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HL7 Implementation Guide for CDA® Release 2: Consolidated CDA Templates for Clinical Notes (US Realm)

Draft Standard for Trial Use Release 2

November 2014

Volume 1 — Introductory Material

Sponsored by: Structured Documents Work Group Patient Care Work Group Child Health work Group

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	resources/solutions-managing-your-practice/coding-billing-
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SNOMED CT	International Healthcare Terminology Standards Developing
	Organization (IHTSDO) http://www.ihtsdo.org/snomed-
	ct/get-snomed-ct or info@ihtsdo.org
Logical Observation Identifiers	Regenstrief Institute
Names & Codes (LOINC)	
International Classification of	World Health Organization (WHO)
Diseases (ICD) codes	

Structure of This Guide

Two volumes comprise this *HL7 Implementation Guide for CDA® Release 2: Consolidated CDA Templates for Clinical Notes.* Volume 1 provides narrative introductory and background material pertinent to this implementation guide, including information on how to understand and use the templates in Volume 2. Volume 2 contains the normative Clinical Document Architecture (CDA) templates for this guide along with lists of all templates, code systems, value sets, and changes from the previous version.

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2 INTRODUCTION

2.1 Purpose

This two-volume implementation guide (IG) contains an overview of Clinical Document Architecture (CDA) markup standards, design, and use (Volume 1) and a consolidated library of CDA templates for clinical notes applicable to the US Realm (Volume 2). These two volumes comprise a Draft Standard for Trial Use (DSTU).

The consolidated library incorporates previous efforts from Health Level Seven (HL7), Integrating the Healthcare Enterprise (IHE), the Health Information Technology Standards Panel (HITSP), the HL7 Health Story guides, HITSP C32, and related components of IHE Patient Care Coordination (IHE PCC). It has additional or enhanced document types for greater expressivity and decrease ambiguity (see Volume 2 Summary of Changes). Volume 1 adds new general guidance (see <a href=Volume 1 Summary of Changes). Volume 2 has a highly detailed change log in Chapter 9 "Changes From Previous Version".

This guide, in conjunction with the HL7 CDA Release 2 (CDA R2) standard, is to be used for implementing the following CDA documents and header constraints for clinical notes.

- Care Plan including Home Health Plan of Care (HHPoC)
- Consultation Note
- Continuity of Care Document (CCD)
- Diagnostic Imaging Reports (DIR)
- Discharge Summary
- History and Physical (H&P)
- Operative Note
- Procedure Note
- Progress Note
- Referral Note
- Transfer Summary
- Unstructured Document
- Patient Generated Document (US Realm Header)

2.2 Audience

The audience for this implementation guide includes architects and developers of healthcare information technology (HIT) systems in the US Realm that exchange patient clinical data. Business analysts and policy managers can also benefit from a basic understanding of the use of CDA templates across multiple implementation use cases.

2.3 Organization of the Guide

This implementation guide is organized into two volumes. Volume 1 contains primarily narrative text describing the Consolidated CDA Release 2 (C-CDA R2) guide, whereas Volume 2 contains normative CDA template definitions.

2.3.1 Volume 1 Introductory Material

This document, Volume 1, provides an overview of Clinical Document Architecture (CDA), summaries of recent changes to the standard, and information on how to understand and use the CDA templates provided in Volume 2.

- **Chapter 1**—Introduction
- **Chapter 2**—CDA R2 Background. This section contains selected background material on the CDA R2 base standard, to aid the reader in conceptualizing the "templated CDA" approach to implementation guide development.
- **Chapter 3**—Design Considerations. This section includes design considerations that describe overarching principles that have been developed and applied across the CDA templates in this guide. Material in this section can be thought of as "heuristics", as opposed to the formal and testable constraints found in Volume 2 of this guide.
- **Chapter 4**—Using This Implementation Guide. This section describes the rules and formalisms used to constrain the CDA R2 standard. It describes the formal representation of CDA templates, the mechanism by which templates are bound to vocabulary, and additional information necessary to understand and correctly implement the normative content found in Volume 2 of this guide.
- **Appendices**. The Appendices include a high-level change log, a summary of extensions to CDA R2, an excerpt of the Claims Attachments Implementation Guide covering Mime Multipart/Related Messages, and additional information.

2.3.2 Volume 2 CDA Templates and Supporting Material

Volume 2 includes CDA templates and prescribes their use for a set of specific document types. The main chapters are:

- **Chapter 1**—Document-Level Templates. This chapter defines the US Realm Header template for that applies across all of the consolidated document types. It defines each of the document types and header constraints specific to each as well as the section-level templates (required and optional) for each.
- **Chapter 2**—Section-Level Templates. This chapter defines the section templates referenced within the document types. Sections are atomic units, and can be reused by future specifications.
- **Chapter 3**—Entry-Level Templates. This chapter defines entry-level templates, called clinical statements. Machine processable data are sent in the entry templates. The entry templates are referenced by one or more section templates. Entry-level templates are always contained in section-level templates, and section-level templates are always contained in a document. Entries are atomic units, and can be reused by future specifications.

- **Chapter 4**—Participation and Other Templates. This chapter defines templates for CDA participants (e.g., author, performer) and other fielded items (e.g., address, name) that cannot stand on their own without being nested in another template.
- **Chapter 5**—Template Ids in This Guide
- Chapter 6—Value Sets in This Guide
- **Chapter 7**—Code Systems in This Guide
- **Chapter 8**—Retired Templates
- **Chapter 9**—Changes from Previous Version. This chapter provides a detailed change log.

2.4 Contents of the Package

The following files comprise this implementation guide package:

Table 1: Contents of the Package

Filename	Description	Standards Applicability
CDAR2_IG_CCDA_CLINNOTES_R2_D1_2014NOV _V1_Introductory_Material.docx	Implementation Guide Introductory Material	Normative
CDAR2_IG_CCDA_CLINNOTES_R2_D1_2014NOV _V2_ Templates_and_Supporting_Material.docx	Implementation Guide Template Library and Supporting Material	Normative
C-CDA_R2_Care_Plan.xml	Care Plan Sample	Informative
C-CDA_R2_CCD.xml C-CDA_R2_CCD_2.xml	Continuity of Care Document sample files	Informative
C-CDA_R2_Consultation_Note.xml	Consultation Note Sample	Informative
C-CDA_R2_Diagnostic_Imaging_Report.xml	Imaging Integration and DICOM Diagnostic Imaging Reports (DIR) Sample	Informative
C-CDA_R2_Discharge_Summary.xml	Discharge Summary Sample	Informative
C-CDA_R2_HHCP485.xml	Home Health Plan of Care Sample based on form 485	informative
C-CDA_R2_History_and_Physical.xml	History and Physical (H&P) Sample	Informative
C-CDA_R2_Operative_Note.xml	Operative Note Sample	Informative
C-CDA_R2_PatientGeneratedDocument.xml	Patient Generated Document Header Sample	Informative

C-CDA_R2_Procedure_Note.xml	Procedure Note Sample	Informative
C-CDA_R2_Progress_Note.xml	Progress Note Sample	Informative
C-CDA_R2_Referral_Note.xml	Referral Note Sample	Informative
C-CDA_R2_Transfer_Summary.xml	Transfer Summary Sample	Informative
C-CDA_R2_Unstructured_Document_embed.xml C-CDA_R2_Unstructured_Document_reference.xml C-CDA_R2_UD_sample.docx C-CDA_R2_UD_sample.pdf	Unstructured Document Samples	Informative
C-CDA-R2.sch voc.xml	C-CDA R2 Schematron file and supporting vocabulary file	Informative
CDA.xsl	Stylesheet for rendering	Informative
schema	Folder containing updated CDA XML Schemas	Informative

3 CDA R2 BACKGROUND

CDA R2 is "... a document markup standard that specifies the structure and semantics of 'clinical documents' for the purpose of exchange" [CDA R2, Section 1.1]¹. Clinical documents, according to CDA, have the following characteristics:

- Persistence
- Stewardship
- Potential for authentication
- Context
- Wholeness
- Human readability

CDA defines a header for classification and management and a document body that carries the clinical record. While the header metadata are prescriptive and designed for consistency across all instances, the body is highly generic, leaving the designation of semantic requirements to implementation.

