
</table>

6.3.3.6.6.1 Parent Template

The parent of this template is Advance Directives.

3345
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.34'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.35'/>
    <id root=' ' extension=' '/>
    <code code='42348-3' displayName='ADVANCE DIRECTIVES'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
  <section>
    <entry>
      <!-- Required if known Advance Directive Observation element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.7'/>
    </entry>
  </section>
</component>

Figure 6.33-397 Sample Coded Advance Directives Section

6.3.3.6.7 Transport Mode Section 1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The transport mode section contains a description of the mode of transport and the time of departure or arrival of the patient to a facility.</td>
</tr>
</tbody>
</table>
6.3.3.7 Administrative and Other Information

6.3.3.7.1 Payers Section 1.3.6.1.4.1.19376.1.5.3.1.1.5.3.7

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.5.3.7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent Template</td>
<td>CCD 3.1 (2.16.840.1.113883.10.20.1.9)</td>
</tr>
<tr>
<td>General Description</td>
<td>The Payers section contains data on the patient’s payers, whether a ‘third party’ insurance, self-pay, other payer or guarantor, or some combination.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>48768-6</td>
<td>R</td>
<td>PAYMENT SOURCES</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.17</td>
<td>R2</td>
<td>Coverage Entry</td>
</tr>
</tbody>
</table>

6.3.3.7.1.1 Parent Template

The parent of this template is CCD 3.1.
<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.9'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.5.3.7'/>
    <id root=' ' extension=' '/>
    <code code='48768-6' displayName='PAYMENT SOURCES'
        codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
  </section>
</component>

Figure 6.33-79 Sample Payers Section

6.3.3.7.2 Referral Source Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.3

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.13.2.3</td>
<td>The Referral Source section shall contain a narrative description of the referral source of the patient. Patients who are not referred by a particular agency or health care provider should be designated as &quot;self referred&quot;.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11293-8</td>
<td>R</td>
<td>ED REFERRAL SOURCE</td>
</tr>
</tbody>
</table>

Figure 6.33-400 Sample Referral Source Section

6.3.3.7.3 Transport Mode Section 1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2

<table>
<thead>
<tr>
<th>Template ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2</td>
<td>The transport mode section contains a description of the mode of transport and the time of departure or arrival of the patient to a facility.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LOINC Code</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>11459-5</td>
<td>R</td>
<td>TRANSPORT MODE</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1</td>
<td>This entry provides coded values giving the mode and time of departure or arrival of the patient to a facility.</td>
</tr>
</tbody>
</table>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.3.2'/>
    <id root=' ' extension=' '/>
    <code code='11459-5' displayName='TRANSPORT MODE'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
    <text>
      Text as described above
    </text>
    <!-- Required Transport element -->
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1'/>
  </section>
</component>

Figure 6.33-411 Sample Transport Mode Section

6.3.3.7.4 ED Disposition Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.10

<table>
<thead>
<tr>
<th>Template ID</th>
<th>1.3.6.1.4.1.19376.1.5.3.1.1.13.2.10</th>
</tr>
</thead>
<tbody>
<tr>
<td>General Description</td>
<td>The ED Disposition section contains descriptions of the various components of ED Disposition, including disposition from the ED, time of disposition, intended transportation mode, time of transport, and the non-ED practitioner the patient's care will be transferred to.</td>
</tr>
<tr>
<td>LOINC Code</td>
<td>Opt</td>
</tr>
<tr>
<td>11302-7</td>
<td>R</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Entries</th>
<th>Opt</th>
<th>Description</th>
</tr>
</thead>
</table>
| 1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2 | R | **Encounter Disposition**
This required entry describes the expected or actual disposition of the patient after the emergency department encounter has been completed. |

Figure 6.33-422 Sample ED Disposition Section
6.3.4 CDA Entry Content Modules

6.3.4.1 Authors and Informants

Each clinical statement that can be made in a CDA Document or HL7 Version 3 message shall be attributable to one or more authors. These are found in <author> elements, either directly within the clinical statement, or in one of its ancestors in the XML document or message.

Each clinical statement may also contain information from zero or more informants. These are found in <informant> elements, again, either directly within the clinical statement, or in one of its ancestors in the XML document or message.

6.3.4.1.1 <author>

Authors shall be described in an <author> element that is either directly on the clinical statement, or which can be reached by one of its ancestors.

6.3.4.1.2 <time value=' '/>

The time of authorship shall be recorded in the <time> element.

6.3.4.1.3 <assignedAuthor> -OR- <assignedEntity1>

<id root=' ' extension=' '>
<addr></addr>
<telecom value=' ' use=' '>

In a CDA document details about the author are provided in the <assignedAuthor> element. In Version 3 messages, they are provided in the <assignedEntity1> element. The semantics are identical even though the element names differ. The identifier of the author, and their address and telephone number shall be present inside the <id>, <addr> and <telecom> elements.

6.3.4.1.4 <assignedPerson><name></name></assignedPerson>
<representedOrganization><name></name></representedOrganization>

The author's and/or the organization's name shall be present when the <author> element is present.

6.3.4.2 Linking Narrative and Coded Entries

This section defines a linking mechanism that allows entries or portions thereof to be connected to the text of the clinical document.

6.3.4.2.1 Standards

RIM HL7 Version 3 Reference Information Model
CDAR2 HL7 Clinical Document Architecture Release 2.0
6.3.4.2.2 Constraints for CDA

Elements within the narrative `<text>` will use the ID attribute to provide a destination for links. Elements within an `<entry>` will be linked to the text via a URI reference using this attribute as the fragment identifier. This links the coded entry to the specific narrative text it is related to within the CDA instance, and can be traversed in either direction. This serves three purposes:

1. It supports diagnostics during software development and testing.
2. It provides a mechanism to enrich the markup that can be supported in the viewing application.
3. It eliminates the need to duplicate content in two places, which prevents a common source of error, and eliminates steps needed to validate that content that should be identical in fact is.

Each narrative content element within CDA may have an ID attribute. This attribute is of type xs:ID. This means that each ID in the document must be unique within that document. Within an XML document, an attribute of type xs:ID must start with a letter, and may be followed one or more letters, digits, hyphens or underscores. Three different examples showing the use of the ID attribute, and references to it appear below:

<table>
<thead>
<tr>
<th>Use of ID</th>
<th>References to ID</th>
</tr>
</thead>
</table>
| `<tr ID='foo'>
  <td ID='bar'>Table Cell 1</td>
  <td>Table Cell 2</td>
</tr>` | `<code>
  <originalText>
    <reference value='#foo'></reference>
  </originalText>
</code>` |
| `<list>
  <item ID='baz'>List item 1</item>
</list>` | `<code>
  <originalText>
    <reference value='#baz'></reference>
  </originalText>
</code>` |
| `<paragraph ID='p-1'>A paragraph
  <content ID='c-1'>with content</content>
</paragraph>` | `<code>
  <originalText>
    <reference value='#p-1'></reference>
  </originalText>
</code>` |

### Table 6.3.4.2.2-1 Example Uses of ID

This allows the text to be located with a special type of URI reference, which simply contains a fragment identifier. This URI is local to the document and so just begins with a hash mark (#), and is followed by the value of the ID being referenced. Given one of these URIs stored in a variable named theURI, the necessary text value can be found via the following XPath expression:

```xml
string(//*[ID=substring-after('#',$theURI)])
```

The table below shows the result of this expression using the examples above:

<table>
<thead>
<tr>
<th>$theURI</th>
<th>Returned Value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>
3535 If your XSLT processor is schema aware, even more efficient mechanisms exist to locate the element than the above expression.

Having identified the critical text in the narrative, any elements using the HL7 CD datatype (e.g., `<code>`) can then contain a `<reference>` to the `<originalText>` found in the narrative. That is why, although CDA allows `<value>` to be of any type in `<entry>` elements, this profile restricts them to always be of xsi:type='CD'.

Now, given an item with an ID stored in a variable named theID all `<reference>` elements referring to it can be found via the following XPath expression:

```
//cda:reference[@URI=concat('#',$theID)]
```

### 6.3.4.2.3 Constraints for HL7 Version 3 Messages

3545 Unlike CDA entries, structured statements in HL7 Version 3 Messages do not have a related narrative text section. Therefore full text representations should be included in the `<text>` element care statement acts.

### 6.3.4.3 Severity 1.3.6.1.4.1.19376.1.5.3.1.4.1

Any condition or allergy may be the subject of a severity observation. This structure is included in the target act using the `<entryRelationship>` element defined in the CDA Schema.

The example below shows the recording the condition or allergy severity, and is used as the context for the following sections.

#### 6.3.4.3.1 Standards

<table>
<thead>
<tr>
<th>PatCareStruct</th>
<th>HL7 Care Provision Domain (DSTU)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
</tbody>
</table>

#### 6.3.4.3.2 Specification
This specification models a severity observation as a separate observation from the condition. While this model is different from work presently underway by various organizations (i.e., SNOMED, HL7, TermInfo), it is not wholly incompatible with that work. In that work, qualifiers may be used to identify severity in the coded condition observation, and a separate severity observation is no longer necessary. The use of qualifiers is not precluded by this specification. However, to support semantic interoperability between EMR systems using different vocabularies, this specification does require that severity information also be provided in a separate observation. This ensures that all EMR systems have equal access to the information, regardless of the vocabularies they support.

6.3.4.3.2.1 <entryRelationship typeCode='SUBJ' inversionInd='true'>

The related statement is made about the severity of the condition (or allergy). For CDA, this observation is recorded inside an <entryRelationship> element occurring in the condition, allergy or medication entry. The containing <entry> is the subject (typeCode='SUBJ') of this new observation, which is the inverse of the normal containment structure, thus inversionInd='true'. For HL7 Version 3 Messages this relationship is represented with a <sourceOf> element, however the semantics, typeCode, and inversionInd is unchanged.

6.3.4.3.2.2 <observation moodCode='EVN' classCode='OBS'>

The related statement is another event (moodCode='EVN') observing (<observation classCode='OBS'>) the severity of the (surrounding) related entry (e.g., a condition or allergy).

6.3.4.3.2.3 <templateId root='2.16.840.1.113883.10.20.1.55'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.1'/>

The <templateId> elements identifies this <observation> as a severity observation, allowing for validation of the content. As a side effect, readers of the CDA can quickly locate and identify severity observations. The templateId elements shown above must be present.
6.3.4.3.2.4 <code code='SEV' codeSystem='2.16.840.1.113883.5.4' 
displayName='Severity' codeSystemName='ActCode' />

This observation is of severity, as indicated by the <code> element listed above. This element is required. The code and codeSystem attributes shall be recorded exactly as shown above.

6.3.4.3.2.5 <text><reference value='#severity-2'/></text>

The <observation> element shall contain a <text> element. For CDA, the <text> elements shall contain a <reference> element pointing to the narrative where the severity is recorded, rather than duplicate text to avoid ambiguity. For HL7 Version 3 Messages, the <text> element should contain the full narrative text.

6.3.4.3.2.6 <statusCode code='completed'/>

The code attribute of <statusCode> for all severity observations shall be completed. While the <statusCode> element is required in all acts to record the status of the act, the only sensible value of this element in this context is completed.

6.3.4.3.2.7 <value xsi:type='CD' code='H|M|L' 
codeSystem='2.16.840.1.113883.5.1063' 
codeSystemName='SeverityObservation'>

The <value> element contains the level of severity. It is always represented using the CD datatype (xsi:type='CD'), even though the value may be a coded or uncoded string. If coded, it should use the HL7 SeverityObservation vocabulary (codeSystem='2.16.840.1.113883.5.1063') containing three values (H, M, and L), representing high, moderate and low severity depending upon whether the severity is life threatening, presents noticeable adverse consequences, or is unlikely substantially effect the situation of the subject.

6.3.4.4 Problem Status Observation 1.3.6.1.4.1.19376.1.5.3.1.4.1.1

Any problem or allergy observation may reference a problem status observation. This structure is included in the target observation using the <entryRelationship> element defined in the CDA Schema. The clinical status observation records information about the current status of the problem or allergy, for example, whether it is active, in remission, resolved, et cetera. The example below shows the recording of clinical status of a condition or allergy, and is used as the context for the following sections.

6.3.4.4.1 Standards

CCD ASTM/HL7 Continuity of Care Document
6.3.4.4.2 Specification

This CCD models a problem status observation as a separate observation from the problem, allergy or medication observation. While this model is different from work presently underway by various organizations (i.e., SNOMED, HL7, TermInfo), it is not wholly incompatible with that work. In that work, qualifiers may be used to identify problem status in the coded condition observation, and a separate clinical status observation is no longer necessary. The use of qualifiers in the problem observation is not precluded by this specification or by CCD. However, to support semantic interoperability between EMR systems using different vocabularies, this specification does require that problem status information also be provided in a separate observation. This ensures that all EMR systems have equal access to the information, regardless of the vocabularies they support.

6.3.4.4.3 <entryRelationship typeCode='REFR' inversionInd='false'>

The related statement is made about the clinical status of the problem or allergy. For CDA, this observation is recorded inside an <entryRelationship> element occurring in the problem or allergy. For HL7 Version 3 Messages, the <entryRelationship> tag name is <sourceOf>, though the typeCode and inversionInd attributes and other semantics remain the same. The containing observation refers to (typeCode='REFR') this new observation.

6.3.4.4.4 <observation moodCode='EVN' classCode='OBS'>

The related statement is another event (moodCode='EVN') observing (<observation classCode='OBS'>) the clinical status of the (surrounding) related observation (e.g., a problem or allergy).

6.3.4.4.5 <templateId root='2.16.840.1.113883.10.20.1.57'/>

These <templateId> elements identify this <observation> as a problem status observation, allowing for validation of the content.
6.3.4.4.6 <code code='33999-4' codeSystem='2.16.840.1.113883.6.1' displayName='Status' codeSystemName='LOINC' />

This observation is of clinical status, as indicated by the <code> element. This element must be present. The code and codeSystem shall be recorded exactly as shown above.

6.3.4.4.7 <text><reference value='#cstatus-2'/></text>

The <observation> element shall contain a <text> element that points to the narrative text describing the clinical status. For CDA, the <text> elements shall contain a <reference> element pointing to the narrative section (see Linking Narrative and Coded Entries), rather than duplicate text to avoid ambiguity. For HL7 Version 3 Messages, the <text> element SHALL contain the full narrative text.

6.3.4.4.8 <statusCode code='completed'/>

The code attribute of <statusCode> for all clinical status observations shall be completed. While the <statusCode> element is required in all acts to record the status of the act, the only sensible value of this element in this context is completed.