3.1 Templated CDA

CDA R2 can be constrained by mechanisms defined in the "Refinement and Localization" section of the *HL7 Version 3 Interoperability Standards*. The mechanism most commonly used to constrain CDA is referred to as "templated CDA". In this approach, a library is created containing modular CDA templates such that the templates can be reused across any number of CDA document types, as shown in the following figure.

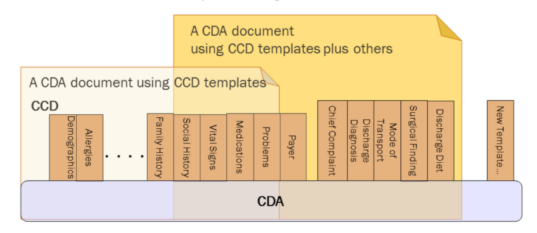


Figure 1: Templated CDA

There are many different kinds of templates that might be created. Among them, the most common are:

¹ HL7 CDA Release 2. http://www.hl7.org/implement/standards/product_brief.cfm?product_id=7

² http://www.hl7.org/v3ballot/html/infrastructure/conformance/conformance.htm

- **Document-level templates:** These templates constrain fields in the CDA header, and define containment relationships to CDA sections. For example, a History and Physical document-level template might require that the patient's name be present, and that the document contain a Physical Exam section.
- **Section-level templates:** These templates constrain fields in the CDA section, and define containment relationships to CDA entries. For example, a Physical Exam section-level template might require that the section/code be fixed to a particular LOINC code, and that the section contain a Systolic Blood Pressure observation.
- **Entry-level templates:** These templates constrain the CDA clinical statement model in accordance with real world observations and acts. For example, a Systolic Blood Pressure entry-level template defines how the CDA Observation class is constrained (how to populate observation/code, how to populate observation/value, etc.) to represent the notion of a systolic blood pressure.
- **Participation and other templates:** These templates group a common set of constraints for reuse in CDA documents. For example, US Realm Date and Time (DTM.US.FIELDED) includes a set of common constraints for recording time. This template is referenced several times throughout the implementation guide in place of repeating constraints.

A CDA implementation guide (such as this one) includes references to those templates that are applicable. On the implementation side, a CDA instance populates the template identifier (templateId) field where it wants to assert conformance to a given template. On the receiving side, the recipient can then not only test the instance for conformance against the CDA Extensible Markup Language (XML) schema, but also test the instance for conformance against asserted templates.

3.2 Current Project

This guide was developed and produced through the joint efforts of HL7, two Sub Work Groups of the Office of the National Coordinator (ONC) Standards and Interoperability (S&I) Framework—Longitudinal Care Plan (LCP) and Long-Term Post-Acute Care (LTPAC) Transition)—and through the SMART C-CDA Collaborative hosted by ONC and Harvard Medical School. This guide builds off of two earlier versions: C-CDA R1 (2011) and C-CDA R1.1 (2012)³.

The ONC Longitudinal Care Coordination Standards and Interoperability (LCC S&I) Work Group and community providers identified a set of priority data elements for shared care and transfer of care for a patient moving from one setting to another. These data elements identified gaps in the existing CDA document types. The current project incorporates these data elements into this implementation guide. Three new document types (Referral Note, Transfer Summary, and Care Plan) and one existing document type (Consultation Note) address the gaps.

The S&I Framework's LCC Long-Term Post-Acute Care (LTPAC) Transition Sub Work Group (SWG) defined the data elements and assisted in the design of the CDA templates

http://www.hl7.org/documentcenter/private/standards/cda/CDAR2_IG_IHE_CONSOL_DSTU_R1dot1_2_012JUL.zip

³ HL7 CDA2 IHE Health Story Consolidation.

to properly express the concepts in the CDA representation of the Referral Note, Transfer Summary, and Consultation Note. The group's related work products can be found at the LCC Long-Term Post-Acute Care (LTPAC) Transition SWG website.4

The S&I Framework's LCC Longitudinal Care Plan (LCP) SWG defined the data elements and assisted in the design of the CDA templates to properly express the concepts in the CDA representation of the Care Plan. The group's related work can be found at the LCC Longitudinal Care Plan (LCP) SWG website.⁵ In addition, the LCP SWG worked with the HL7 Patient Care Work Group (PCWG) during development of the HL7 Care Plan Domain Analysis Model (DAM). The HL7 PCWG's related work can be found at the HL7 Care Plan DAM website.⁶ The IHE PCC Work Group products were reviewed and many participants in the Work Group also participated in the S&I design sessions. The key work reviewed can be found in the IHE Patient Care Plan Content Profile.7 All of these works and group discussions created the design of the CDA Care Plan, which is a static reflection of a dynamic care plan at a point in time.

Diverse document types meet varied types of patient care and coordination needs. The Transfer Summary document is exchanged by healthcare providers in instances when a patient moves between health care settings and care teams temporarily or permanently (e.g., long term care facility to hospital, hospital to skilled nursing facility or home health agency, or from one Primary Care Physician to a new Primary Care Physician). The Transfer Summary provides comprehensive information regarding the patient's history, current status, and care plan.

The Continuity of Care Document (CCD) is a subset of the Transfer Summary and contains just the most clinically important patient information. It is a snapshot in time and may be generated for a single visit or a set of visits. The CCD can be used as an alternative to the Transfer Summary when minimal information needs to be conveyed, or for reporting updates to clinical registries and centralized data repositories.

In cases when a provider requests consultation from another provider, a Referral Note document is exchanged to communicate the referral request and pertinent patient information. When the consultation is completed, the consulting provider may generate a Consultation Note to inform the requesting clinician of her opinion or advice. A patient with complex needs requires the care of multiple providers in various settings. In this situation, a Care Plan document provides a snapshot in time of current health concerns, goals, interventions and care coordination activities amongst providers, the patient, and the patient's caregivers.

The ONC-sponsored SMART (Substitutable Medical Apps and Reusable Technologies) project8, in an effort to make C-CDA instances more reliably consumable by SMART applications for mobile devices, has analyzed real world C-CDA instances and identified common sources of ambiguity and misinterpretation. Many of the clarifications to C-

Acute+Care+%28LTPAC%29+Transition+SWG).

http://wiki.siframework.org/LCC+Longitudinal+Care+Plan+%28LCP%29+SWG

http://www.ihe.net/uploadedFiles/Documents/PCC/IHE_PCC_Suppl_PtCP.pdf

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⁴ S&I Framework, LCC LTPAC Transition SWG. http://wiki.siframework.org/LCC+Long-Term+Post-

⁵ S&I Framework, LCC LCP SWG.

⁶ HL7, Care Plan. http://wiki.hl7.org/index.php?title=Care_Plan

⁷ IHE, Patient Care Plan Content Profile.

⁸ SMART Platforms. http://smartplatforms.org

CDA R1.1 templates analysis.	s, included here as C-	-CDA R2, are as a	direct result of the	SMART

4 DESIGN CONSIDERATIONS

Design considerations describe overarching principles that have been developed and applied across the CDA templates in this guide. Material in this section can be thought of as "heuristics", as opposed to the formal and testable constraints found in Volume 2 of this guide.

4.1 CDA Participations

A CDA participant (e.g., Author, Informant), per the Reference Information Model (RIM), is "an association between an Act and a Role with an Entity playing that Role. Each Entity (in a Role) involved in an Act in a certain way is linked to the Act by one Participation-instance. The kind of involvement in the Act is specified by the Participation.typeCode."

CDA principles when asserting participations include:

- **Participation persistence:** An object's participations (and participation time stamps) don't change just because that object is reused. For instance, authorship of an object doesn't change just because that object is now included in a summary document.
- **Participation evolution:** Additional participations (and participation time stamps) can be ascribed to an object over its lifetime. (In some cases, an electronic health record (EHR) system will create a new object instead of adding participants to an existing object, such as when an EHR has imported a CCD and the receiving clinician chooses to create a local problem list entry corresponding to a problem in the CCD).
- **Device participation:** Devices do not participate as legally responsible entities, but can participate as authors in some scenarios.

Meaningful Use Stage 29 criterion §170.314(b)(4) Clinical Information Reconciliation requires a system to "simultaneously display (i.e., in a single view) the data from at least two list sources in a manner that allows a user to view the data and their attributes, which must include, at a minimum, the **source** and **last modification date**."

CDA addresses this requirement via the Author Participation and its time stamp. CDA requires that Author and Author time stamp be asserted in the document header. From there, authorship propagates to contained sections and contained entries, unless explicitly overridden. Thus, all entries in CDA implicitly include Author and Author time stamp.

In this version of CDA, we have added a new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119) to better ensure consistent representation. This template should be used to explicitly assert authorship and author time stamps, unless the values propagated from the document header hold true.

⁹ US HHS, HIT. http://www.gpo.gov/fdsys/pkg/FR-2012-09-04/pdf/2012-20982.pdf

4.2 Determining a Clinical Statement's Status

A general recipient requirement is to be able to determine the status of an entry—whether it (a problem, a medication administration, etc.) is active, completed, or in some other state. Often complicating the determination is the interplay between an act's various components (such as statusCode and effectiveTime), and inconsistent modeling between different objects.

This guide uses general rules for formalizing the representation of an object's status. We then show how those rules are applied to example templates (Problem Concern Act, Problem Observation).

Principles of C-CDA's approach to status include:

- Act.statusCode specifies the state of the entry: Per the RIM, the statusCode "reflects the state of the activity. In the case of an Observation, this is the status of the activity of observing, not the status of what is being observed".
- Act.moodCode and Act.statusCode are inter-related: Generally, an Observation in EVN (event) mood is a discrete event (you look, listen, measure; you record what you see; you're done), so generally an Observation in EVN mood will have a statusCode of "completed". An exception is a prolonged period of observation, where potentially you'd have an observation in EVN mood that is "active". For an Observation in RQO (request) mood, the statusCode generally remains "active" until the request is complete, at which time the statusCode changes to "completed". For an Observation in GOL (goal) mood, the statusCode generally remains "active" as long as the observation in question is still an active goal for the patient.
- Act.effectiveTime and Act.statusCode are inter-related: Per the RIM, the effectiveTime, also referred to as the "biologically relevant time", is the time at which the observation holds for the patient. So, whereas the effectiveTime is the biologically relevant time, the statusCode is the state of the activity. For a provider seeing a patient in the clinic today, observing a history of heart attack that occurred five years ago, the status of the observation is completed, and the effectiveTime is five years ago.
- **"Status" observations**: C-CDA 1.1 included an optional Problem Status observation. In this guide, we have deprecated the template.

The Problem Concern Act (V2) (templateId 2.16.840.1.113883.10.20.22.4.3:2014-06-09) reflects an ongoing concern on behalf of the provider that placed the concern on a patient's problem list. So long as the underlying condition is of concern to the provider (i.e., as long as the condition, whether active or resolved, is of ongoing concern and interest to the provider), the statusCode is "active". Only when the underlying condition is no longer of concern is the statusCode set to "completed". The effectiveTime of a Problem Concern Act reflects the time that the underlying condition was felt to be a concern—it may or may not correspond to the effectiveTime of the condition (e.g., even five years later, the clinician may remain concerned about a prior heart attack).

A Problem Concern Act can contain many Problem Observations (templateId 2.16.840.1.113883.10.20.22.4.4:2014-06-09). Each Problem Observation is a discrete observation of a condition and therefore will have a statusCode of "completed".

The statusCode of the Problem Concern Act is the definitive indication of the status of the concern. The effectiveTime of the Problem Observation is the definitive indication of whether or not the underlying condition is resolved. This is shown graphically in the following figure.

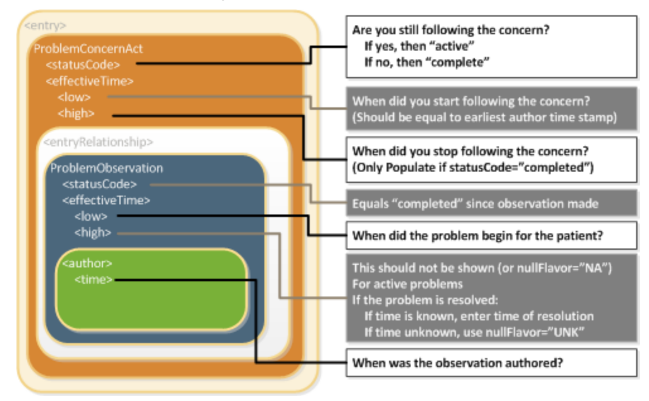


Figure 2: Problem Concern Act

4.3 Rendering Header Information for Human Presentation

Metadata carried in the header may already be available for rendering from EHRs or other sources external to the document. An example of this would be a doctor using an EHR that already contains the patient's name, date of birth, current address, and phone number. When a CDA document is rendered within that EHR, those pieces of information may not need to be displayed since they are already known and displayed within the EHR's user interface.

Good practice recommends that the following be present whenever the document is viewed:

- Document title and document dates
- Service and encounter types, and date ranges as appropriate

- Names of all persons along with their roles, participations, participation date ranges, identifiers, address, and telecommunications information
- Names of selected organizations along with their roles, participations, participation date ranges, identifiers, address, and telecommunications information
- Date of birth for recordTarget(s)
- Patient identifying information

In Operative and Procedure Notes, the following information is typically displayed in the EHR and/or rendered directly in the document:

- The performers of the surgery or procedure, including any assistants
- The surgery or procedure performed (serviceEvent)
- The date of the surgery or procedure

4.4 Unknown and No Known Information

Information technology solutions store and manage data, but sometimes data are not available. An item may be unknown, not relevant, or not computable or measureable, such as where a patient arrives at an emergency department unconscious and with no identification.

In many cases, the C-CDA standard will stipulate that a piece of information is required (e.g., via a SHALL conformance verb). However, in most of these cases, the standard provides an "out", allowing the sender to indicate that the information isn't known.

Here, we provide guidance on representing unknown information. Further details can be found in the HL7 V3 Data Types Release 1 specification that accompanies the CDA R2 normative standard. However, it should be noted that the focus of Consolidated CDA is on the unambiguous representation of known data, and that in general, the often subtle nuances of unknown information representation are less relevant to the recipient.

Many fields in C-CDA contain a "@nullFlavor" attribute, used to indicate an exceptional value. Some flavors of Null are used to indicate that the known information falls outside of value set binding constraints. Not all uses of the @nullFlavor attribute are associated with a case in which information is unknown. Allowable values for populating the attribute give details about the reason the information is unknown, as shown in the following example.

Figure 3: nullFlavor Example

Use null flavors for unknown, required, or optional attributes:

- NI No information. This is the most general and default null flavor.
- NA Not applicable. Known to have no proper value (e.g., last menstrual period for a male).

UNK Unknown. A proper value is applicable, but is not known.