6.3.4.4.9 <value xsi:type='CE' code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>

The <value> element contains the clinical status. It is always represented using the CE datatype (xsi:type='CE'). It shall contain a code from the following set of values from SNOMED CT.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>55561003</td>
<td>Active</td>
</tr>
<tr>
<td>73425007</td>
<td>Inactive</td>
</tr>
<tr>
<td>90734009</td>
<td>Chronic</td>
</tr>
<tr>
<td>7087005</td>
<td>Intermittent</td>
</tr>
<tr>
<td>255227004</td>
<td>Recurrent</td>
</tr>
<tr>
<td>415684004</td>
<td>Rule out</td>
</tr>
<tr>
<td>410516002</td>
<td>Ruled out</td>
</tr>
<tr>
<td>413322009</td>
<td>Resolved</td>
</tr>
</tbody>
</table>

6.3.4.5 Health Status 1.3.6.1.4.1.19376.1.5.3.1.4.1.2

A problem observation may reference a health status observation. This structure is included in the target observation using the <entryRelationship> element defined in the CDA Schema. The health status observation records information about the current health status of the patient. The example below shows the recording the health status, and is used as the context for the following sections.
6.3.4.5.1 Specification

This specification models a health status observation as a separate observation about the patient.

6.3.4.5.2 <entryRelationship typeCode='REFR'>

The related statement is made about the health status of the patient. For CDA, this observation is recorded inside an <entryRelationship> element occurring in the observation. The contained observation is referenced (typeCode='REFR') by the observation entry. For HL7 Version 3 Messages, the entryRelationship tagName is sourceOf, though the typeCode and inversionInd attributes and other semantics remain the same.

6.3.4.5.3 <observation moodCode='EVN' classCode='OBS'>

The related statement is another event (moodCode='EVN') observing (<observation classCode='OBS'>) the health status of the patient.

6.3.4.5.4 <templateId root='2.16.840.1.113883.10.20.1.51'/>

The <templateId> element identifies this <observation> as a health status observation, allowing for validation of the content.

6.3.4.5.5 <code code='11323-3' displayName='Health Status' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />

This observation is of health status, as indicated by the <code> element. This element must be present. The code and codeSystem attributes shall be recorded exactly as shown above.
6.3.4.5.6 <text><reference value='#hstatus-2'/></text>

The <observation> element shall contain a <text> element that contains the narrative text describing the clinical status. For CDA, the <text> elements shall contain a <reference> element pointing to the narrative section (see Linking Narrative and Coded Entries, rather than duplicate text to avoid ambiguity. For HL7 Version 3 Messages, the <text> element shall contain the full narrative text.

6.3.4.5.7 <statusCode code='completed'/>

The code attribute of <statusCode> for all health status observations shall be completed. While the <statusCode> element is required in all acts to record the status of the act, the only sensible value of this element in this context is completed.

6.3.4.5.8 <value xsi:type='CE' code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>

The <value> element contains the clinical status. It is always represented using the CE datatype (xsi:type='CE').

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>81323004</td>
<td>Alive and well</td>
</tr>
<tr>
<td>313386006</td>
<td>In remission</td>
</tr>
<tr>
<td>162467007</td>
<td>Symptom free</td>
</tr>
<tr>
<td>161901003</td>
<td>Chronically ill</td>
</tr>
<tr>
<td>271593001</td>
<td>Severely ill</td>
</tr>
<tr>
<td>21134002</td>
<td>Disabled</td>
</tr>
<tr>
<td>161045001</td>
<td>Severely disabled</td>
</tr>
<tr>
<td>419099009</td>
<td>Deceased</td>
</tr>
</tbody>
</table>

6.3.4.6 Comments 1.3.6.1.4.1.19376.1.5.3.1.4.2

This entry allows for a comment to be supplied with each entry. For CDA this structure is usually included in the target act using the <entryRelationship> element defined in the CDA Schema, but can also be used in the <component> element when the comment appears within an <organizer>. The example below shows recording a comment for an <entry>, and is used as context for the following sections. For HL7 Version 3 Messages, this relationship is represented with the element <sourceOf>, although the remainder of the typecodes and semantics are unchanged.

Any condition or allergy may be the subject of a comment.

6.3.4.6.1 Standards

<table>
<thead>
<tr>
<th>CareStruct</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
</tbody>
</table>
6.3.4.6.3 <entryRelationship typeCode='SUBJ' inversionInd='true'> or <component typeCode='COMP'/>

A related statement is made about an act, or a cluster or battery of results. In CDA the former shall be recorded inside an <entryRelationship> element occurring at the end of the entry. The containing act is the subject (typeCode='SUBJ') of this new observation, which is the inverse of the normal containment structure, thus inversionInd='true'.

For HL7 Version 3 Messages, the relationship element is <sourceOf>, however the typeCode and inversionInd remain the same.

In the latter case, the comment shall be recorded inside a <component> element contained within the <organizer> element.
6.3.4.6.4 <act classCode='ACT' moodCode='EVN'>
The related statement is an event (moodCode='EVN') describing the act (classCode='ACT') of making an arbitrary comment or providing instruction on the related entry.

6.3.4.6.5 <templateId root='2.16.840.1.113883.10.20.1.40'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.2'/>
These <templateId> elements identify this <act> as a comment, allowing for validation of the content.

6.3.4.6.6 <code code='48767-8' displayName='Annotation Comment'
    codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
The <code> element indicates that this is a comment and shall be recorded as shown above. The codeSystem and code attributes shall use the values specified above.

6.3.4.6.7 <text><reference value='#comment-2'/></text>
The <text> element provides a way to represent the <reference> to the text of the comment in the narrative portion of the document. For CDA, this SHALL be represented as a <reference> element that points to the narrative text section of the CDA. The comment itself is not the act being coded, so it appears in the <text> of the <observation>, not as part of the <code>. For HL7 Version 3 Messages, the <text> element SHALL contain the full narrative text.

6.3.4.6.8 <statusCode code='completed' />
The code attribute of <statusCode> for all comments must be completed.

6.3.4.6.9 <author>
The comment may have an author.

6.3.4.6.10<time value=' '>
The time of the comment creation shall be recorded in the <time> element when the <author> element is present.

6.3.4.6.11<assignedAuthor>
    <id root=' ' extension=' '>
    <addr/>
    <telecom value=' ' use=' '>
    The identifier of the author, and their address and telephone number must be present inside the <id>, <addr> and <telecom> elements when the <author> element is present.
6.3.4.6.12<assignedPerson><name></name></assignedPerson>
<representedOrganization><name></name></representedOrganization>

The author's and/or the organization's name must be present when the <author> element is present.

6.3.4.7 Patient Medication Instructions 1.3.6.1.4.1.19376.1.5.3.1.4.3

Any medication may be the subject of further instructions to the patient, for example to indicate that it should be taken with food, et cetera.

This structure is included in the target substance administration or supply act using the <entryRelationship> element defined in the CDA Schema. The example below shows the recording of patient medication instruction for an <entry>, and is used as context for the following section.

6.3.4.7.1 Standards

Pharmacy  HL7 Pharmacy Domain (Normative)

6.3.4.7.2 Specification

```xml
<entry>
  <substanceAdministration classCode='SBADM' moodCode='EVN'>
    <entryRelationship typeCode='SUBJ' inversionInd='true'>
      <act classCode='ACT' moodCode='INT'>
        <templateId root='2.16.840.1.113883.10.20.1.49'/>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3'/>
        <code code='PINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'
          codeSystemName='IHEActCode' />
        <text><reference value='#comment-2'/></text>
        <statusCode code='completed' />
      </act>
    </entryRelationship>
  </substanceAdministration>
</entry>
```

6.3.4.7.3 <entryRelationship typeCode='SUBJ' inversionInd='true'>

Again, a related statement is made about the medication or immunization. This observation is recorded inside an <entryRelationship> element occurring at the end of the substance administration or supply entry. The containing <entry> is the subject (typeCode='SUBJ') of this new observation, which is the inverse of the normal containment structure, thus inversionInd='true'.

6.3.4.7.4 <act classCode='ACT' moodCode='INT'>

The related statement is the intent (moodCode='INT') on how the related entry is to be performed.
6.3.4.7.5 <templateId root='2.16.840.1.113883.10.20.1.49'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3'/>

These <templateId> elements identify this <act> as a medication instruction, allowing for validation of the content. As a side effect, readers of the CDA can quickly locate and identify medication instructions.

6.3.4.7.6 <code code='PINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'
  codeSystemName='IHEActCode'/>

The <code> element indicates that this is a patient medication instruction. This element shall be recorded exactly as specified above.

  Note: These values will be sent to HL7 for harmonization with the HL7 Act Vocabulary.

6.3.4.7.7 <text><reference value='#comment-2'/></text>

The <text> element indicates the text of the comment. For CDA, this SHALL be represented as a <reference> element that points at the narrative portion of the document. The comment itself is not the act being coded, so it appears in the <text> of the <observation>, not as part of the <code>. For HL7 Version 3 Messages, the full text SHALL be represented here.

6.3.4.7.8 <statusCode code='completed' />

The code attribute of <statusCode> for all comments must be completed.

6.3.4.8 Medication Fulfillment Instructions 1.3.6.1.4.1.19376.1.5.3.1.4.3.1

Any medication may be the subject of further instructions to the pharmacist, for example to indicate that it should be labeled in Spanish, et cetera.

This structure is included in the target substance administration or supply act using the <entryRelationship> element defined in the CDA Schema. The figure below is an example of recording an instruction for an <entry>, and is used as context for the following sections.

6.3.4.8.1 Standards

Pharmacy

HL7 Pharmacy Domain (Normative)

6.3.4.8.2 Specification
Again, a related statement is made about the medication or immunization. In CDA, this observation is recorded inside an \texttt{<entryRelationship>> element occurring at the end of the substance administration or supply entry. The containing \texttt{<act>> is the subject (typeCode='SUBJ') of this new observation, which is the inverse of the normal containment structure, thus inversionInd='true'. For HL7 Version 3 Messages, this relationship is represented with the \texttt{<sourceOf>> element however the semantics, typeCode, and inversionInd remain the same.

### 6.3.4.8.4 \texttt{<act classCode='ACT' moodCode='INT'>}

The related statement is the intent (moodCode='INT') on how the related entry is to be performed.

### 6.3.4.8.5 \texttt{<templateId root='2.16.840.1.113883.10.20.1.43'/>}

\texttt{<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3.1'/>}

These \texttt{<templateId>> elements identify this \texttt{<act>> as a medication fulfillment instruction, allowing for validation of the content.

### 6.3.4.8.6 \texttt{<code code='FINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2' codeSystemName='IHEActCode'/>}

The \texttt{<code>> element indicates that this is a medication fulfillment instruction. This element shall be recorded exactly as specified above.

\texttt{Note:} These values will be sent to HL7 for harmonization with the HL7 Act Vocabulary.

### 6.3.4.8.7 \texttt{<text><reference value='#comment-2'/></text>}

The \texttt{<text>> element contains a free text representation of the instruction. For CDA this SHALL contain a provides a \texttt{<reference>>element to the link text of the comment in the narrative portion of the document. The comment itself is not the act being coded, so it appears in the \texttt{<text>> of the \texttt{<observation>>}, not as part of the \texttt{<code>>. For HL7 Version 3 Messages, the full text SHALL be represented here.
6.3.4.8.8 <statusCode code='completed' />

The code attribute of <statusCode> for all comments must be completed.

6.3.4.9 External References 1.3.6.1.4.1.19376.1.5.3.1.4.4

CDA Documents may reference information contained in other documents. While CDA Release 2.0 supports references in content via the <linkHtml> element, this is insufficient for many EMR systems as the link is assumed to be accessible via a URL, which is often not the case. In order to link an external reference, one needs the document identifier, and access to the clinical system wherein the document resides. For a variety of reasons, it is desirable to refer to the document by its identity, rather than by linking through a URL.

1. The identity of a document does not change, but the URLs used to access it may vary depending upon location, implementation, or other factors.
2. Referencing clinical documents by identity does not impose any implementation specific constraints on the mechanism used to resolve these references, allowing the content to be implementation neutral. For example, in the context of an XDS Affinity domain the clinical system used to access documents would be an XDS Registry and one or more XDS Repositories where documents are stored. In other contexts, access might be through a Clinical Data Repository (CDR), or Document Content Management System (DCMS). Each of these may have different mechanisms to resolve a document identifier to the document resource.
3. The identity of a document is known before the document is published (e.g., in an XDS Repository, Clinical Data Repository, or Document Content Management System), but its URL is often not known. Using the document identity allows references to existing documents to be created before those documents have been published to a URL. This is important to document creators, as it does not impose workflow restrictions on how links are created during the authoring process.

Fortunately, CDA Release 2.0 also provides a mechanism to refer to external documents in an entry, as shown below.
6.3.4.9.1 Specification

6.3.4.9.2 <act classCode='ACT' moodCode='EVN'>

The external reference is an act that refers to documentation of an <act> (classCode='ACT'), that previously occurred (moodCode='EVN').

6.3.4.9.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4'/>

The <templateId> element identifies this <act> as a reference act, allowing for validation of the content. As a side effect, readers of the CDA can quickly locate and identify reference acts. The templateId must have root='1.3.6.1.4.1.19376.1.5.3.1.4.4'.

6.3.4.9.4 <id root=' ' extension=' '/>

The reference is an act of itself, and must be uniquely identified. If there is no explicit identifier for this act in the source EMR system, a GUID may be used for the root attribute, and the extension may be omitted. Although HL7 allows for multiple identifiers, this profile requires that one and only one be used.

6.3.4.9.5 <code nullFlavor='NA'/> 

The reference act has no code associated with it.

6.3.4.9.6 <text><reference value='#study-1'/></text>

In order to connect this external reference to the narrative text which it refers, the value of the <reference> element in the <text> element is a URI to an element in the CDA narrative of this document.

6.3.4.9.7 <reference typeCode='SPRT|REFR'>
<externalDocument classCode='DOC' moodCode='EVN'>

External references are listed as either supporting documentation (typeCode='SPRT') or simply reference material (typeCode='REFR') for the reader. If this distinction is not
supported by the source EMR system, the value of typeCode should be REFR. For CDA, the reference is indicated by a <reference> element containing an <externalDocument> element which documents (classCode='DOC') the event (moodCode='EVA'). For HL7 Version 3 Messages, the reference is represented with the element <sourceOf> and the external document is represented with a <act> element, however semantics, and attributes remain otherwise without change.

6.3.4.9.8 <id extension=' ' root=' '/>

The identifier of the document is supplied in the <id> element.

6.3.4.9.9 <text><reference value=' '/></text>

A link to the original document may be provided here. This shall be a URL where the referenced document can be located. For CDA, the link should also be present in the narrative inside the CDA Narrative in a <linkHTML> element.

6.3.4.10 Internal References 1.3.6.1.4.1.19376.1.5.3.1.4.4.1

CDA and HL7 Version 3 Entries may reference (point to) information contained in other entries within the same document or message as shown below.

6.3.4.10.1 Specification

For CDA the act being referenced appears inside a related entryRelationship. The type (typeCode) and direction (inversionInd) attributes will be specified in the entry content module that contains the reference. For HL7 Version 3 Messages, the relationship is indicated with a <sourceOf> element, however typeCodes and semantics remain unchanged.

6.3.4.10.2 <entryRelationship typeCode=' ' inversionInd='true|false'>

For CDA the act being referenced appears inside a related entryRelationship. The type (typeCode) and direction (inversionInd) attributes will be specified in the entry content module that contains the reference. For HL7 Version 3 Messages, the relationship is indicated with a <sourceOf> element, however typeCodes and semantics remain unchanged.

6.3.4.10.3 <act classCode=' ' moodCode=' '> The act being referred to can be any CDA Clinical Statement element type (act, procedure, observation, substanceAdministration, supply, et cetera). For compatibility with the Clinical Statement model the internal reference shall always use the <act> class, regardless of the XML element type of the act it refers to.

6.3.4.10.4 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4.1'/>

The <templateId> element identifies this as an internal reference that conforms to all rules specified in this section.
6.3.4.10.5 <id root=' ' extension=' '/>

This element shall be present. The root and extension attributes shall identify an element defined elsewhere in the same document.

6.3.4.10.6 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>

This element shall be present. It shall be valued when the internal reference is to element that has a <code> element, and shall have the same attributes as the <code> element in the act it references. If the element it references does not have a <code> element, then the nullFlavor attribute should be set to "NA".

6.3.4.11 Concern Entry 1.3.6.1.4.1.19376.1.5.3.1.4.5.1

This event (moodCode='EVN') represents an act (<act classCode='ACT'> of being concerned about a problem, allergy or other issue. The <effectiveTime> element describes the period of concern. The subject of concern is one or more observations about related problems (see 1.3.6.1.4.1.19376.1.5.3.1.4.5.2) or allergies and intolerances (see 1.3.6.1.4.1.19376.1.5.3.1.4.5.3). Additional references can be provided having additional information related to the concern. The concern entry allows related acts to be grouped. This allows representing the history of a problem as a series of observation over time, for example.

6.3.4.11.1 Standards

<table>
<thead>
<tr>
<th>CCD</th>
<th>ASTM/HL7 Continuity of Care Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>ClinStat</td>
<td>ClinStat HL7 Clinical Statement (DRAFT)</td>
</tr>
</tbody>
</table>
6.3.4.11.2 Specification

```xml
<act classCode='ACT' moodCode='EVN'>
  <templateId root='2.16.840.1.113883.10.20.1.27'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.1'/>
  <id root='' extension=''/>
  <code nullFlavor='NA'/>
  <statusCode code='active|suspended|aborted|completed'/>
  <effectiveTime>
    <low value=''/>
    <high value=''/>
  </effectiveTime>
  <!-- one or more entry relationships identifying problems of concern -->
  <entryRelationship typeCode='SUBJ' inversionInd='false'>
    <sourceOf typeCode='SUBJ' inversionInd='false'>
      ...</sourceOf>
  </entryRelationship>
  <!-- For HL7 Version 3 Messages
  <sourceOf typeCode='SUBJ' inversionInd='false'>
    ...</sourceOf>
  -->
  <!-- optional entry relationship providing more information about the concern -->
  <entryRelationship typeCode='REFR'>
    <sourceOf typeCode='REFR' inversionInd='false'>
      ...</sourceOf>
  </entryRelationship>
  <!-- For HL7 Version 3 Messages
  <sourceOf typeCode='REFR' inversionInd='false'>
    ...</sourceOf>
  -->
</act>
```

6.3.4.11.3 <act classCode='ACT' moodCode='EVN'>
All concerns reflect the act of recording (<act classCode='ACT'>) the event (moodCode='EVN') of being concerned about a problem, allergy or other issue about the patient condition.

6.3.4.11.4 <templateId root='2.16.840.1.113883.10.20.1.27'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.1'/>
These template identifiers indicates this entry conforms to the concern content module. This content module inherits constraints from the HL7 CCD Template for problem acts, and so also includes that template identifier.

6.3.4.11.5 <id root='' extension=''/>
This required element identifies the concern.

6.3.4.11.6 <code nullFlavor='NA'/>
The code is not applicable to a concern act, and so shall be recorded as shown above.

6.3.4.11.7 <statusCode code='active|suspended|aborted|completed'/>
The statusCode associated with any concern must be one of the following values:

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>active</td>
<td>A concern that is still being tracked.</td>
</tr>
</tbody>
</table>
suspended
A concern that is active, but which may be set aside. For example, this value might be used to suspend concern about a patient problem after some period of remission, but before assumption that the concern has been resolved.

aborted
A concern that is no longer actively being tracked, but for reasons other than because the problem was resolved. This value might be used to mark a concern as being aborted after a patient leaves care against medical advice.

completed
The problem, allergy or medical state has been resolved and the concern no longer needs to be tracked except for historical purposes.

**Note:** A concern in the "active" state represents one for which some ongoing clinical activity is expected, and that no activity is expected in other states. Specific uses of the suspended and aborted states are left to the implementation.

4.6.3.11.8 `<effectiveTime><low value=' '/><high value=' '/></effectiveTime>`

The `<effectiveTime>` element records the starting and ending times during which the concern was active. The `<low>` element shall be present. The `<high>` element shall be present for concerns in the completed or aborted state, and shall not be present otherwise.

4.6.3.11.9 <!-- 1..* entry relationships identifying problems of concern -->

`<entryRelationship type='SUBJ' inversionInd='false'>`

Each concern is about one or more related problems or allergies. This entry shall contain one or more problem or allergy entries that conform to the specification in section Problem Entry or Allergies and Intolerances. This is how a series of related observations can be grouped as a single concern.

For CDA this SHALL be represented with the `<entryRelationship>` element. For HL7 Version 3 Messages, this SHALL be represented as a `<sourceOf>` element. The typeCode SHALL be ‘SUBJ’ for both HL7 Version 3 and CDA. HL7 Version 3 additionally requires that inversionInd SHALL be ‘false’.

**Note:** The Allergy and Intolerances entry is a refinement of the Problem entry.

4.6.3.11.10 <!-- 0..n optional entry relationship providing more information about the concern -->

`<entryRelationship type='REFR' inversionInd='false'>`

Each concern may have 0 or more related references. These may be used to represent related statements such related visits. This may be any valid CDA clinical statement, and SHOULD be an IHE entry template. For CDA this SHALL be represented with the `<entryRelationship>` element. For HL7 Version 3 Messages, this SHALL be represented as a `<subjectOf>` element. The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘false’.

4.6.3.12 Problem Concern Entry 1.3.6.1.4.1.19376.1.5.3.1.4.5.2

This entry is a specialization of the Concern Entry, wherein the subject of the concern is focused on a problem. Elements shown in the example below in gray are explained in the Concern Entry.
6.3.4.12.1 Standards

<table>
<thead>
<tr>
<th>CCD</th>
<th>ASTM/HL7 Continuity of Care Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>ClinStat</td>
<td>HL7 Clinical Statement Pattern (Draft)</td>
</tr>
</tbody>
</table>

6.3.4.12.2 Parent Template

The parent of this template is Concern Entry. This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.27

6.3.4.12.3 Specification

```xml
<act classCode='ACT' moodCode='EVN'>
<templateId root='2.16.840.1.113883.10.20.1.27'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.1'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2'/>
<id root=' ' extension=' '/>
<code nullFlavor='NA'/>
<statusCode code='active|suspended|aborted|completed'/>
<effectiveTime>
<low value=' '/>
<high value=' '/>
</effectiveTime>
<!-- 1..* entry relationships identifying problems of concern -->
<entryRelationship type='SUBJ'>
<observation classCode='OBS' moodCode='EVN'/>
<templateID root='1.3.6.1.4.1.19376.1.5.3.1.4.5'/>
</entryRelationship>
<!-- optional entry relationship providing more information about the concern -->
<entryRelationship type='REFR'>
</act>
```

6.3.4.12.4

This entry has a template identifier of 1.3.6.1.4.1.19376.1.5.3.1.4.5.1, and is a subtype of the Concern Entry, and so must also conform to that specification, with the template identifier of 1.3.6.1.4.1.19376.1.5.3.1.4.5.2. These elements are required and shall be recorded exactly as shown above.

6.3.4.12.5

This entry shall contain one or more problem entries that conform to the Problem Entry template 1.3.6.1.4.1.19376.1.5.3.1.4.5. For CDA this SHALL be represented with the <observation> element. For HL7 Version 3 Messages, this SHALL be represented...
as a `<subjectOf>` element. The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘false’

**6.3.4.13 Allergy and Intolerance Concern 1.3.6.1.4.1.19376.1.5.3.1.4.5.3**

4230 This entry is a specialization of the Concern Entry, wherein the subject of the concern is focused on an allergy or intolerance. Elements shown in the example below in gray are explained in that entry.
6.3.4.13.1 Standards

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>ClinStat</td>
<td>HL7 Clinical Statement Pattern (Draft)</td>
</tr>
</tbody>
</table>

6.3.4.13.2 Parent Template

The parent of this template is [Concern Entry](#). This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.27

6.3.4.13.3 Specification

```xml
<act classCode='ACT' moodCode='EVN'>
  <templateId root='2.16.840.1.113883.10.20.1.27'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.1'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.3'/>
  <id root=' ' extension=' '/>
  <code nullFlavor='NA'/>
  <statusCode code='active|suspended|aborted|completed'/>
  <effectiveTime>
    <low value=' '/>
    <high value=' '/>
  </effectiveTime>
  <!-- 1..* entry relationships identifying allergies of concern -->
  <entryRelationship typeCode='SUBJ'>
    <observation classCode='OBS' moodCode='EVN'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.6'/>
  </entryRelationship>
  <!-- optional entry relationship providing more information about the concern -->
  <entryRelationship type='REFR'>
    ...
  </entryRelationship>
</act>
```

6.3.4.13.4

This entry has a template identifier of 1.3.6.1.4.1.19376.1.5.3.1.4.5.3, and is a subtype of the Concern entry, and so must also conform to the rules of the [Concern Entry](#). These elements are required and shall be recorded exactly as shown above.

6.3.4.13.5

This entry shall contain one or more allergy or intolerance entries that conform to the [Allergy and Intolerance Entry](#). For CDA this SHALL be represented with the
<entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element. The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘false’

6.3.4.14 Problem Entry 1.3.6.1.4.1.19376.1.5.3.1.4.5

This section makes use of the linking, severity, clinical status and comment content specifications defined elsewhere in the technical framework. In HL7 RIM parlance, observations about a problem, complaint, symptom, finding, diagnosis, or functional limitation of a patient is the event (moodCode='EVN') of observing (<observation classCode='OBS'>) that problem. The <value> of the observation comes from a controlled vocabulary representing such things. The <code> contained within the <observation> describes the method of determination from yet another controlled vocabulary. An example appears below in the figure below.

6.3.4.14.1 Standards

<table>
<thead>
<tr>
<th>Standard</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>ClinStat</td>
<td>HL7 Clinical Statement Pattern (Draft)</td>
</tr>
</tbody>
</table>

6.3.4.14.2 Parent Template

This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.28

6.3.4.14.3 Specification
6.3.4.14.4 <observation classCode='OBS' moodCode='EVN' negationInd='false|true'>

The basic pattern for reporting a problem uses the CDA <observation> element, setting the classCode='OBS' to represent that this is an observation of a problem, and the moodCode='EVN', to represent that this is an observation that has in fact taken place. The negationInd attribute, if true, specifies that the problem indicated was observed to not have occurred (which is subtly but importantly different from having not been observed). The value of negationInd should not normally be set to true. Instead, to record that there is "no prior history of chicken pox", one would use a coded value indicated exactly that.

However, it is not always possible to record problems in this manner, especially if using a controlled vocabulary that does not supply pre-coordinated negations, or which do not allow the negation to be recorded with post-coordinated coded terminology.

6.3.4.14.5 <templateId root='2.16.840.1.113883.10.20.1.28'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5'/>

These <templateId> elements identify this <observation> as a problem, under both IHE and CCD specifications. This SHALL be included as shown above.

6.3.4.14.6 <id root='' extension=' '/>

The specific observation being recorded must have an identifier (<id>) that shall be provided for tracking purposes. If the source EMR does not or cannot supply an intrinsic identifier, then a GUID shall be provided as the root, with no extension (e.g., <id root='CE1215CD-69EC-4C7B-805F-569233C5E159'/>). At least one identifier must be present, more than one may appear.

6.3.4.14.7 <code code=' ' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>

The <code> describes the process of establishing a problem. The code element should be used, as the process of determining the value is important to clinicians (e.g., a diagnosis
is a more advanced statement than a symptom). The recommended vocabulary for
describing problems is shown in the table below. Subclasses of this content module may
specify other vocabularies. When the list below is used, the codeSystem is
'2.16.840.1.113883.6.96' and codeSystemName is SNOMED CT.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>64572001</td>
<td>Condition</td>
</tr>
<tr>
<td>418799008</td>
<td>Symptom</td>
</tr>
<tr>
<td>404684003</td>
<td>Finding</td>
</tr>
<tr>
<td>409586006</td>
<td>Complaint</td>
</tr>
<tr>
<td>248536006</td>
<td>Functional limitation</td>
</tr>
<tr>
<td>55607006</td>
<td>Problem</td>
</tr>
<tr>
<td>282291009</td>
<td>Diagnosis</td>
</tr>
</tbody>
</table>

6.3.4.14.8 <text><reference value=' '/></text>

The <text> element is required and points to the text describing the problem being
recorded; including any dates, comments, et cetera. The <reference> contains a URI in
value attribute. This URI points to the free text description of the problem in the
document that is being described.

6.3.4.14.9 <statusCode code='completed'/>

A clinical document normally records only those condition observation events that have
been completed, not observations that are in any other state. Therefore, the <statusCode>
shall always have code='completed'.

6.3.4.14.10 <effectiveTime><low value=' '/><high value=' '/></effectiveTime>

The <effectiveTime> of this <observation> is the time interval over which the
<observation> is known to be true. The <low> and <high> values should be no more
precise than known, but as precise as possible. While CDA allows for multiple
mechanisms to record this time interval (e.g. by low and high values, low and width, high
and width, or center point and width), we are constraining Medical summaries to use only
the low/high form. The <low> value is the earliest point for which the condition is known
to have existed. The <high> value, when present, indicates the time at which the
observation was no longer known to be true. Thus, the implication is made that if the
<hight> value is specified, that the observation was no longer seen after this time, and it
thus represents the date of resolution of the problem. Similarly, the <low> value may
seem to represent onset of the problem. Neither of these statements is necessarily precise,
as the <low> and <high> values may represent only an approximation of the true onset
and resolution (respectively) times. For example, it may be the case that onset occurred
prior to the <low> value, but no observation may have been possible before that time to
discern whether the condition existed prior to that time. The <low> value should
normally be present. There are exceptions, such as for the case where the patient may be
able to report that they had chicken pox, but are unsure when. In this case, the
<effectiveTime> element shall have a <low> element with a nullFlavor attribute set to 'UNK'. The <high> value need not be present when the observation is about a state of the patient that is unlikely to change (e.g., the diagnosis of an incurable disease).