ASKU Asked, but not known. Information was sought, but not found (e.g., the patient was asked but did not know).

NAV Temporarily unavailable. The information is not available, but is expected to be available later.

NASK Not asked. The patient was not asked.

MSK There is information on this item available but it has not been provided by the sender due to security, privacy, or other reasons. There may be an alternate mechanism for gaining access to this information.

OTH The actual value is not an element in the value domain of a variable. (e.g., concept not provided by required code system).

The list above contains those null flavors that are commonly used in clinical documents. For the full list and descriptions, see the nullFlavor vocabulary domain in the CDA R2 normative edition. 10

Any **SHALL**, **SHOULD** or **MAY** conformance statement may use nullFlavor, unless the nullFlavor is explicitly disallowed (e.g., through another conformance statement which includes a **SHALL** conformance for a vocabulary binding to the @code attribute, or through an explicit **SHALL NOT** allow use of nullFlavor conformance).

Figure 4: Attribute Required (nullFlavor not allowed)

```
1. SHALL contain exactly one [1..1] code (CONF:15407).

a. This code SHALL contain exactly one [1..1] @code="11450-4" Problem List (CodeSystem: LOINC 2.16.840.1.113883.6.1) (CONF:15408).

or

2. SHALL contain exactly one [1..1] effectiveTime/@value (CONF:5256).
```

Figure 5: Allowed nullFlavors When Element is Required (with xml examples)

```
1. SHALL contain at least one [1..*] id
2. SHALL contain exactly one [1..1] code
3. SHALL contain exactly one [1..1] effectiveTime
<entrv>
 <observation classCode="OBS" moodCode="EVN">
   <id nullFlavor="NI"/>
   <code nullFlavor="OTH">
     <originalText>New Grading system
   </code>
   <statusCode code="completed"/>
   <effectiveTime nullFlavor="UNK"/>
   <value xsi:type="CD" nullFlavor="OTH">
     <originalText>Spiculated mass grade 5</originalText>
    </value>
  </observation>
</entry>
```

-

¹⁰ HL7 CDA Release 2. http://www.hl7.org/implement/standards/product_brief.cfm?product_id=7

If a sender wants to state that a piece of information is unknown, the following principles apply:

1. If the sender doesn't know an attribute of an act, that attribute can be null.

Figure 6: Unknown Medication Example

2. If the sender doesn't know if an act occurred, the nullFlavor is on the act (detail could include specific allergy, drug, etc.).

Figure 7: Unknown Medication Use of Anticoagulant Drug Example

3. If the sender wants to state "no known", a negationInd can be used on the corresponding act (substanceAdministration, Procedure, etc.)

Previously, CCD, IHE, and HITSP recommended using specific codes to assert no known content, for example 160244002 No known allergies or 160245001 No current problems or disability. Specific codes are still allowed; however, use of these codes is not recommended.

These next examples illustrate nuances of representing information in coded fields when information is a negative assertion, for example it is not the case that the patient has an allergy or it is not the case that a patient takes a medication. The phrases "no known allergies" or "no known medications" are typically associated with this type of negative assertion.

Figure 8: No Known Medications Example

```
<entry>
 <substanceAdministration moodCode="EVN" classCode="SBADM" negationInd="true">
   <text>No known medications</text>
    <consumable>
      <manufacturedProduct>
        <manufacturedLabeledDrug>
          <code code="410942007" displayName="drug or medication"</pre>
                codeSystem="2.16.840.1.113883.6.96"
                codeSystemName="SNOMED CT"/>
         </manufacturedLabeledDrug>
       </manufacturedProduct>
     </consumable>
  </substanceAdministration>
</entry>
```

Figure 9: Value Known, Code for Value Not Known

```
<entry>
  <observation classCode="OBS" moodCode="EVN">
    <value xsi:type="CD" nullFlavor="OTH">
      <originalText>Spiculated mass grade 5</originalText>
    </value>
  </observation>
</entrv>
```

Figure 10: Value Completely Unknown

```
<entry>
  <observation classCode="OBS" moodCode="EVN">
   <value xsi:type="CD" nullFlavor="UNK"/>
  </observation>
</entry>
```

Figure 11: Value Known, Code in Required Code System Not Known But Code from Another Code System is Known

5 USING THIS IMPLEMENTATION GUIDE

This chapter describes the rules and formalisms used to constrain the CDA R2 standard. It describes the formal representation of CDA templates, the mechanism by which templates are bound to vocabulary, and additional information necessary to understand and correctly implement the normative content found in Volume 2 of this guide.

5.1 Levels of Constraint

The CDA standard describes conformance requirements in terms of three general levels corresponding to three different, incremental types of conformance statements:

- Level 1 requirements impose constraints upon the CDA Header. The body of a Level 1 document may be XML or an alternate allowed format. If XML, it must be CDA-conformant markup.
- Level 2 requirements specify constraints at the section level of a CDA XML document: most critically, the section code and the cardinality of the sections themselves, whether optional or required.
- Level 3 requirements specify constraints at the entry level within a section. A specification is considered "Level 3" if it requires any entry-level templates.

Note that these levels are rough indications of what a recipient can expect in terms of machine-processable coding and content reuse. They do not reflect the level or type of clinical content, and many additional levels of reusability could be defined.

The contexts table for each document type lists the sections defined in the document template.

5.2 Conformance Conventions Used in This Guide

5.2.1 Templates and Conformance Statements

Conformance statements within this implementation guide are presented as constraints from Trifolia Workbench, a template repository. An algorithm converts constraints recorded in Trifolia to a printable presentation. Each constraint is uniquely identified by an identifier at or near the end of the constraint (e.g., CONF:86-7345). The digits in the conformance number before the hyphen identify which implementation guide the template belongs to and the number after the hyphen is unique to the owning implementation guide. Together, these two numbers uniquely identify each constraint. These identifiers are persistent but not sequential. Conformance numbers in this guide associated with a conformance statement that is carried forward from a previous version of this guide will carry the same conformance number from the previous version. This is true even if the previous conformance statement has been edited. If a conformance statement is entirely new it will have a new conformance number.

Bracketed information following each template title indicates the template type (section, observation, act, procedure, etc.), the object identifier (OID) or uniform resource

name (URN), and whether the template is open or closed. The identifier OID is the templateId/@root value; all templateIds have an @root value. Versioned templates also have an @extension value, which is a date identifying the version of this template; such templates are identified by URN and the HL7 version (urn:hl7ii). The URN identifier includes both the @root and @extension value for the templateId (for example, identifier urn:hl7ii:2.16.840.1.113883.10.20.5.5.41:2014-06-09).

Each section and entry template in Volume 2 of this guide includes a context table. The "Contained By" column indicates which templates use this template, and if the template is optional or required in the containing template. The "Contains" column indicates any templates that the template uses.

Table 2: Contexts Table Example—Allergy Concern Act (V2)

Contained By:	Contains:
Allergies and Intolerances Section (entries optional) (V2) (optional)	Allergy - Intolerance Observation (V2)
Allergies and Intolerances Section (entries required) (V2) (required)	Author Participation

Each entry template also includes a constraints overview table to summarize the constraints in the template.

Table 3: Constraints Overview Example—Allergy Concern Act (V2)

XPath	Card.	Verb	Data Type	CONF#	Value
act (identifier: urn	:hl7ii:2.16	.840.1.1138	383.10.20	0.22.4.30:2014-	06-09)
@classCode	11	SHALL		1098-7469	2.16.840.1.113883.5.6 (HL7ActClass) = ACT
@moodCode	11	SHALL		1098-7470	2.16.840.1.113883.5.1001 (ActMood) = EVN
templateId	11	SHALL		1098-7471	
@root	11	SHALL		1098-10489	2.16.840.1.113883.10.20.22. 4.30
@extension	11	SHALL		1098-32543	2014-06-09

The expression "such that it" at the end of one conformance statement links that conformance statement to the following subordinate conformance statement to further constrain the first conformance statement. To understand the full effect of this conformance construct, the two conformances must be considered as a single compound requirement. The subordinate conformance statement functions as a subordinate clause (like a "where" clause), which is being applied on the first conformance statement.