6.3.4.14.11  <value xsi:type='CD' code=' ' codeSystem=' ' codeSystemName=' ' displayName=' '>

The <value> is the condition that was found. This element is required. While the value may be a coded or an un-coded string, the type is always a coded value (xsi:type='CD'). If coded, the code and codeSystem attributes shall be present. The codeSystem should reference a controlled vocabulary describing problems, complaints, symptoms, findings, diagnoses, or functional limitations, e.g., ICD-9, SNOMED-CT or MEDCIN, or others.

The table below is an incomplete listing of acceptable values for the codeSystem attribute, along with the codeSystemName.

<table>
<thead>
<tr>
<th>CodeSystem</th>
<th>codeSystemName</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.16.840.1.113883.6.96</td>
<td>SNOMED-CT</td>
<td>SNOMED Controlled Terminology</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.103</td>
<td>ICD-9CM (diagnoses)</td>
<td>International Classification of Diseases, Clinical Modifiers, Version 9</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.26</td>
<td>MEDCIN</td>
<td>A classification system from MEDICOMP Systems.</td>
</tr>
</tbody>
</table>

It is recommended that the codeSystemName associated with the codeSystem, and the displayName for the code also be provided for diagnostic and human readability purposes, but this is not required by this profile.

If uncoded, all attributes other than xsi:type='CD' must be absent.

In cases where information about a problem or allergy is unknown or where there are no problems or allergies, an entry shall use codes from the table below to record this fact:

<table>
<thead>
<tr>
<th>Entry Type</th>
<th>Code</th>
<th>Display Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Problem</td>
<td>396782006</td>
<td>Past Medical History Unknown</td>
<td>To indicate unknown medical history</td>
</tr>
<tr>
<td>Problem</td>
<td>407559004</td>
<td>Family History Unknown</td>
<td>To indicate that the patient's family history is not known.</td>
</tr>
<tr>
<td>Problem</td>
<td>160243008</td>
<td>No Significant Medical History</td>
<td>To indicate no relevant medical history</td>
</tr>
<tr>
<td>Problem</td>
<td>160245001</td>
<td>No current problems or disability</td>
<td>To indicate that the patient has no current problems (as distinct from no history).</td>
</tr>
<tr>
<td>Allergy</td>
<td>409137002</td>
<td>No Known Drug Allergies</td>
<td>To indicate that there are no known Drug allergies for this patient.</td>
</tr>
<tr>
<td>Allergy</td>
<td>160244002</td>
<td>No Known Allergies</td>
<td>To indicate that there are no known allergies for this patient.</td>
</tr>
<tr>
<td>Allergy</td>
<td>64970000</td>
<td>Substance Type Unknown</td>
<td>To indicate the state where there is a known allergy or intolerance to an unknown substance</td>
</tr>
</tbody>
</table>
6.3.4.14.12  <originalText><reference value=' /'></originalText>

The <value> contains a <reference> to the <originalText> in order to link the coded value to the problem narrative text (minus any dates, comments, et cetera). The <reference> contains a URI in value attribute. This URI points to the free text description of the problem in the document that is being described.

6.3.4.14.13 <!-- zero or one <entryRelationship typeCode='SUBJ' inversionInd='true'> elements containing severity -->

An optional <entryRelationship> element may be present indicating the severity of the problem. When present, this <entryRelationship> element shall contain a severity observation conforming to the Severity entry template (1.3.6.1.4.1.19376.1.5.3.1.4.1).

For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <subjectOf> element. The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘true’.

6.3.4.14.14 <!-- zero or one <entryRelationship typeCode='REFR' inversionInd='false'> elements containing clinical status -->

An optional <entryRelationship> may be present indicating the clinical status of the problem, e.g., resolved, in remission, active. When present, this <entryRelationship> element shall contain a clinical status observation conforming to the Problem Status Observation template (1.3.6.1.4.1.19376.1.5.3.1.4.1).

For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element. The typeCode SHALL be ‘REFR’ and inversionInd SHALL be ‘false’.

6.3.4.14.15 <!-- zero or one <entryRelationship typeCode='REFR' inversionInd='false'> elements identifying the health status of concern -->

An optional <entryRelationship> may be present referencing the health status of the patient, e.g., resolved, in remission, active. When present, this <entryRelationship> element shall contain a clinical status observation conforming to the Health Status Observation template (1.3.6.1.4.1.19376.1.5.3.1.4.1). The typeCode SHALL be ‘REFR’ and inversionInd SHALL be ‘false’.

For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element.

6.3.4.14.16 <!-- zero to many <entryRelationship typeCode='SUBJ' inversionInd='true'> element containing comments -->

One or more optional <entryRelationship> elements may be present providing an additional comments (annotations) for the condition. When present, this <entryRelationship> element shall contain a comment observation conforming to the Comment entry template (1.3.6.1.4.1.19376.1.5.3.1.4.2). The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘true’.
For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element.

### 6.3.4.15 Allergies and Intolerances 1.3.6.1.4.1.19376.1.5.3.1.4.6

Allergies and intolerances are special kinds of problems, and so are also recorded in the CDA <observation> element, with classCode='OBS'. They follow the same pattern as the problem entry, with exceptions noted below.

#### 6.3.4.15.1 Standards

<table>
<thead>
<tr>
<th>CCD</th>
<th>ASTM/HL7 Continuity of Care Document</th>
</tr>
</thead>
<tbody>
<tr>
<td>CareStruct</td>
<td>HL7 Care Provision Care Structures (DSTU)</td>
</tr>
<tr>
<td>ClinStat</td>
<td>HL7 Clinical Statement Pattern (Draft)</td>
</tr>
</tbody>
</table>

#### 6.3.4.15.2 Specification

```xml
<observation classCode='OBS' moodCode='EVN' negationInd='false'>
  <templateId root='2.16.840.1.113883.10.20.1.18'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.6'/>
  <id root=' ' extension=' '/>
  <code code='ALG|OINT|DALG|EALG|FALG|DINT|EINT|FINT|DNAINT|ENAINT|FNAINT' codeSystem='2.16.840.1.113883.5.4'
    codeSystemName='ObservationIntoleranceType'/>
  <text><reference value=' '/></text>
  <statusCode code='completed'/>
  <effectiveTime>
    <low value=' '/>
    <high value=' '/>
  </effectiveTime>
  <participant typeCode='CSM'>
    <participantRole classCode='MANU'>
      <playingEntity classCode='MMAT'>
        <code code=' ' codeSystem=' '>
          <originalText><reference value='#substance'/></orginalText>
        </code>
      </playingEntity>
    </participantRole>
  </participant>
</observation>
```

This entry has a template identifier of 1.3.6.1.4.1.19376.1.5.3.1.4.6, and is a subtype of the **Problem Entry**, and so must also conform to the rules of the problem entry, which has the template identifier of 1.3.6.1.4.1.19376.1.5.3.1.4.5.5. These elements are required and shall be recorded exactly as shown above.
6.3.4.15.4 <code>

code='ALG|OINT|DINT|EINT|FINT|DALG|EALG|FALG|DNAINT|ENAINT|FNAINT' displayName=' ' codeSystem='2.16.840.1.113883.5.4'

codeSystemName='ObservationIntoleranceType'/>

The <code> element represents the kind of allergy observation made, to a drug, food or environmental agent, and whether it is an allergy, non-allergy intolerance, or unknown class of intolerance (not known to be allergy or intolerance). The <code> element of an allergy entry shall be provided, and a code and codeSystem attribute shall be present. The example above uses the HL7 ObservationIntoleranceType vocabulary domain, which does provide suitable observation codes. Other vocabularies may be used, such as SNOMED-CT or MEDCIN. The displayName and codeSystemName attributes should be present.

6.3.4.15.5 <value xsi:type='CD' code=' ' codeSystem=' ' codeSystemName=' '>

The <value> is a description of the allergy or adverse reaction. While the value may be a coded or an uncoded string, the type is always a coded value (xsi:type='CD'). If coded, the code and codeSystem attributes must be present. The codingSystem should reference a controlled vocabulary describing allergies and adverse reactions, see Table 5.4 12 above. If uncoded, all attributes other than xsi:type='CD' must be absent. The allergy or intolerance may not be known, in which case that fact shall be recorded appropriately. This might occur in the case where a patient experiences an allergic reaction to an unknown substance.

6.3.4.15.6 <participant typeCode='CSM'>

The substance that causes the allergy or intolerance may be specified in the <participant> element.

6.3.4.15.7 <code code=' ' codeSystem=' '>

The <code> element shall be present. It may contain a code and codeSystem attribute to indicate the code for the substance causing the allergy or intolerance. It shall contain a <reference> to the <originalText> in the narrative where the substance is named.

6.3.4.15.8 <!-- zero to many <entryRelationship> elements containing reactions -->

An allergy entry can record the reactions that are manifestations of the allergy or intolerance as shown below.
This is a related entry (typeCode=MFST) that indicates the manifestations (typeCode=MFST) the reported allergy or intolerance. These are events that may occur, or have occurred in the past as a reaction to the allergy or intolerance.

The entry contained with this entry relationship is some sort of problem that is a manifestation of the allergy. It is recorded using the Problem Entry structure, with the additional template identifier (2.16.840.1.113883.10.20.1.54) indicating that this problem is a reaction.

An optional <entryRelationship> element may be present indicating the severity of the problem. When present, this <entryRelationship> element shall contain a severity observation conforming to the Severity entry template (1.3.6.1.4.1.19376.1.5.3.1.4.1). For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element. The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘true’.

An optional <entryRelationship> may be present indicating the clinical status of the allergy, e.g., resolved, in remission, active. When present, this <entryRelationship> element shall contain a clinical status observation conforming to the Problem Status Observation template (1.3.6.1.4.1.19376.1.5.3.1.4.1.1). The typeCode SHALL be ‘REFR’ and inversionInd SHALL be ‘false’. For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element.

One or more optional <entryRelationship> elements may be present providing an additional comments (annotations) for the allergy. When present, this
<entryRelationship> element shall contain an entry conforming to the Comment entry template (1.3.6.1.4.1.19376.1.5.3.1.4.2). The typeCode SHALL be ‘SUBJ’ and inversionInd SHALL be ‘true’.

For CDA this SHALL be represented with the <entryRelationship> element. For HL7 Version 3 Messages, this SHALL be represented as a <sourceOf> element.

6.3.4.16 Medications 1.3.6.1.4.1.19376.1.5.3.1.4.7

This content module describes the general structure for a medication. All medication administration acts will be derived from this content module.

6.3.4.16.1 Standards

<table>
<thead>
<tr>
<th>Pharmacy</th>
<th>HL7 Pharmacy Domain (Normative)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCD</td>
<td>ASTM/HL7 Continuity of Care Document</td>
</tr>
</tbody>
</table>
This section makes use of the linking, severity and instruction entries.

Medications are perhaps the most difficult data elements to model due to variations in the ways that medications are prescribed.

This profile identifies the following relevant fields of a medication as being important to be able to generate in a medical summary. The table below identifies and describes these fields, and indicates the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

### 6.3.4.16.2.1 Medication Fields

<table>
<thead>
<tr>
<th>Field</th>
<th>Opt</th>
<th>CDA Tag</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start and Stop Date</td>
<td>R2</td>
<td>&lt;effectiveTime&gt;</td>
<td>The date (and time if available) when the medication regimen began and is expected to finish. The first component of the &lt;effectiveTime&gt;</td>
</tr>
</tbody>
</table>
encodes the lower and upper bounds over which the `<substanceAdministration>` occurs, and the start time is determined from the lower bound. If the medication has been known to be stopped, the high value must be present, but expressed as a flavor of null (e.g., Unknown).

| Frequency | R2 | `<effectiveTime>` | The frequency indicates how often the medication is to be administered. It is often expressed as the number of times per day, but which may also include information such as 1 hour before/after meals, or in the morning, or evening. The second `<effectiveTime>` element encodes the frequency. In cases where split or tapered doses are used, these may be found in subordinate `<substanceAdministration>` elements. |
| Route | R2 | `<routeCode>` | The route is a coded value, and indicates how the medication is received by the patient (by mouth, intravenously, topically, et cetera). |
| Dose | R2 | `<doseQuantity>` | The amount of the medication given. This should be in some known and measurable unit, such as grams, milligrams, et cetera. It may be measured in "administration" units (such as tablets or each), for medications where the strength is relevant. In this case, only the unit count is specified, no units are specified. It may be a range. |
| Site | O | `<approachSiteCode>` | The site where the medication is administered, usually used with IV or topical drugs. |
| Rate | R2 | `<rateQuantity>` | The rate is a measurement of how fast the dose is given to the patient over time (e.g., .5 liter / 1 hr), and is often used with IV drugs. |
| Product | R¹ | `<consumable>` `<name>` `<consumable>` | The name of the substance or product. This should be sufficient for a provider to identify the kind of medication. It may be a trade name or a generic name. This information is required in all medication entries. If the name of the medication is unknown, the type, purpose or other description may be supplied. The name should not include packaging, strength or dosing information. Note: Due to restrictions of the CDA schema, there is no way to explicitly link the name to the narrative text. |
| Strength | R2 | `<consumable>` `<code>` `<originalText>` `<code>` `<consumable>` | The name and strength of the medication. This information is only relevant for some medications, as the dose of the medication is often sufficient to indicate how much medication the patient receives. For example, the medication Percocet comes in a variety of strengths, which indicate specific amounts of two different medications being received in single tablet. Another example is eye-drops, where the medication is in a solution of a particular strength, and the dose quantity is some number of drops. The `originalText` referenced by the `<code>` element in the consumable should refer to the name and strength of the medication in the narrative text. Note: Due to restrictions of the CDA schema, there is no way to separately record the strength. |
| Code | R2 | `<consumable>` `<code>` `<consumable>` | A code describing the product from a controlled vocabulary, such as RxNorm, First DataBank, et cetera. |
| Instructions | R2 | `<entryRelationship>` | A place to put free text comments to support additional relevant information, or to deal with specialized dosing instructions. For example, "take with food", or tapered dosing. |
| Indication | O | `<entryRelationship>` | A link to supporting clinical information about the reason for providing the medication (e.g., a link to the relevant diagnosis). |

¹ A consumable is not necessary when the substanceAdministration code indicates none or unknown medications.
The general model is to record each prescribed medication in a \(<\text{substanceAdministration}>\) intent (moodCode='INT'). Medications that have been reported by the patient or administered (instead of prescribed), are recorded in the same element, except that this is now an event (moodCode='EVN'). The \(<\text{substanceAdministration}>\) element may contain subordinate \(<\text{substanceAdministration}>\) elements in a related component entry to deal with special cases (see the section below on Special Cases). These cases include split, tapered, or conditional dosing, or combination medications. The use of subordinate \(<\text{substanceAdministration}>\) elements to deal with these cases is optional. The comment field should always be used in these cases to provide the same information as free text in the top level \(<\text{substanceAdministration}>\) element. There are a variety of special cases for dosing that need to be accounted for. These are described below. Most of these special cases involve changing the dosage or frequency over time, or based on some measurement. When the dosage changes, then additional entries are required for each differing dosage. The last case deals with combination medications.