The following example shows a compound conformance statement made up of two conformance statements joined by a "such that it" clause. The effect of this syntax can be interpreted as a "where" clause. Thus...

- 1. **SHALL** contain exactly one [1..1] **templateId** (CONF:81-7899) such that it
 - a. **SHALL** contain exactly one [1..1] @root="2.16.840.1.113883.10.20.22.4.31" (CONF:81-10487).

...is understood as:

This template **SHALL** contain exactly one [1..1] **templateId** where it contains exactly one [1..1] **@root="2.16.840.1.113883.10.20.22.4.31"**.

This means that you must have a template id with @root="2.16.840.1.113883.10.20.22.4.31", but you can also have other template ids with different valued attributes.

The following figure shows a typical template's set of constraints presented in this guide. The next chapters describe specific aspects of conformance statements—open vs. closed templates, conformance verbs, cardinality, vocabulary conformance, containment relationships, and null flavors.

Figure 12: Constraint Conformance Including "such that it" Syntax Example

```
Age Observation:

[observation: identifier urn:oid:2.16.840.1.113883.10.20.22.4.31 (open)]

1. SHALL contain exactly one [1..1] @classCode="OBS" Observation (CodeSystem: HL7ActClass 2.16.840.1.113883.5.6 STATIC) (CONF:81-7613).

2. SHALL contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001 STATIC) (CONF:81-7614).

3. SHALL contain exactly one [1..1] templateId (CONF:81-7899) such that it
```

a. **SHALL** contain exactly one [1..1] @root="2.16.840.1.113883.10.20.22.4.31" (CONF:81-10487).

5.2.2 Template Versioning

A new version of an existing implementation guide reuses templates from the previous version. During the ballot phase or update phase, templates carry the designation "Published" to indicate the template is unchanged from the previous version or "Draft" to indicate a new or revised template. Substantial revisions to previously published templates are indicated by the version number (V2, V3, etc.) in all phases: ballot, update, and published guides.

If there are no substantive changes to a template that has been successfully published, the template will carry the same templateId/@root (identifier oid) and templateId/@extension as in the previous implementation guide. (In the case of older templates, the @extension attribute will not be present.) During a new ballot or update phase, "Published" is appended to the main heading for the template to indicate that the template cannot be commented on in the ballot or update. The "Published" designation is removed in the final publication versions.

A revised version of a previously published template keeps the same templateId/@root as the previous version but is assigned a new templateId/@extension. The notation "(Vn)" (V2, V3, etc.) is also added to the template name. Versions are not necessarily forward or backward compatible. A versioning may be due to substantive changes in the template or because a contained template has changed. The "(Vn)" designation is persistent; it appears with that template when it is used in subsequent guides. During a new ballot or update phase, "Draft" is appended to the main heading for the template to indicate that it may be voted on in the ballot or commented on in the update; the "Draft" designation is removed in the final publication versions.

A new version of a template is explicitly linked to the prior version, enabling the automatic generation of the detailed change log found in Volume 2, Chapter 9 "Changes From Previous Version".

An example the change log for a versioned template is shown in the following figure. In this example, Medication Activity (2.16.840.1.113883.10.20.22.4.16) has versioned to Medication Activity (V2) (2.16.840.1.113883.10.20.22.4.16:2014-06-09).

Figure 13: Versioned Template Change Log Example

Change	Old	New
Name	Medication Activity	Medication Activity (V2)
Oid	urn:oid:2.16.840.1.113883.1 0.20.22.4.16	urn:hl7ii:2.16.840.1.113883. 10.20.22.4.16:2014-06-09
Description	A medication activity describes	A medication activity describes
CONF #: 1098-30822 Added		SHALL contain exactly one [11] Drug Monitoring Act (NEW) (identifier: urn:oid:2.16.840.1.113883.1 0.20.22.4.123) (CONF:1098-30822).
CONF #: 81-7511 Removed	SHALL contain exactly one [11] low (CONF:81-7511).	
CONF #: 1098-7516 Modified	SHOULD contain zero or one [01] doseQuantity	SHALL contain exactly one [11] doseQuantity

Structured Documents Working Group collaborated with Templates Working Group to establish template versioning recommendations, recently published in the following specification: <a href="https://document.com/https://doc

5.2.3 Open and Closed Templates

In open templates, all of the features of the CDA R2 base specification are allowed except as constrained by the templates. By contrast, a closed template specifies everything that is allowed and nothing further may be included.

Estimated Date of Delivery (templateId 2.16.840.1.113883.10.20.15.3.1) is an example of a closed template in this guide.

Open templates allow HL7 implementers to develop additional structured content not constrained within this guide. HL7 encourages implementers to bring their use cases forward as candidate requirements to be formalized in a subsequent version of the standard to maximize the use of shared semantics.

5.2.4 Conformance Verbs (Keywords)

The keywords **shall**, **should**, **may**, **need not**, **should not**, and **shall not** in this document are to be interpreted as described in the HL7 Version 3 Publishing Facilitator's Guide.¹¹

- **SHALL**: an absolute requirement
- SHALL NOT: an absolute prohibition against inclusion
- **SHOULD/SHOULD NOT**: best practice or recommendation. There may be valid reasons to ignore an item, but the full implications must be understood and carefully weighed before choosing a different course
- **MAY/NEED NOT**: truly optional; can be included or omitted as the author decides with no implications

The keyword "SHALL" allows the use of nullFlavor unless the requirement is on an attribute or the use of nullFlavor is explicitly precluded.

When conformance statements are nested (or have subordinate clauses) the conformance statements are to be read and interpreted in hierarchical order. These hierarchical clauses can be interpreted as "if then, else" clauses. Thus...

- a. This structuredBody **SHOULD** contain zero or one [0..1] **component** (CONF:1098-29066) such that it
 - i. SHALL contain exactly one [1..1] Plan of Treatment Section (V2) (identifier: urn:h17ii:2.16.840.1.113883.10.20.22.2.10:2014-06-09) (CONF:1098-29067).

...is understood as:

- a. It is recommended (**SHOULD**) that the structureBody contains a component.
 - i. **If** the component exists, **then** it must contain a Plan of Treatment Section (V2),
 - ii. **else** the component does not exist, and the conformance statement about the Plan of Treatment Section (V2) should be skipped.

¹¹ HL7, Version 3 Publishing Facilitator's Guide. http://www.hl7.org/v3ballot/html/help/pfg/pfg.htm

In the case where the higher level conformance statement is a **SHALL**, there is no conditional clause. Thus...

- a. This structuredBody **SHALL** contain exactly one [1..1] **component** (CONF:1098-29086) such that it
 - i. SHALL contain exactly one [1..1] <u>Problem Section (entries required) (V2)</u> (identifier: urn:h17ii:2.16.840.1.113883.10.20.22.2.5.1:2014-06-09) (CONF:1098-29087).

...means that the structuredBody is always required to have a component.

5.2.5 Cardinality

The cardinality indicator (0..1, 1..1, 1..*, etc.) specifies the allowable occurrences within a document instance. The cardinality indicators are interpreted with the following format "m...n" where m represents the least and n the most:

- 0..1 zero or one
- 1..1 exactly one
- 1..* at least one
- 0..* zero or more
- 1..n at least one and not more than n

When a constraint has subordinate clauses, the scope of the cardinality of the parent constraint must be clear. In the next figure, the constraint says exactly one participant is to be present. The subordinate constraint specifies some additional characteristics of that participant.

Figure 14: Constraints Format – only one allowed

```
1. SHALL contain exactly one [1..1] participant (CONF:2777).

a. This participant SHALL contain exactly one [1..1] @typeCode="LOC"

(CodeSystem: 2.16.840.1.113883.5.90 HL7ParticipationType)

(CONF:2230).
```

In the next figure, the constraint says only one participant "like this" is to be present. Other participant elements are not precluded by this constraint.

Figure 15: Constraints Format - only one like this allowed

```
1. SHALL contain exactly one [1..1] participant (CONF:2777) such that it

a. SHALL contain exactly one [1..1] @typeCode="LOC" (CodeSystem: 2.16.840.1.113883.5.90 HL7ParticipationType) (CONF:2230).
```

5.2.6 Optional and Required with Cardinality

The terms optional and required describe the lower bound of cardinality as follows:

Optional means that the number of allowable occurrences of an element may be 0; the cardinality will be expressed as [0..1] or [0..*] or similar. In these cases, the element

may not be present in the instance. Conformances formulated with **MAY** or **SHOULD** are both considered "optional" conformances.

Required means that the number of allowable occurrences of an element must be at least 1; the cardinality will be expressed as [m..n], where $m \ge 1$ and $n \ge 1$ (for example, [1..1] or [1..*]). In these cases, the element must be present in the instance. Conformance statements formulated with SHALL are required conformances. If an element is required but it is not known, the @nullFlavor attribute must be used. See Unknown and No Known Information.

5.2.7 Vocabulary Conformance

The templates in this document use terms from several code systems. These vocabularies are defined in various supporting specifications and may be maintained by other bodies, as is the case for the LOINC® and SNOMED CT® vocabularies.

Note that value set identifiers (e.g., ValueSet 2.16.840.1.113883.1.11.78 Observation Interpretation (HL7) **DYNAMIC**) used in the binding definitions of template conformance statements do not appear in the XML instance of a CDA document. The definition of the template must be referenced to determine or validate the vocabulary conformance requirements of the template.

Value set bindings adhere to HL7 Vocabulary Working Group best practices, and include both an indication of stability and of coding strength for the binding. Value set bindings can be **STATIC**, meaning that they bind to a specified version of a value set, or **DYNAMIC**, meaning that they bind to the most current version of the value set. If a **STATIC** binding is specified, a date **SHALL** be included to indicate the value set version. If a **DYNAMIC** binding is specified, the value set authority and link to the base definition of the value set **SHALL** be included, if available, so implementers can access the current version of the value set. When a vocabulary binding binds to a single code, the stability of the binding is implicitly **STATIC**.

Figure 16: Binding to a Single Code

- 2. SHALL contain exactly one [1..1] code (CONF:15403).
 - a) This code **SHALL** contain exactly one [1..1] @code="11450-4" Problem List (CONF:15408).
 - b) This code **SHALL** contain exactly one [1..1] @codeSystem="2.16.840.1.113883.6.1" (CodeSystem: LOINC 2.16.840.1.113883.6.1 **STATIC**) (CONF: 31141).

The notation conveys the actual code (11450-4), the code's displayName (Problem List), the OID of the codeSystem from which the code is drawn (2.16.840.1.113883.6.1), and the codeSystemName (LOINC).

HL7 Data Types Release 1 requires the codeSystem attribute unless the underlying data type is "Coded Simple" or "CS", in which case it is prohibited. The displayName and the codeSystemName are optional, but recommended, in all cases.

The above example would be properly expressed as follows.

Figure 17: XML Expression of a Single-Code Binding

A full discussion of the representation of vocabulary is outside the scope of this document; for more information, see the *HL7 V3 Normative Edition 2010*¹² sections on Abstract Data Types and XML Data Types R1.

There is a discrepancy between the HL7 R1 Data Types and this guide in the in the implementation of translation code versus the original code. The R1 data type requires the original code in the root. The convention agreed upon for this implementation guide specifies a code from the required value set be used in the element and other codes not included in the value set are to be represented in a translation for the element. This discrepancy is resolved in HL7 Data Types R2.

In the next example, the conformant code is SNOMED-CT code 206525008.

Figure 18: Translation Code Example

```
<code code='206525008'
    displayName='neonatal necrotizing enterocolitis'
    codeSystem='2.16.840.1.113883.6.96'
    codeSystemName='SNOMED CT'>
    <translation code='NEC-1'
    displayName='necrotizing enterocolitis'
    codeSystem='2.16.840.1.113883.19'/>
</code>
```

Value set tables are present below a template, or are referenced if they occur elsewhere in the specification, when there are value set bindings in the template. The value set table provides the value set identifier, a description, and a link to the source of the value set when possible. Ellipses in the last row indicate the value set members shown are examples and the true source must be accessed to see all members.

If a value set binding has a **DYNAMIC** stability, implementers creating a CDA document must go to the location in the URL to check for the most current version of the value set expansion.

¹² HL7 Version 3 Interoperability Standards, http://www.hl7.org/memonly/downloads/v3edition.cfm - V32010

Table 4: Example Value Set Table (Referral Types)

Value Set: Referral Types 2.16.840.1.113883.11.20.9.56

A value set of SNOMED-CT codes descending from "3457005" patient referral (procedure).

Value Set Source:

http://vtsl.vetmed.vt.edu/TerminologyMgt/RF2Browser/ISA.cfm?SCT ConceptID=3457005

Code	Code System	Code System OID	Print Name
44383000	SNOMED CT	2.16.840.1.113883.6.96	Patient referral for consultation
391034007	SNOMED CT	2.16.840.1.113883.6.96	Refer for falls assessment (procedure)
86395003	SNOMED CT	2.16.840.1.113883.6.96	Patient referral for family planning (procedure)
306106002	SNOMED CT	2.16.840.1.113883.6.96	Referral to intensive care service (procedure)
306140002	SNOMED CT	2.16.840.1.113883.6.96	Referral to clinical oncology service (procedure)
396150002	SNOMED CT	2.16.840.1.113883.6.96	Referral for substance abuse (procedure)
	•	•	

5.2.8 Data Types

All data types used in a CDA document are described in the CDA R2 normative edition. ¹³ All attributes of a data type are allowed unless explicitly prohibited by this specification.

5.2.9 Document-Level Templates "Properties" Heading

In Volume 2 of this implementation guide, each document-level template has a "Properties" heading for ease of navigation. The Properties heading is an organizational construct, underneath which relevant CDA act-relationships and roles are called out as headings in the document.

5.3 XML Conventions Used in This Guide

5.3.1 XPath Notation

Instead of the traditional dotted notation used by HL7 to represent RIM classes, this document uses XML Path Language (XPath) notation¹⁴ in conformance statements and elsewhere to identify the XML elements and attributes within the CDA document instance to which various constraints are applied. The implicit context of these expressions is the root of the document. This notation provides a mechanism that will be familiar to developers for identifying parts of an XML document.

XPath statements appear in this document in a monospace font.

¹³ HL7 CDA Release 2. http://www.hl7.org/implement/standards/product_brief.cfm?product_id=7

¹⁴ W3C, XML Path Language. http://www.w3.org/TR/xpath/

XPath syntax selects nodes from an XML document using a path containing the context of the node(s). The path is constructed from node names and attribute names (prefixed by a '@') and catenated with a '/' symbol.

Figure 19: XML Document Example

In the above example, the code attribute of the code could be selected with the XPath expression in the next figure.