6.3.4.16.3.1 Normal Dosing 1.3.6.1.4.1.19376.1.5.3.1.4.7.1

This template identifier is used to identify medication administration events that do not require any special processing. The parent template is 1.3.6.1.4.1.19376.1.5.3.1.4.7. Medications that use this template identifier shall not use subordinate \(<\text{substanceAdministration}>\) acts.

6.3.4.16.3.2 Tapered Doses 1.3.6.1.4.1.19376.1.5.3.1.4.8

This template identifier is used to identify medication administration events that require special processing to handle tapered dosing. The parent template is 1.3.6.1.4.1.19376.1.5.3.1.4.7. A tapered dose is often used for certain medications where abrupt termination of the medication can have negative consequences. Tapered dosages may be done by adjusting the dose frequency, the dose amount, or both.

When merely the dose frequency is adjusted, (e.g., Prednisone 5mg b.i.d. for three days, then 5mg. daily for three days, and then 5mg every other day), then only one medication entry is needed, multiple frequency specifications recorded in \(<\text{effectiveTime}>\) elements. When the dose varies (e.g. Prednisone 15mg daily for three days, then 10 mg daily for three days, the 5 mg daily for three days), subordinate medication entries should be created for each distinct dosage.

6.3.4.16.3.3 Split Dosing 1.3.6.1.4.1.19376.1.5.3.1.4.9

This template identifier is used to identify medication administration events that require special processing to handle split dosing. The parent template is 1.3.6.1.4.1.19376.1.5.3.1.4.7. A split dose is often used when different dosages are given at different times (e.g., at different times of day, or on different days). This may be to account for different metabolism rates at different times of day, or to simply address drug
packaging deficiencies (e.g., and order for Coumadin 2mg on even days, 2.5mg on odd
days is used because Coumadin does not come in a 2.25mg dose form).
In this case a subordinate <substanceAdministration> entry is required for each separate
dosage.

6.3.4.16.3.4 Conditional Dosing 1.3.6.1.4.1.19376.1.5.3.1.4.10
This template identifier is used to identify medication administration events that require
special processing to handle conditional dosing. The parent template is
1.3.6.1.4.1.19376.1.5.3.1.4.7. A conditional dose is often used when the dose amount
differs based on some measurement (e.g., an insulin sliding scale dose based on blood
sugar level). In this case a subordinate <substanceAdministration> entry is required for
each different dose, and the condition should be recorded.

6.3.4.16.3.5 Combination Medications 1.3.6.1.4.1.19376.1.5.3.1.4.11
This template identifier is used to identify medication administration events that require
special processing to handle combination medications. The parent template is
1.3.6.1.4.1.19376.1.5.3.1.4.7. A combination medication is made up of two or more other
medications. These may be prepackaged, such as Percocet, which is a combination of
Acetaminophen and oxycodone in predefined ratios, or prepared by a pharmacist, such as
a GI cocktail.
In the case of the prepackaged combination, it is sufficient to supply the name of the
combination drug product, and its strength designation in a single
<substanceAdministation> entry. The dosing information should then be recorded as
simply a count of administration units.
In the latter case of a prepared mixture, the description of the mixture should be provided
as the product name (e.g., "GI Cocktail"), in the <substanceAdministration> entry. That
entry may, but is not required, to have subordinate <substanceAdministration> entries
included beneath it to record the components of the mixture.

6.3.4.16.4 <templateId root='2.16.840.1.113883.10.20.1.24'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7'/>
All medications entries use the <templateId> elements specified above to indicate that
they are medication acts. This element is required. In addition, a medication entry shall
further identify itself using one of the template identifiers detailed in the next section.

6.3.4.16.5 <templateId root='' />
The <templateId> element identifies this <entry> as a particular type of medication event,
allowing for validation of the content. As a side effect, readers of the CDA can quickly
locate and identify medication events. The templateId must use one of the values in the
table below for the root attribute.

<table>
<thead>
<tr>
<th>root</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.3.6.1.4.1.19376.1.5.3.1.4.7.1</td>
<td>A &quot;normal&quot; &lt;substanceAdministration&gt; act that may not contain any subordinate &lt;substanceAdministration&gt; acts.</td>
</tr>
</tbody>
</table>

2009-08-10 149 Copyright © 2009: IHE International
1.3.6.1.4.1.19376.1.5.3.1.4.8 A `<substanceAdministration>` act that records tapered dose information in subordinate `<substanceAdministration>` act.

1.3.6.1.4.1.19376.1.5.3.1.4.9 A `<substanceAdministration>` act that records split dose information in subordinate `<substanceAdministration>` acts.

1.3.6.1.4.1.19376.1.5.3.1.4.10 A `<substanceAdministration>` act that records conditional dose information in subordinate `<substanceAdministration>` acts.

1.3.6.1.4.1.19376.1.5.3.1.4.11 A `<substanceAdministration>` act that records combination medication component information in subordinate `<substanceAdministration>` acts.

6.3.4.16.6 `<id root=' ' extension=' '/>`

A top level `<substanceAdministration>` element must be uniquely identified. If there is no explicit identifier for this observation in the source EMR system, a GUID may be used for the root attribute, and the extension may be omitted. At least one identifier must be present, more than one may appear. Subordinate `<substanceAdministration>` elements may, but need not be uniquely identified.

6.3.4.16.7 `<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '>`

The `<code>` element is used to supply a code that describes the `<substanceAdministration>` act, not the medication being administered or prescribed. This may be a procedure code, such as those found in CPT-4 (and often used for billing), or may describe the method of medication administration, such as by intravenous injection. The type of medication is coded in the consumable, do not supply the code for the medication in this element. This element is optional.

One of the following values from SNOMED CT shall be used in the `<code>` element to record that a patient is either not on medications, or that medications are not known.
### Entry Type | Code  | Display Name                              | Description                                                                 |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Medication</td>
<td>182904002</td>
<td>Drug Treatment Unknown</td>
<td>To indicate lack of knowledge about drug therapy</td>
</tr>
<tr>
<td>Medication</td>
<td>182849000</td>
<td>No Drug Therapy Prescribed</td>
<td>To indicate the absence of any prescribed medications</td>
</tr>
<tr>
<td>Medication</td>
<td>408350003</td>
<td>Patient Not On Self-Medications</td>
<td>To indicate no treatment</td>
</tr>
</tbody>
</table>

#### 6.3.4.16.8 `<reference value='' />'</reference>`

4720 The URI given in the value attribute of the `<reference>` element points to an element in the narrative content that contains the complete text describing the medication. In a CDA document, the URI given in the value attribute of the `<reference>` element points to an element in the narrative content that contains the complete text describing the medication. In an HL7 message, the content of the text element shall contain the complete text describing the medication.

#### 6.3.4.16.9 `<statusCode code='completed' />`

4725 The status of all `<substanceAdministration>` elements must be "completed". The act has either occurred, or the request or order has been placed.

#### 6.3.4.16.10 `<effectiveTime xsi:type='IVL_TS'>`

4730 The first `<effectiveTime>` element encodes the start and stop time of the medication regimen. This an interval of time (xsi:type='IVL_TS'), and must be specified as shown. This is an additional constraint placed upon CDA Release 2.0 by this profile, and simplifies the exchange of start/stop and frequency information between EMR systems.

#### 6.3.4.16.11 `<low value='' />' <high value='' />'`

4735 The `<low>` and `<high>` values of the first `<effectiveTime>` element represent the start and stop times for the medication. The `<low>` value represents the start time, and the `<high>` value represents the stop time. If either the `<low>` or the `<high>` value is unknown, this shall be recorded by setting the nullFlavor attribute to UNK. The `<high>` value records the end of the medication regimen according to the information provided in the prescription or order. For example, if the prescription is for enough medication to last 30 days, then the high value should contain a date that is 30 days later then the `<low>` value. The rationale is that a provider, seeing an un-refilled prescription would normally assume that the medication is no longer being taken, even if the intent of the treatment plan is to continue the medication indefinitely.

4740

#### 6.3.4.16.12 `<effectiveTime operator='A' xsi:type='TS|PIVL_TS|EIVL_TS|PIVL_PPD_TS|SXPR_TS' />`

4745 The second `<effectiveTime>` element records the frequency of administration. This `<effectiveTime>` element must be intersected with the previous time specification...
(operator='A'), producing the bounded set containing only those time specifications that fall within the start and stop time of the medication regimen. Several common frequency expressions appear in the table below, along with their XML representations.

### 6.3.4.16.12.1 Specifying Medication Frequency

<table>
<thead>
<tr>
<th>Freq</th>
<th>Description</th>
<th>XML Representation</th>
</tr>
</thead>
<tbody>
<tr>
<td>b.i.d.</td>
<td>Twice a day</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_TS' institutionSpecified='true' operator='A'&gt; &lt;period value='12' unit='h'/&gt;&lt;/effectiveTime&gt;</code></td>
</tr>
<tr>
<td>q12h</td>
<td>Every 12 hours</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_TS' institutionSpecified='false' operator='A'&gt; &lt;period value='12' unit='h'/&gt;&lt;/effectiveTime&gt;</code></td>
</tr>
<tr>
<td>Once</td>
<td>Once, on 2005-09-01 at 1:18am.</td>
<td><code>&lt;effectiveTime xsi:type='TS' value='200509010118'/&gt;</code></td>
</tr>
<tr>
<td>t.i.d.</td>
<td>Three times a day, at times determined by the person administering the medication</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_TS' institutionSpecified='true' operator='A'&gt; &lt;period value='8' unit='h'/&gt;&lt;/effectiveTime&gt;</code></td>
</tr>
<tr>
<td>q6h</td>
<td>Every 8 hours</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_TS' institutionSpecified='false' operator='A'&gt; &lt;period value='8' unit='h'/&gt;&lt;/effectiveTime&gt;</code></td>
</tr>
<tr>
<td>qam</td>
<td>In the morning</td>
<td><code>&lt;effectiveTime xsi:type='EIVL' operator='A'&gt; &lt;event code='ACM'/&gt;&lt;/effectiveTime&gt;</code></td>
</tr>
<tr>
<td>Every day at 8 in the morning for 10 minutes</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_TS' operator='A'&gt; &lt;phase&gt; &lt;low value=&quot;198701010800&quot; inclusive=&quot;true&quot;/&gt; &lt;width value=&quot;10&quot; unit=&quot;min&quot;/&gt; &lt;/phase&gt; &lt;period value='1' unit='d'/&gt;&lt;/effectiveTime&gt;</code></td>
<td></td>
</tr>
<tr>
<td>q4-6h</td>
<td>Every 4 to 6 hours.</td>
<td><code>&lt;effectiveTime xsi:type='PIVL_PPD TS' institutionSpecified='false' operator='A'&gt; &lt;period value='5' unit='h'/ &gt; &lt;standardDeviation value='1' unit='h'/&gt; &lt;/effectiveTime&gt;</code></td>
</tr>
</tbody>
</table>

The last frequency specification is about as bad as it gets, but can still be represented accurately within the HL7 V3 datatypes. The mean (average) of the low and high values is specified for the period. The mean of 4 and 6 is 5. The standard deviation is recorded as one half the difference between the high and low values, with an unspecified distribution. The type attribute of the `<effectiveTime>` element describes the kind of frequency specification it contains. More detail is given for each type in the table below.

### 6.3.4.16.12.2 Data types used in Frequency Specifications

<table>
<thead>
<tr>
<th>xsi:type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>TS</td>
<td>An xsi:type of TS represents a single point in time, and is the simplest of all to represent. The value attribute of the <code>&lt;effectiveTime&gt;</code> element specifies the point in time in HL7 date-time format (CCYYMMDDHHMMSS)</td>
</tr>
<tr>
<td>PIVL_TS</td>
<td>An xsi:type of PIVL TS is the most commonly used, representing a periodic interval of time. The <code>&lt;low&gt;</code> element of <code>&lt;phase&gt;</code> may be present. If so it specifies the starting point, and only the lower order components of this value are relevant with respect to the <code>&lt;period&gt;</code>. The <code>&lt;width&gt;</code> element represents the duration of the dose administration (e.g., for IV administration). The <code>&lt;period&gt;</code> indicates how often the dose is given. Legal values for the unit attribute of <code>&lt;period&gt;</code> are s, min, h, d, wk and mo representing seconds, minutes, hours, days, weeks, and months respectively.</td>
</tr>
<tr>
<td>EIVL_TS</td>
<td>An xsi:type of EIVL TS represents an event based time interval, where the event is not a precise time, but is often used for timing purposes (e.g. with meals, between meals, before breakfast, before sleep). Refer to the HL7 TimingEvent vocabulary for the codes to use for the <code>&lt;event&gt;</code> element. This interval may specify an <code>&lt;offset&gt;</code> which provides information about the time offset from the specified event (e.g., <code>&lt;offset&gt;&lt;low value='-1' unit='h'/&gt;&lt;width value='10' unit='min'/&gt; &lt;/offset&gt;</code></td>
</tr>
</tbody>
</table>
means 1 hour before the event. In that same example, the `<width>` element indicates the duration for
the dose to be given.

| PIVL_PPD_TS | An xsi:type of PIVL_PPD_TS represents a probabilistic time interval and is used to represent
dosing frequencies like q4-6h. This profile requires that the distributionType of this interval be left
unspecifed. The `<period>` element specifies the average of the time interval, and the value of the
`<standardDeviation>` shall be computed as half the width of the interval. The unit attributes of the
`<period>` and `<standardDeviation>` elements shall be the same. |
| SXPR_TS     | An xsi:type of SXPR_TS represents a parenthetical set of time expressions. This type is used when
the frequency varies over time (e.g., for some cases of tapered dosing, or to handle split dosing). The
`<comp>` elements of this `<effectiveTime>` element are themselves time expressions (using any of
the types listed above). Each `<comp>` element may specify an operator (e.g. to intersect or form the
union of two sets). |

6.3.4.16.13 `<routeCode code=' ' displayName=' ' codeSystem='2.16.840.1.113883.5.112'
  codeSystemName='RouteOfAdministration'>`

The `<routeCode>` element specifies the route of administration using the HL7
RouteOfAdministration vocabulary. A code must be specified if the route is known, and
the displayName attribute should be specified. If the route is unknown, this element shall
not be sent.

6.3.4.16.14 `<approachSiteCode code=' ' codeSystem=' '>`

The `<approachSiteCode>` element describes the site of medication administration. It may
be coded to a controlled vocabulary that lists such sites (e.g., SNOMED-CT). In CDA
documents, this element contains a URI in the value attribute of the `<reference>` that
points to the text in the narrative identifying the site. In a message, the `<originalText>`
element shall contain the text identifying the site.

6.3.4.16.15 `<doseQuantity> `<low value=' ' unit=' '/>` `<high value=' ' unit=' '/>`</doseQuantity>

The dose is specified if the `<doseQuantity>` element. If a dose range is given (e.g., 1-2
tablets, or 325-750mg), then the `<low>` and `<high>` bounds are specified in their
respective elements, otherwise both `<low>` and `<high>` have the same value. If the dose is
in countable units (tablets, caplets, "eaches"), then the unit attribute is not sent. Otherwise
the units are sent. The unit attribute should be derived from the HL7
UnitsOfMeasureCaseSensitive vocabulary.

6.3.4.16.16 `<low|high value=' '> `<translation> `<originalText><reference value=' '/></originalText>`
  `<translation>`</low|high >

Any `<low>` and `<high>` elements used for `<doseQuantity>` or `<rateQuantity>` should
contain a `<translation>` element that provides a `<reference>` to the `<originalText>` found
in the narrative body of the document. In a CDA document, any `<low>` and `<high>`
elements used for `<doseQuantity>` or `<rateQuantity>` should contain a `<translation>`
element that provides a `<reference>` to the `<originalText>` found in the narrative body of
the document. In a message, the <originalText> may contain the original text used to describe dose quantity.

6.3.4.16.17 <rateQuantity><low value=' ' unit=' '/><high value=' '
unit=' '/></rateQuantity>

The rate is specified in the <rateQuantity> element. The rate is given in units that have measure over time. In this case, the units should be specified as a string made up of a unit of measure (see doseQuantity above), followed by a slash (/), followed by a time unit (s, min, h or d).

Again, if a range is given, then the <low> and <high> elements contain the lower and upper bound of the range, otherwise, they contain the same value.

6.3.4.16.18 <consumable>

The <consumable> element shall be present, and shall contain a <manufacturedProduct> entry conforming to the Product Entry template

6.3.4.16.19 <entryRelationship typeCode='REFR'>

The top level <substanceAdministration> element may contain a reference (typeCode='REFR') to related prescription activity as described in the Supply Entry.

6.3.4.16.20 <entryRelationship typeCode='COMP'>

A top level <substanceAdministration> element may contain one or more related components, either to handle split, tapered or conditional dosing, or to support combination medications.

In the first three cases, the subordinate components shall specify only the changed <frequency> and/or <doseAmount> elements. For conditional dosing, each subordinate component shall have a <precondition> element that specifies the <observation> that must be obtained before administration of the dose. The value of the <sequenceNumber> shall be an ordinal number, starting at 1 for the first component, and increasing by 1 for each subsequent component. Components shall be sent in <sequenceNumber> order.

6.3.4.16.21 <entryRelationship typeCode='SUBJ' inversionInd='true'/>

At most one instruction may be provided for each <substanceAdministration> entry. If provided, it shall conform to the requirements listed for Patient Medication Instructions. The instructions shall contain any special case dosing instructions (e.g., split, tapered, or conditional dosing), and may contain other information (take with food, et cetera).

6.3.4.16.22 <entryRelationship typeCode='RSON'>

<act classCode='ACT' moodCode='EVN'>

<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.1.4.1'/>

{id root=' ' extension=' '/>
A `<substanceAdministration>` event may indicate one or more reasons for the use of the medication. These reasons identify the concern that was the reason for use via the `Internal Reference` entry content module.

The extension and root of each observation present must match the identifier of a concern entry contained elsewhere within the CDA document.

A consumer of the Medical Summary is encouraged, but not required to maintain these links on import.

6.3.4.16.23 `<precondition><criterion>`
   `<text><reference value=' '></text>`
   `<criterion></precondition>`

In a CDA document, the preconditions for use of the medication are recorded in the `<precondition>` element. The value attribute of the `<reference>` element is a URL that points to the CDA narrative describing those preconditions.

6.3.4.16.24 `<condition typeCode='PRCN'>`
   `<criterion>
   `<text></text>
   `<value nullFlavor='UNK'/>`
   `<interpretationCode nullFlavor='UNK'/>`
   `<criterion>
   </condition>`

In a message, the preconditions for use of the medication are recorded in the `<condition>` element. The typeCode shall be PRCN. The `<text>` element of the criterion shall contain a text description of the precondition. The `<value>` element is required, and may be recorded in a structured data type if known, and if not, may be recorded using a nullFlavor as shown above. The same is true for `<interpretationCode>`.

6.3.4.17 Immunizations 1.3.6.1.4.1.19376.1.5.3.1.4.12

An immunizations entry is used to record the patient's immunization history.
6.3.4.17.1 Specification

```xml
<substanceAdministration typeCode='SBADM' moodCode='EVN' negationInd='true{{!}}false'>
    <!-- The reasonCode would normally provide a reason why the immunization was not performed. It isn't supported by CDA R2, and so comments will have to suffice. -->
    <reasonCode code='' codeSystem='' codeSystemName='ActNoImmunizationReasonIndicator'/>
    <reasonCode code='' codeSystem='' codeSystemName='ActNoImmunizationReasonIndicator'/><id root='' extension=''/>
    <code code='IMMUNIZ' codeSystem='2.16.840.1.113883.5.4' codeSystemName='ActCode'/>
    <text><reference value='#xxx'/></text>
    <statusCode code='completed'/>
    <templateId root='2.16.840.1.113883.10.20.1.24'/>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.12'/>
    <effectiveTime value=''/>
    <routeCode code='' codeSystem='' codeSystemName='RouteOfAdministration'/>
    <approachSiteCode code='' codeSystem='' codeSystemName='HumanSubstanceAdministrationSite'/>
    <doseQuantity value='' units=''/>
    <consumable typeCode='CSM'>
        <manufacturedProduct classCode='MANU'>
            <manufacturedLabeledDrug classCode='MMAT' determinerCode='KIND'>
                <code code='' codeSystem='' codeSystemName=''>
                    <originalText><reference value='#yyy'/></originalText>
            </code>
        </manufacturedLabeledDrug>
    </manufacturedProduct>
</consumable>
</entryRelationship>
</entryRelationship>
</entryRelationship>
</entryRelationship>
</substanceAdministration>
```

6.3.4.17.2 <substanceAdministration typeCode='SBADM' moodCode='EVN' negationInd='true{{!}}false'>

An immunization is a substance administration event. An immunization entry may be a record of why a specific immunization was not performed. In this case, negationInd shall be set to "true", otherwise, it shall be false.
6.3.4.17.3 <templateId root='2.16.840.1.113883.10.20.1.24'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.12'/>

The <templateId> elements identifies this <substanceAdministration> as an immunization. Both elements shall be present as shown above.

6.3.4.17.4 <id root=' ' extension=' '/>

This shall be the identifier for the immunization event.

6.3.4.17.5 <code code='IMMUNIZ' codeSystem='2.16.840.1.113883.5.4' codeSystemName='ActCode'/>

This required element records that the act was an immunization. The substance administration act must have a <code> element with code and codeSystem attributes present. If no coding system is used by the source, then simply record the code exactly as shown above. Another coding system that may be used for codes for immunizations are the CPT-4 codes for immunization procedures. This <code> element shall not be used to record the type of vaccine used from a vocabulary of drug names.

<table>
<thead>
<tr>
<th>codeSystem</th>
<th>codeSystemName</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.16.840.1.113883.5.4</td>
<td>IMMUNIZ</td>
<td>The IMMUNIZ term from the HL7 ActCode vocabulary.</td>
</tr>
</tbody>
</table>

6.3.4.17.6 <text><reference value='#xxx'/></text>

In a CDA document, the URI given in the value attribute of the <reference> element points to an element in the narrative content that contains the complete text describing the immunization activity. In an HL7 message, the content of the text element shall contain the complete text describing the immunization activity.

6.3.4.17.7 <statusCode code='completed'/>

The statusCode shall be set to "completed" for all immunizations.

6.3.4.17.8 <effectiveTime value=' '/>

The effectiveTime element shall be present and should contain a time value that indicates the date of the substance administration. If the date is unknown, this shall be recorded using the nullFlavor attribute, with the reason that the information is unknown being specified. Otherwise, the date shall be recorded, and should have precision of at least the day.

6.3.4.17.9 <routeCode code=' ' codeSystem=' ' codeSystemName='RouteOfAdministration'/>

See routeCode under Medications.
6.3.4.17.10  <approachSiteCode code=' ' codeSystem=' ' codeSystemName='HumanSubstanceAdministrationSite'/>
See approachSiteCode under Medications.

6.3.4.17.11  <doseQuantity value=' ' units=' '/>
See doseQuantity under Medications.

6.3.4.17.12  <consumable typeCode='CSM'>
See consumable under Medications.

6.3.4.17.13  <entryRelationship typeCode='REFR'>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3'/>
The top level <substanceAdministration> element may contain a reference (typeCode='REFR') to related Supply entry

6.3.4.17.14  <entryRelationship typeCode='SUBJ'>
<observation classCode='OBS' moodCode='EVN'>
<templateId root='2.16.840.1.113883.10.20.1.46'/>
This optional entry relationship may be present to indicate that position of this immunization in a series of immunizations.

6.3.4.17.15  <code code='30973-2' displayName='Dose Number' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
The <code> element shall be present and must be recorded with the code and codeSystem attributes shown above. This element indicates that the observation describes the dose number for the immunization.

6.3.4.17.16  <statusCode code='completed'/> The <statusCode> element shall be present, and must be recorded exactly as shown above. This element indicates that the observation has been completed.

6.3.4.17.17  <value xsi:type='INT' value=' '/> The <value> element shall be present, and shall indicate the immunization series number in the value attribute.

6.3.4.17.18  <entryRelationship inversionInd='false' typeCode='CAUS'> This repeatable element should be used to identify adverse reactions caused by the immunization.

6.3.4.17.19  <observation classCode='OBS' moodCode='EVN'> This element is required, and provides a pointer to the adverse reaction caused by the immunization.
6.3.4.17.20  <templateId root='2.16.840.1.113883.10.20.1.28'/>
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5'/>
            <templateId root='2.16.840.1.113883.10.20.1.54'/>

It shall contain a conforming Problem Entry that also conform to the CCD Reaction template.

6.3.4.17.21  <id root=' ' extension=' '/>

This element is required, and gives the identifier of the adverse reaction. The adverse reaction pointed to by this element shall be described in more detail using the Allergies entry, elsewhere in the document where this element was found.

6.3.4.17.22  <!-- Optional <entryRelationship> element containing comments -->

An immunization entry can have negationInd set to true to indicate that an immunization did not occur. In this case, it shall have at least one comment that provides an explanation for why the immunization did not take place. Other comments may also be present.

6.3.4.18 Supply Entry 1.3.6.1.4.1.19376.1.5.3.1.4.7.3

The supply entry describes a prescription activity.
6.3.4.18.1 Specification

```xml
<substanceAdministration classCode='SBADM' moodCode='INT|EVN'>
  ...
  <entryRelationship typeCode='REFR' inversionInd='false'>
    <sequenceNumber value=''/>
    <supply classCode='SPLY' moodCode='INT|EVN'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3'/>
      <id root='' extension=''/>
      <repeatNumber value=''/>
      <quantity value='' unit=''/>
      <author>
        <time value=''/>
        <assignedAuthor>
          <id root='' extension=''/>
          <addr/>
          <telecom use='' value=''/>
          <assignedPerson><name></name></assignedPerson>
          <representedOrganization><name></name></representedOrganization>
        </assignedAuthor>
      </author>
      <performer typeCode='PRF'>
        <time value=''/>
        <assignedEntity>
          <id root='' extension=''/>
          <addr/>
          <telecom use='' value=''/>
          <assignedPerson><name></name></assignedPerson>
          <representedOrganization><name></name></representedOrganization>
        </assignedEntity>
      </performer>
      <!-- Optional Fulfillment instructions -->
    </supply>
  </entryRelationship>
</substanceAdministration>
```

6.3.4.18.2 <entryRelationship typeCode='REFR' inversionInd='false'>

A `<substanceAdministration>` act may reference (typeCode='REFR') a prescription activity in an `<entryRelationship>` element in a CDA document. In a message, the relationship is recorded using a `<sourceOf>` element instead of the `<entryRelationship>` element. The typeCode and inversionInd attributes, and the semantics remain identical.

6.3.4.18.3 <sequenceNumber value='' />

The prescription activity may have a `<sequenceNumber>` element to indicate the fill number. A value of 1, 2 or N indicates that it is the first, second, or Nth fill respectively of a specific prescription. This element should be present when the embedded `<supply>` element has a moodCode attribute of EVN.

6.3.4.18.4 <supply classCode='SPLY' moodCode='INT|EVN'>

The `<supply>` element shall be present. The moodCode attribute shall be INT to reflect that a medication has been prescribed, or EVN to indicate that the prescription has been filled.
6.3.4.18.5 <templateId root='2.16.840.1.113883.10.20.1.34'/>
   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3'/>

The <templateId> elements shown above shall be present, and identify this supply act as a Supply Entry.

6.3.4.18.6 <id root=' ' extension=' '/>

Each supply act shall have an identifier to uniquely identify the supply entry.

6.3.4.18.7 <repeatNumber value=' '/>

Each supply entry should have a <repeatNumber> element that indicates the number of times the prescription can be filled.

6.3.4.18.8 <quantity value=' ' unit=' '/>

The supply entry should indicate the quantity supplied. The value attribute shall be present and indicates the quantity of medication supplied. If the medication is supplied in dosing units (tablets or capsules), then the unit attribute need not be present (and should be set to 1 if present). Otherwise, the unit element shall be present to indicate the quantity (e.g., volume or mass) of medication supplied.

6.3.4.18.9 <author>

A supply entry that describes an intent (<supply classCode='SPLY' moodCode='INT'>) may include an <author> element to identify the prescribing provider.

6.3.4.18.10 <time value=' '/>

The <time> element must be present to indicate when the author created the prescription. If this information is unknown, it shall be recorded by setting the nullFlavor attribute to UNK.