Figure 20: XPath Expression Example

```
author/assignedAuthor/code/@code
```

5.3.2 XML Examples and Sample Documents

Extensible Mark-up Language (XML) examples appear in figures in this document in this monospace font. XML elements (code, assignedAuthor, etc.) and attribute names (SNOMED CT, 17561000, etc.) also appear in this monospace font. Portions of the XML content may be omitted from the content for brevity, marked by an ellipsis (...) as shown in the example below.

Figure 21: ClinicalDocument Example

```
<ClinicalDocument xmls="urn:h17-org:v3">
    ...
</ClinicalDocument>
```

6 REFERENCES

- American Recovery And Reinvestment Act of 2009, US Public Law 111-5, 123
 Stat. 115, 516 (Feb. 19, 2009). http://www.gpo.gov/fdsys/pkg/PLAW-111publ5/content-detail.html
- CDC. (May 2012). Epidemiology and Prevention of Vaccine-Preventable Diseases (The Pink Book), Appendix D: Vaccine Administration Guidelines.
 http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/appendices/D/vaccadmin.pdf
- Electronic Health Record Incentive Program; Final Rule (Stage 1 Meaningful Use), (July 28, 2010), http://edocket.access.gpo.gov/2010/pdf/2010-17207.pdf
- GS1 website. http://www.gs1.org/
- Health Industry Business Communications Council (HIBBCC) website. http://www.hibcc.org/
- Health Information Technology: Initial Set of Standards, Implementation Specifications, and Certification Criteria for Electronic Health Record Technology; 45 CFR Part 170, Final Rule, (July 28, 2010). http://edocket.access.gpo.gov/2010/pdf/2010-17210.pdf
- Health Information Technology: Standards, Implementation Specifications, and Certification Criteria for Electronic Health Record Technology, 2014 Edition; Revisions to the Permanent Certification Program for Health Information Technology. 45 CFR Part 170, Final rule, (September 4, 2012).
 http://www.gpo.gov/fdsys/pkg/FR-2012-09-04/pdf/2012-20982.pdf
- HITSP Summary Documents Using HL7 Continuity of Care Document (CCD)
 Component (HITSP/C32) webpage, Versions 2.1, 2.2, 2.3, 2.5. (December 13, 2007 July 8, 2009).

 http://www.hitsp.org/ConstructSet_Details.aspx?&PrefixAlpha=4&PrefixNumeric=32
- HL7 Ballot Desk Top webpage.
 http://www.hl7.org/ctl.cfm?action=ballots.loginchoice (Must be an HL7 member to view.)
- HL7 Care Plan webpage. http://wiki.hl7.org/index.php?title=Care_Plan
- *HL7 Clinical Document Architecture, Release 2 (CDA R2).* (May 2005). http://www.hl7.org/implement/standards/product_brief.cfm?product_id=7
- HL7 Implementation Guide for CDA Release 2: Additional Information Specification Implementation Guide, Release 3.0. (July 31, 2007).
 http://www.hl7.org/documentcenter/public/wg/ca/CDAR2AIS0000R030_ImplementationGuideDraft.pdf
- *HL7 Implementation Guide for CDA Release 2: Consultation Notes*, (U.S. Realm), Draft Standard for Trial Use, Release 1, Levels 1, 2, and 3, DSTU Updated: January 2010.

- http://www.hl7.org/documentcenter/public/standards/dstu/CDAR2_CONSNOTE DSTU R1 2010JAN.zip
- *HL7 Implementation Guide for CDA Release 2: History and Physical (H&P) Notes* (U.S. Realm) Draft Standard for Trial Use, Release 1, Levels 1, 2, and 3 A CDA Implementation guide for History and Physical Notes, DSTU Updated: January 2010.
 - http://www.hl7.org/implement/standards/product_brief.cfm?product_id=19
- HL7 Implementation Guide for CDA Release 2: Imaging Integration, Levels 1, 2, and 3, Basic Imaging Reports in CDA and DICOM Diagnostic Imaging Reports (DIR) Universal Realm, Release 1.0. (March 2009).
 http://www.hl7.org/documentcenter/private/standards/cda/igs/cdar2_ii_bimg rpts r1 inf 2009mar.zip
- HL7 Implementation Guide for CDA Release 2: Procedure Note (Universal Realm),
 Draft Standard for Trial Use, Release 1, Levels 1, 2, and 3, July 2010.
 http://www.hl7.org/documentcenter/public/standards/dstu/CDAR2_IG_PROCNOTE_DSTU_R1_2010JUL.zip
- *HL7 Implementation Guide for CDA Release 2: Unstructured Documents*, Release 1, Level 1 (Universal Realm), Draft Standard for Trial Use, September 2010. http://www.hl7.org/implement/standards/product_brief.cfm?product_id=259
- *HL7 Implementation Guide for CDA Release 2.0 Operative Note*, (U.S. Realm), Draft Standard for Trial Use, Release 1, Levels 1, 2 and 3, Published. (March 2009).
- HL7 Implementation Guide for CDA Release 2.0, Care Record Summary Release 2, Discharge Summary, (U.S. Realm) Draft Standard for Trial Use, Levels 1, 2 and 3. (December 2009).
 http://www.hl7.org/implement/standards/product_brief.cfm?product_id=233
- *HL7 Implementation Guide for CDA Release 2.0, Progress Note* (U.S. Realm), Draft Standard for Trial Use, Levels 1, 2, and 3. (January 2011). http://www.hl7.org/implement/standards/product_brief.cfm?product_id=188
- HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation (US Realm), Release 1.1. (July 2012).
 http://www.hl7.org/documentcenter/private/standards/cda/CDAR2_IG_IHE_CONSOL_DSTU_R1dot1_2012JUL.zip
- HL7 Implementation Guide: CDA Release 2 Continuity of Care Document (CCD),
 Release 1, A CDA implementation of ASTM E2369-05 Standard Specification for
 Continuity of Care Record© (CCR). (April 01, 2007).
 http://www.hl7.org/documentcenter/private/standards/cda/igs/HL7_CCD_final.zip
- HL7 Version 3 Domain Analysis Model: Pressure Ulcer Prevention, Release 1.
 (August 2013).
 http://www.hl7.org/documentcenter/private/standards/v3/V3_DAM_PRULCE
 <a href="http://www.hl7.org/documentcenter/private/standards/v3/V3_
- *HL7 Version 3 Interoperability Standards*, Normative Edition 2010. http://www.hl7.org/memonly/downloads/v3edition.cfm - V32010

- HL7 Version 3 Publishing Facilitator's Guide, Release 1. (2005).
 http://www.hl7.org/v3ballot/html/help/pfg/pfg.htm
- HL7 Version 3 Standard: Refinement, Constraint and Localization to Version 3
 Messages, Release 2. (September 2012).
 http://www.hl7.org/v3ballot/html/infrastructure/conformance/conformance.ht
 m
- IHE Patient Care Coordination (PCC) Technical Framework Supplement, Patient Care Plan Content Profile (PtCP), Trial Implementation (October 4, 2013). http://www.ihe.net/uploadedFiles/Documents/PCC/IHE_PCC_Suppl_PtCP.pdf
- IHE. (2009). Cross Transaction Specifications and Content Specifications. IHE ITI Technical Framework, Volume 3 (ITI TF-3) (see 5.2 Scanned Documents Content Model). http://www.ihe.net/Technical_Framework/upload/IHE_ITI_TF_6-0_Vol3_FT_2009-08-10.pdf
- International Council for Commonality in Blood Bank Automation (ICCBBA) website. http://iccbba.org/
- Joint Commission Requirements for Discharge Summary (JCAHO IM.6.10 EP7).
- Mosby's Medical Dictionary, 8th edition. © 2009, Elsevier.
- Pressure Ulcer Prevention Domain Analysis Model, current, accessed May 2, 2012.
- S&I Framework, LCC Long-Term Post-Acute Care (LTPAC) Transition SWG webpage.
 http://wiki.siframework.org/LCC+Long-Term+Post-Acute+Care+%28LTPAC%29+Transition+SWG
- S&I Framework, LCC Longitudinal Care Plan (LCP) SWG web page.
 http://wiki.siframework.org/LCC+Longitudinal+Care+Plan+%28LCP%29+SWG
- SMART Platforms website. http://smartplatforms.org
- Taber's Cyclopedic Medical Dictionary, 21st Edition, F.A. Davis Company. http://www.tabers.com
- Terminfo. http://www.hl7.org/special/committees/terminfo/index.cfm
- Trifolia Workbench (Consolidation Project Edition). Open source template database. https://trifolia.lantanagroup.com/
- W3C, Extensible Markup Language (XML) 1.0 (Fifth Edition), http://www.w3.org/TR/2008/REC-xml-20081126/
- W3C, XML Path Language (XPath), Version 1.0, (1999).
 http://www.w3.org/TR/xpath/