6.3.4.18.11 <assignedAuthor>

The <assignedAuthor> element shall be present, and identifies the author.

6.3.4.18.12 <id root=' ' extension=' '/>

One or more <id> elements should be present. These identifiers identify the author of the act. When the author is the prescribing physician they may include local identifiers or regional identifiers necessary for prescribing.

6.3.4.18.13 <assignedPerson><name/></assignedPerson>
   <representedOrganization><name/></representedOrganization>

An <assignedPerson> and/or <representedOrganization> element shall be present. This element shall contain a <name> element to identify the prescriber or their organization.
6.3.4.18.14  <performer typeCode='PRF'>
The <performer> element may be present to indicate who is intended (moodCode='INT'),
or actually filled (moodCode='EVN') the prescription.

6.3.4.18.15  <time value=' '/>
The <time> element shall be present to indicate when the prescription was filled
(moodCode='EVN'). If this information is unknown, it shall be recorded by setting the
nullFlavor attribute to UNK.
The <time> element should be present to indicate when the prescription is intended to be
filled (moodCode='INT').

6.3.4.18.16  <assignedEntity>
The <assignedEntity> element shall be present, and identifies the filler of the
prescription.

6.3.4.18.17  <id root=' ' extension=' '/>
One or more <id> elements should be present. These identify the performer.

6.3.4.18.18  <assignedPerson><name/></assignedPerson>
<representedOrganization><name/></representedOrganization>
An <assignedPerson> and/or <representedOrganization> element shall be present. This
element shall contain a <name> element to identify the filler or their organization.

6.3.4.18.19  <!-- Optional Fulfillment instructions -->
<entryRelationship typeCode='SUBJ'>
</entryRelationship>
An entry relationship may be present to provide the fulfillment instructions. When
present, this entry relationship shall contain a Medication Fulfillment Instructions entry.

6.3.4.19 Product Entry 1.3.6.1.4.1.19376.1.5.3.1.4.7.2
The product entry describes a medication or immunization used in a
<substanceAdministration> or <supply> act. It adopts the constraints of the ASTM/HL7
Continuity of Care Document.
6.3.4.19.1 Specification

```xml
<!-- Within a CDA Document -->
<manufacturedProduct>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.2'/>
  <templateId root='2.16.840.1.113883.10.20.1.53'/>
  <manufacturedMaterial>
    <code code='' displayName='' codeSystem='' codeSystemName=''>
      <originalText><reference value=''/></originalText>
    </code>
    <name></name>
  </manufacturedMaterial>
</manufacturedProduct>

<!-- Within a message -->
<administerableMaterial>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.2'/>
  <templateId root='2.16.840.1.113883.10.20.1.53'/>
  <administerableMaterial>
    <code></code>
    <desc></desc>
  </administerableMaterial>
</administerableMaterial>
```

In a CDA document, the name and strength of the medication are specified in the elements under the `<manufacturedMaterial>` element. In a message, they are contained within the `<administerableMaterial>` element, inside another `<administerableMaterial>` element\(^1\). The templateId elements are required and identify this as a product entry.

\(^1\) This duplication of element names is an artifact of the standard.

6.3.4.19.2 `<manufacturedProduct> -OR- `<administerableMaterial>`

```xml
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.2'/>
<templateId root='2.16.840.1.113883.10.20.1.53'/>
<manufacturedMaterial> -OR- `<administerableMaterial>`
```

In a CDA document, the name and strength of the medication are specified in the elements under the `<manufacturedMaterial>` element. In a message, they are contained within the `<administerableMaterial>` element, inside another `<administerableMaterial>` element\(^1\). The templateId elements are required and identify this as a product entry.

6.3.4.19.3 `<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '>`

```xml
<originalText><reference value=''/></originalText>
</code>
```

The `<code>` element of the `<manufacturedMaterial>` describes the medication. This may be coded using a controlled vocabulary, such as RxNorm, First Databank, or other vocabulary system for medications, and should be the code that represents the generic medication name and strength (e.g., acetaminophen and oxycodone -5/325), or just the generic medication name alone if strength is not relevant (Acetaminophen).

In a CDA document, the `<originalText>` shall contain a `<reference>` whose URI value points to the generic name and strength of the medication, or just the generic name alone if strength is not relevant. Inside a message, the `<originalText>` may contain the actual text that describes the medication in similar fashion.

**Note:** When the text is supplied from the narrative, the implication is that if you supply the components of a combination medication in an entry, you must also display these in the narrative text, otherwise you would not be able to break the combination medication down into its component parts. This is entirely consistent with the CDA Release 2.0 requirements that the narrative supply the necessary and relevant human readable information content.
The `<code>` element is also used to support coding of the medication. If coded, it must provide a code and codeSystem attribute using a controlled vocabulary for medications. The displayName for the code and codeSystemName should be provided as well for diagnostic and human readability purposes, but are not required. The table below provides the codeSystem and codeSystemName for several controlled terminologies that may be used to encode medications and/or immunizations.

<table>
<thead>
<tr>
<th>codeSystem</th>
<th>codeSystemName</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.16.840.1.113883.6.88</td>
<td>RxNorm</td>
<td>RxNorm</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.69</td>
<td>NDC</td>
<td>National Drug Codes</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.63</td>
<td>FDDC</td>
<td>First DataBank Drug Codes</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.96</td>
<td>SNOMED-CT</td>
<td>SNOMED Controlled Terminology</td>
</tr>
<tr>
<td>2.16.840.1.113883.6.59</td>
<td>CVX</td>
<td>CDC Vaccine Codes</td>
</tr>
</tbody>
</table>

The code used for an immunization may use code systems other than what might be used for other medications, such as the CDC maintained CVX codes. Code systems that describe vaccination procedures (such as CPT-4) shall not be used to describe the vaccine entry.

**6.3.4.19.4<name> -OR- <desc>**

In a CDA document, the `<name>` element should contain the brand name of the medication (or active ingredient in the case of subordinate `<substanceAdministration>` elements used to record components of a medication). Within a message, this information shall be provided in the `<desc>` element.

**6.3.4.20 Simple Observations 1.3.6.1.4.1.19376.1.5.3.1.4.13**

The simple observation entry is meant to be an abstract representation of many of the observations used in this specification. It can be made concrete by the specification of a few additional constraints, namely the vocabulary used for codes, and the value representation. A simple observation may also inherit constraints from other specifications (e.g., ASTM/HL7 Continuity of Care Document).
6.3.4.20.1 Specification

These acts are simply observations that have occurred, and so are recorded using the <observation> element as shown above.

6.3.4.20.2 <observation classCode='OBS' moodCode='EVN'>

These acts are simply observations that have occurred, and so are recorded using the <observation> element as shown above.

6.3.4.20.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'/>

The <templateId> element identifies this <observation> as a simple observation, allowing for validation of the content. The templateId must appear as shown above.

6.3.4.20.4 <id root='' extension='' />

Each observation shall have an identifier.

6.3.4.20.5 <code code='' displayName='' codeSystem='' codeSystemName='' />

Observations shall have a code describing what was measured. The code system used is determined by the vocabulary constraints on the types of measurements that might be recorded in a section. Content modules that are derived from the Simple Observation content module may restrict the code system and code values used for the observation.

6.3.4.20.6 <text><reference value='#xxx'/></text> -OR- <text>text</text>

Each observation measurement entry may contain a <text> element providing the free text that provides the same information as the observation within the narrative portion of the document with a <text> element. For CDA based uses of Simple Observations, this
element SHALL be present, and SHALL contain a <reference> element that points to the related string in the narrative portion of the document. For HL7 Version 3 based uses, the <text> element MAY be included.

6.3.4.20.7 <statusCode code='completed'/>
The status code of all observations shall be completed.

6.3.4.20.8 <effectiveTime value=' '/>
The <effectiveTime> element shall be present in standalone observations, and shall record the date and time when the measurement was taken. This element should be precise to the day. If the date and time is unknown, this element should record that using the nullFlavor attribute.

6.3.4.20.9 <value xsi:type=' '/>
The value of the observation shall be recording using a data type appropriate to the observation. Content modules derived from the Simple Observation content module may restrict the allowable data types used for the observation.

6.3.4.20.10 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/>
If there is an interpretation that can be performed using an observation result (e.g., high, borderline, normal, low), these may be recorded within the interpretationCode element.

6.3.4.20.11 <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>
The methodCode element may be used to record the specific method used to make an observation when this information is not already pre-coordinated with the observation code.

6.3.4.20.12 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>
The targetSiteCode may be used to record the target site where an observation is made when this information is not already pre-coordinated with the observation code.

6.3.4.20.13 <author><assignedAuthor classCode='ASSIGNED'>...<assignedAuthor></author>
In CDA uses, SimpleObservations are assumed to be authored by the same author as the document through context conduction. However specific authorship of observation may be represented by listing the author in the header and referencing the author in a <author> relationship. If authors are explicitly listed in documents, an <id> element SHOULD reference the ID of the author in the header through an assignedAuthor Role. If the author of the observation is not an author of the document the <person> object including a name and ID SHALL be included.

For HL7 Version 3 purposes, the <author> element SHOULD be present unless it can be determined by conduction from organizers or higher level structures. When used for HL7
Version 3 the role element name is <assignedEntity> and the author is represented a <assignedPerson> element.

6.3.4.21 Vital Signs Organizer 1.3.6.1.4.1.19376.1.5.3.1.4.13.1

A vital signs organizer collects vital signs observations.

6.3.4.21.1 Specification

```xml
<organizer classCode='CLUSTER' moodCode='EVN'>
  <templateId root='2.16.840.1.113883.10.20.1.32'/>
  <templateId root='2.16.840.1.113883.10.20.1.35'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.1'/>
  <id root='' extension=''/>
  <code code='46680005' displayName='Vital signs'
    codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'/>
  <statusCode code='completed'/>
  <effectiveTime value=''/>
</organizer>
```

6.3.4.21.2 <organizer classCode='CLUSTER' moodCode='EVN'>

The vital signs organizer is a cluster of vital signs observations.

6.3.4.21.3 <templateId root='2.16.840.1.113883.10.20.1.32'/>

The vital signs organizer shall have the <templateId> elements shown above to indicate that it inherits constraints from the ASTM/HL7 CCD Specification for Vital signs, and the constraints of this specification.

6.3.4.21.4 <id root=' ' extension=' '/>

The organizer shall have an <id> element.

6.3.4.21.5 <code code='46680005' displayName='Vital signs'
  codeSystem='2.16.840.1.113883.6.96'
  codeSystemName='SNOMED CT'/>
6.3.4.21.6 `<statusCode code='completed'/>`

The observations have all been completed.

6.3.4.21.7 `<effectiveTime value=' '/>`

The effective time element shall be present to indicate when the measurement was taken.

6.3.4.21.8 `<author typeCode='AUT'><assignedEntity1 typeCode='ASSIGNED'>...</assignedEntity1></author>`

For use with HL7 Version 3, Vital Sign organizers SHALL contain an `<author>` element to represent the person or device.

6.3.4.21.9 `<-- one or more vital signs observations --> <component typeCode='COMP'>`

The organizer shall have one or more `<component>` elements that are `<observation>` elements using the Vital Signs Observation template.

6.3.4.22 Vital Signs Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.2

A vital signs observation is a simple observation that uses a specific vocabulary, and inherits constraints from CCD.

6.3.4.22.1 Specification

```xml
<observation classCode='OBS' moodCode='EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'/>
  <templateId root='2.16.840.1.113883.10.20.1.31'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.2'/>
  <id root=' ' extension=' '/>
  <code code=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'/>
  <text><reference value='#xxx'/></text>
  <statusCode code='completed'/>
  <effectiveTime value=' '/>
  <repeatNumber value=' '/>
  <value xsi:type='PQ' value=' ' unit=' '/>
  <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/>
  <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>
  <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>
</observation>
```

6.3.4.22.2 `<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'/>`

`<templateId root='2.16.840.1.113883.10.20.1.31'/>`

`<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.2'/>`

A vital signs observation shall have the `<templateId>` elements shown above to indicate that it inherits constraints from the ASTM/HL7 CCD Specification for Vital signs, and the constraints of this specification.
A vital signs observation entry shall use one of the following LOINC codes, with the specified data types and units.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>9279-1</td>
<td>RESPIRATION RATE</td>
<td>/min</td>
</tr>
<tr>
<td>8867-4</td>
<td>HEART BEAT</td>
<td></td>
</tr>
<tr>
<td>2710-2</td>
<td>OXYGEN SATURATION</td>
<td>%</td>
</tr>
<tr>
<td>8480-6</td>
<td>INTRAVASCULAR SYSTOLIC</td>
<td>mm[Hg]</td>
</tr>
<tr>
<td>8462-4</td>
<td>INTRAVASCULAR DIASTOLIC</td>
<td></td>
</tr>
<tr>
<td>8310-5</td>
<td>BODY TEMPERATURE</td>
<td>Cel or [degF]</td>
</tr>
<tr>
<td>8302-2</td>
<td>BODY HEIGHT (MEASURED)</td>
<td>m, cm,[in_us]</td>
</tr>
<tr>
<td>8306-3</td>
<td>BODY HEIGHT^LYING</td>
<td>[in_uk]</td>
</tr>
<tr>
<td>8287-5</td>
<td>CIRCUMFRENCE.OCCIPITAL-FRONTAL (TAPE MEASURE)</td>
<td></td>
</tr>
<tr>
<td>3141-9</td>
<td>BODY WEIGHT (MEASURED)</td>
<td>kg, g, [lb_av]</td>
</tr>
</tbody>
</table>

The <value> element shall be present, and shall be of the appropriate data type specified for measure in the table above.

The interpretation code may be present to provide an interpretation of the vital signs measure (e.g., High, Normal, Low, et cetera).

The <methodCode> element may be present to indicate the method used to obtain the measure. Note that method used is distinct from, but possibly related to the target site.

The target site of the measure may be identified in the <targetSiteCode> element (e.g., Left arm [blood pressure], oral [temperature], et cetera).

The family history organizer collects the problems of a patient's family member.
6.3.4.23.1 Specification

Each family history entry is organized (classCode='CLUSTER') into a group of observations about a family member.

6.3.4.23.2 <organizer classCode='CLUSTER' moodCode='EVN'>

The organizer is identified by the <templateId> elements, which shall be present as shown above.

6.3.4.23.3 <templateId root='2.16.840.1.113883.10.20.1.23'/>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.15'/>

The organizer is identified by the <templateId> elements, which shall be present as shown above.

6.3.4.23.4 <subject typeCode='SUBJ'>

The <subject> element shall be present and relates the subject of the observations to the patient. It shall contain a <relatedSubject> element that is a personal relation of the patient (classCode='PRS').

6.3.4.23.5 <code code='' displayName='' codeSystem='2.16.840.1.113883.5.111' codeSystemName='RoleCode'/>

The <code> element shall be present, and give the relationship of the subject to the patient. The code attribute shall be present, and shall contain a value from the HL7
6.3.4.23.6 <subject>
The <subject> element contains information about the relation.

6.3.4.23.7 <sdtc:id root=' ' extension=' '/>
The <sdtc:id> element should be present. It is used to identify the patient relation to create a pedigree graph.

6.3.4.23.8 <administrativeGenderCode code=' '/>
The <administrativeGenderCode> element should be present. It gives the gender of the relation.

6.3.4.23.9 <participant typeCode='IND'>
   <participantRole classCode='PRS'>
The <participant> element may be present to record the relationship of the subject to other family members to create a pedigree graph. It shall contain a <participantRole> element showing the relationship of the subject to other family members (classCode='PRS').

6.3.4.23.10 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>
The <code> element shall be present, and gives the relationship of the participant to the subject. The code attribute shall be present, and shall contain a value from the HL7 FamilyMember vocabulary. The codeSystem attribute shall be present and shall use the value shown above.

6.3.4.23.11 <playingEntity classCode='PSN'>
The <playingEntity> element identifies the related person. It shall be recorded as shown above.

6.3.4.23.12 <sdtc:id root=' ' extension=' '/>
The <sdtc:id> element shall be present. It must have the same root and extension attributes of the <subject> of a separate family history organizer. See Appendix C of PCC-TF for definition of this extension to CDA.

6.3.4.23.13 <component typeCode='COMP'>
   <observation classCode='OBS' moodCode='EVN'>
   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.3'/>
The family history organizer shall contain one or more components using the <component> element shown above. These components must conform the Family History Observation template.
6.3.4.24 Social History Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.4

A social history observation is a simple observation that uses a specific vocabulary, and inherits constraints from CCD.

6.3.4.24.1 Standards

| CCD | ASTM/HL7 Continuity of Care Document |

6.3.4.24.2 Parent Template

The parent of this template is Simple Observation. This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.33

6.3.4.24.3 Specification

Figure 6.3-43 Social History Observation Example

```xml
<observation typeCode='OBS' moodCode='EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'/>
  <templateId root='2.16.840.1.113883.10.20.1.33'/>
  <id root=' ' extension=' '/>
  <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>
  <text><reference value='#xxx'/></text>
  <statusCode code='completed'/>
  <effectiveTime value=' '/>
  <repeatNumber value=' '/>
  <value xsi:type=' '/>
  <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/>
  <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>
  <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>
</observation>
```

6.3.4.24.4 <templateId root='2.16.840.1.113883.10.20.1.33'/>

These <templateId> elements identify this as a Social History observation.

6.3.4.24.5 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>

The <code> element identifies the type social history observation.
### 6.3.4.24.6 `<repeatNumber>` value=""/>

The `<repeatNumber>` element should not be used in a social history observation.

### 6.3.4.24.7 `<value xsi:type=' ' ... />

The `<value>` element reports the value associated with the social history observation. The data type to use for each observation should be drawn from the table above.

Observations in the table above using the PQ data type have a unit in the form `{xxx}/d, {xxx}/wk or {xxx}/a represent the number of items per day, week or year respectively.

The value attribute indicates the number of times of the act performed, and the units represent the frequency. The example below shows how to represent 1 drink per day.

```xml
<code code='160573003' displayName='ETOH Use'
  codeSystem='2.16.840.1.113883.6.96'
  codeSystemName='SNOMED CT'/>

<value xsi:type='PQ' value='1' unit='{drink}/d'/>
```

Observations in the table using the CD data type should include coded values from an appropriate vocabulary to represent the social history item. The example below shows the encoding to indicate drug use of cannabis.

```xml
<code code='363908000' displayName='Drug Use'
  codeSystem='2.16.840.1.113883.6.96'
  codeSystemName='SNOMED CT'/>

<value xsi:type='CD' code='398705004' displayName='cannabis'
  codeSystem='2.16.840.1.113883.6.96'
  codeSystemName='SNOMED CT'/>
```

Other social history observations may use any appropriate data type.
The <interpretationCode>, <methodCode>, and <targetSiteCode> elements should not be used in a social history observation.

6.4 HL7 Version 2.0 Content Modules For care management

6.5 PCC Value Sets

6.5.1 APR H&P History of Past Illness Value Set

6.5.2 APR H&P Social History Value Set

Examples Using PCC Content Profiles

Example documents conforming to each profile can be found on the IHE wiki at the following URLs.

<table>
<thead>
<tr>
<th>Profile and Content</th>
<th>URL</th>
</tr>
</thead>
<tbody>
<tr>
<td>XDS-MS</td>
<td></td>
</tr>
<tr>
<td>Referral Summary</td>
<td>XDSMS Example1</td>
</tr>
<tr>
<td>Discharge Summary</td>
<td>XDSMS Example1</td>
</tr>
<tr>
<td>XPHR</td>
<td></td>
</tr>
<tr>
<td>XPHR Content</td>
<td>XPHR Example1</td>
</tr>
<tr>
<td>XPHR Update</td>
<td>XPHR Example2</td>
</tr>
<tr>
<td>(EDR) ED Referral</td>
<td>EDR Example</td>
</tr>
<tr>
<td>(APS) Antepartum Summary</td>
<td>APS Example</td>
</tr>
<tr>
<td>(EDES)</td>
<td></td>
</tr>
<tr>
<td>Triage Note</td>
<td>EDES Example1</td>
</tr>
<tr>
<td>ED Nursing Note</td>
<td>EDES Example2</td>
</tr>
<tr>
<td>Composite Triage and Nursing Note</td>
<td>EDES Example3</td>
</tr>
<tr>
<td>ED Physician Note</td>
<td>EDES Example4</td>
</tr>
<tr>
<td>(FSA) Functional Status Section</td>
<td>FSA Example</td>
</tr>
</tbody>
</table>

Validating CDA Documents using the Framework
Many of the constraints specified by the content modules defined in the PCC Technical Framework can be validated automatically by software. Automated validation is a very desirable capability, as it makes it easier for implementers to test the correctness of their implementations. With regard to validation of the content module, the PCC Technical Framework narrative is the authoritative specification, not any automated software tool. Having said that, it is still very easy to create a validation framework for the IHE PCC Technical Framework using a XML validation tool such as Schematron. Since each content module has a name (the template identifier), any XML instance that reports itself to be of that "class" can be validated by creating assertions that must be true for each constraint indicated for the content module. In the XML representation, the <templateId> element is a child of the element that is claiming conformance to the template named. Thus the general pattern of a Schematron that validates a specific template is shown below:

```
<schema xmlns="http://www.ascc.net/xml/schematron" xmlns:cda="urn:hl7-org:v3">
  <ns prefix="cda" uri="urn:hl7-org:v3" />
  <pattern name='ReferralSummary'>
    <rule context='*[cda:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.1.3"]'>
      <!-- one or more assertions made by the content module -->
    </rule>
  </pattern>
</schema>
```

### A.1 Validating Documents

For document content modules, the pattern can be extended to support common document content module constraints as shown below:
<schema xmlns="http://www.ascc.net/xml/schematron" xmlns:cda="urn:hl7-org:v3">
  <ns prefix="cda" uri="urn:hl7-org:v3"/>
  <pattern name='ReferralSummary'>
    <rule context='*[templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.1.3"]'>
      <!-- Verify that the template id is used on the appropriate type of object -->
      <assert test='..//ClinicalDocument'>
        Error: The referral content module can only be used on Clinical Documents.
      </assert>
      <!-- Verify that the parent templateId is also present. -->
      <assert test='templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.1.2"]'>
        Error: The parent template identifier for medical summary is not present.
      </assert>
      <!-- Verify the document type code -->
      <assert test='@code = "34133-9"'>
        Error: The document type code of a referral summary must be 34133-9 SUMMARIZATION OF EPISODE NOTE.
      </assert>
      <assert test='@codeSystem = "2.16.840.1.113883.6.1"'>
        Error: The document type code must come from the LOINC code system (2.16.840.1.113883.6.1).
      </assert>
      <!-- Verify that all required data elements are present -->
      <assert test='//templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.1"]'>
        Error: A referral summary must contain a reason for referral.
      </assert>
      <!-- Alert on any missing required if known elements -->
      <assert test='//templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.8"]'>
        Warning: A referral summary should contain a list of history of past illnesses.
      </assert>
      <!-- Note any missing optional elements -->
      <assert test='//templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.18"]'>
        Note: This referral summary does not contain the pertinent review of systems.
      </assert>
    </rule>
  </pattern>
</schema>
A.2 Validating Sections

The same pattern can be also applied to sections with just a few minor alterations.

```
<schema xmlns="http://www.ascc.net/xml/schematron" xmlns:cda="urn:hl7-org:v3">
  <ns prefix="cda" uri="urn:hl7-org:v3" />
  <pattern name='ReasonForReferralUncoded'>
    <rule context='*[templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.3.1"]'>
      <!-- Verify that the template id is used on the appropriate type of object -->
      <assert test='section'>
        Error: The coded reason for referral module can only be used on a section.
      </assert>
      <assert test='false'>
        Manual: Manually verify that this section contains narrative providing the reason for referral.
      </assert>
      <!-- Verify that the parent templateId is also present. -->
      <assert test='templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.3.1"]'>
        Error: The parent template identifier for the reason for referral module is not present.
      </assert>
      <!-- Verify the section type code -->
      <assert test='code[@code = "42349-1"]'>
        Error: The section type code of the reason for referral section must be 42349-1 REASON FOR REFERRAL.
      </assert>
      <assert test='code[@codeSystem = "2.16.840.1.113883.6.1"]'>
        Error: The section type code must come from the LOINC code system (2.16.840.1.113883.6.1).
      </assert>
    </rule>
  </pattern>
  <pattern name='ReasonForReferralCoded'>
    <rule context='*[templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.3.2"]'>
      <!-- The parent template will have already verified the type of object -->
      <!-- Verify that the parent templateId is also present. -->
      <assert test='templateId[@root="1.3.6.1.4.1.19376.1.5.3.1.3.1"]'>
        Error: The parent template identifier for the reason for referral module is not present.
      </assert>
      <!-- Don't bother with the section type code, as the parent template caught it -->
      <!-- Verify that all required data elements are present -->
      <assert test='*[templateId[@root = "1.3.6.1.4.1.19376.1.5.3.1.4.13"]]'>
        Error: A coded reason for referral section must contain an simple observation.
      </assert>
      <!-- Alert on any missing required if known elements -->
      <!-- Note any missing optional elements -->
    </rule>
  </pattern>
</schema>
```

A similar pattern can also be followed for Entry and Header content modules, and these are left as an exercise for the reader.
A.3 Phases of Validation and Types of Errors

Note that each message in the Schematrons shown above start with a simple text string that indicates whether the message indicates one of the following conditions:

- An error, e.g., the failure to transmit a required element,
- A warning, e.g., the failure to transmit a required if known element,
- A note, e.g., the failure to transmit an optional element.
- A manual test, e.g., a reminder to manually verify some piece of content.

Schematron supports the capability to group sets of rules into phases by the pattern name, and to specify which phases of validation should be run during processing. To take advantage of this capability, one simply breaks each <pattern> element above up into separate patterns depending upon whether the assertion indicates an error, warning, note or manual test, and then associate each pattern with a different phase. This is shown in the figure below.

Using these simple "templates" for template validation one can simply create a collection of Schematron patterns that can be used to validate the content modules in the PCC.
Technical Framework. Such Schematrons are expected to be made available as part of the MESA test tools that are provided to IHE Connectathon participants, and which will also be made available to the general public after connectathon.

Extensions to CDA Release 2.0

This section describes extensions to CDA Release 2.0 that are used by the IHE Patient Care Coordination Technical Framework.

A.4 IHE PCC Extensions

All Extensions to CDA Release 2.0 created by the IHE PCC Technical Committee are in the namespace urn:ihe:pcc:hl7v3.

The approach used to create extension elements created for the PCC Technical Framework is the same as was used for the HL7 Care Record Summary (see Appendix E) and the ASTM/HL7 Continuity of Care Document (see section 7.2).

A.4.1 replacementOf

The <replacementOf> extension element is applied to a section appearing in a PHR Update Document to indicate that that section's content should replace that of a previously existing section. The identifier of the previously existing section is given so that the PHR Manager receiving the Update content will know which section to replace. The model for this extension is shown below.

![Figure 6.5-1Model for replacementOf](image)

Use of this extension is shown below. The <replacementOf> element appears after all other elements within the <section> element. The <id> element appearing in the <externalDocumentSection> element shall provide the identifier of the section being replaced in the parent document.
A.5 Extensions Defined Elsewhere used by IHE PCC

A.5.1 Entity Identifiers

There is often a need to record an identifier for an entity so that it can be subsequently referenced. This extension provides a mechanism to store that identifier. The element appears after any <realm>, <typeId> or <templateId> elements, but before all others in the entity where it is used:

```
<playingEntity classCode='ENT' determinerCode='INSTANCE'>
  <sdtc:id root='1.3.6.4.1.4.1.2835.2' extension='EntityID'/>
</playingEntity>
```

A.5.2 Patient Identifier

There is a need to record the identifier by which a patient is known to another healthcare provider. This extension provides a role link between the assigned, related or associated entity, and the patient role.

Use of this extension to record the identifier under which the patient is known to a provider is shown below.

```
<assignedEntity>
  <id extension='1' root='1.3.6.4.1.4.1.2835.1'/>
  <code code='59058001'
    codeSystem='2.16.840.1.113883.6.96'
    codeSystemName='SNOMED CT'
    displayName='General Physician'/
  >
  <addr>
    <streetAddressLine>21 North Ave</streetAddressLine>
    <city>Burlington</city>
    <state>MA</state>
    <postalCode>01803</postalCode>
    <country>USA</country>
  </addr>
  <telecom value='tel:(999)555-1212' use='WP'/>
  <assignedPerson>
    <name>
      <prefix>Dr.</prefix><given>Bernard</given><family>Wiseman</family><suffix>Sr.</suffix>
    </name>
  </assignedPerson>
  <sdtc:patient xmlns:sdtc='urn:hl7-org:sdtc' >
    <sdtc:id root='1.3.6.4.1.4.1.2835.2' extension='PatientMRN'/>
  </sdtc:patient>
</assignedEntity>
```

The <patient> element records the link between the related, assigned or associated entity and the patient. The <id> element provides the identifier for the patient. The root attribute
of the <id> should be the namespace used for patient identifiers by the entity. The
extension attribute of the <id> element shall be the patient's medical record number or
other identifier used by the entity to identify the patient.