APPENDIX A — ACRONYMS AND ABBREVIATIONS

ADL Activities of Daily Living

C-CDA R1, R1.1, R2 Consolidated CDA (Release 1, 1.1, and 2)

CCD Continuity of Care Document

CDA, CDA R2 Clinical Document Architecture (Release 2)
CDC Centers for Disease Control and Prevention

CFR Code of Federal Regulations

CID content identifier

CPT Current Procedural Terminology
CVX Codes for Vaccine Administered

DAM Domain Analysis Model

DI device identifier

DICOM Digital Imaging and Communications in Medicine

CMET Common Message Element Type

DIR Diagnostic Imaging Report

DME durable medical equipment

DRIV is derived from

DSTU Draft Standard for Trial Use

EHR electronic health record

EMR electronic medical record

EVN event

FDA Food and Drug Administration

FIPS Federal Information Processing Standards

GOL goal

H&P History and Physical

HCT/P Human Cell & Tissue Products

HHPoC Home Health Plan of Care

HIBCC Health Industry Business Communications Council

HIE health information exchange

HISP health information service provider
HIT healthcare information technology

HITSP Health Information Technology Standards Panel

HL7 Health Level Seven

HTML Hypertext Markup Language

IADL Instrumental Activities of Daily Living

ICCBBA International Council for Commonality in Blood Banking

Automation, Inc.

ICD International Classification of Diseases

ICF International Classification of Functioning, Disability and Health

IG implementation guide

IHE Integrating the Healthcare Enterprise

PCC Patient Care Coordination

IHTSDO International Health Terminology Standard Development

Organisation

RFC Request for Comments

ITI information technology infrastructure

LCC Longitudinal Care Coordination

LCP Longitudinal Care Plan

LOINC Logical Observation Identifiers Names and Codes

LTPAC Long-Term Post-Acute Care

MHTML MIME HTML

MIME Multipurpose Internet Mail Extensions

MPHO Medical Products of Human Origin

NA not applicable

NDC National Drug Code

NDFRT National Drug File Reference Terminology

NEMA National Electrical Manufacturers Association

NHSN National Healthcare Safety Network

NI no information

NLM National Library of Medicine

NPI National Provider Identifier

NPO nothing by mouth

NPP non-physician provider

NUBC National Uniform Billing Committee

NUCC National Uniform Claim Committee

OID object identifier

ONC Office of National Coordinator

OTH not an element in the value domain

PCDATA Parsed Character Data

PCWG Patient Care Work Group

PDF Portable Document Format

PGD Patient Generated Document

PGP Pretty Good Privacy

PHQ Patient Health Questionnaire

PHR personal health record
PI Production Identifier

PKCS#7 public-key cryptography standard seven (Cryptographic Message

Syntax Standard

RFC request for comment

RIM Reference Information Model

RMIM Refined Message Information Model

RQO request

RSNA Radiological Society of North America

S&I Standards and Interoperability

sdtc Standard Duty Title Code

SDWG Structured Documents Working Group

SMART Substitutable Medical Applications & Reusable Technology

SNF Skilled Nursing Facility

SNOMED CT Systemized Nomenclature for Medicine – Clinical Terms

SOP Service Object Pair

SPL Structured Product Labeling

SR Structured Report

SSN Social Security Number

SWG Sub Work Group

TPN Total Parenteral Nutrition

UCUM Unified Code for Units of Measure

UD Unstructured Document

UDI Unique Device Identification
UML Unified Modeling Language
UNII Unique Ingredient identifier

UNK unknown

URL uniform resource locator

URN uniform resource name

UUID universally unique identifier

VIS Vaccine Information Statement

WADO Web Access to Persistent DICOM Objects

XDS-SD Cross Enterprise Sharing of Scanned Documents

XML eXtensible Markup language

XML-DSIG XML digital signature
XPath XML Path Language

APPENDIX B — HIGH-LEVEL CHANGE LOG

This implementation guide builds on *HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation (US Realm), Release 1.1.* (July 2012). Please refer to Volume 2 Chapter 9, "Changes From Previous Version", for a detailed change log.

Volume 1 Summary of Changes

- Chapter 3.1 CDA Participations—New text describing CDA approach to participations.
- Chapter 3.2 Determining a Clinical Statement's Status—New text describing how to know if an entry is active, resolved, etc.
- Chapter 3.4 Unknown and No Known Information—Merged and revised Consolidated CDA R1.1 "Section 1.8.8 Null Flavor" and "Section 1.8.9 Unknown Information"; additional examples.
- Chapter 4.2.2 Template Versioning—New text describing C-CDA approach to template versioning.

Volume 2 Summary of Changes

Table 5: High-Level Change Log

Type of Template	Template	Summary of New Content or Update to Template
US Realm Header	US Realm Header (V2) 2.16.840.1.113883.10.20.22.1.1:2014-06-09	Tighten constraint for patientRole/patient/raceCode (MAY to SHOULD) to support MU Stage 2 requirements for exchange documents. Patient race is included in the Common MU Data Set and has associated vocabulary requirement.
		Tighten constraint for patientRole/patient/ethnicGroupCode (MAY to SHOULD) to support MU Stage 2 requirements for exchange documents. Patient ethnicity is included in the Common MU Data Set and has associated vocabulary requirement.
		Clarify constraint for patientRole/patient/sdtc:raceCode to require raceCode to be present.
		Added extension for stdc: signatureText extension for digital signature
		Constrain Language value set for patientRole/patient/ languageCommunication to include ISO-639-2 alpha-3 codes with corresponding alpha-2 codes as required by MU Stage 2 for recording patient language. Language value set includes all codes from RFC-4646, which include ISO-639 codes.
		Tighten constraint for patientRole/patient/languageCommunication/ preferenceInd (MAY to SHOULD) to support MU Stage 2 requirements for exchange documents and patient engagement initiatives.
Document	Care Plan (NEW) 2.16.840.1.113883.10.20.22.1.15	A Care Plan (including Home Health Plan of Care (HHPoC)) is a consensus-driven dynamic plan that represents a patient's and Care Team Members' prioritized concerns, goals, and planned interventions. The CDA Care Plan is an instance in time of a dynamic Care Plan.
Document	Consultation Note (V2) 2.16.840.1.113883.10.20.22.1.4:2014-06-09	Consultation Note is generated as a result of a request from a clinician for an opinion or advice from another clinician.
Document	Continuity of Care Document (CCD) (V2) 2.16.840.1.113883.10.20.22.1.2:2014-06-09	Tighten constraint for Vital Signs Section (entries required) (MAY to SHALL) to support MU Stage 2 requirements for exchange documents. Patient vital signs are included in the Common MU Data Set. Tighten constraint for Procedures Section (entries required) (MAY to SHALL) to
		support MU Stage 2 requirements for exchange documents. Patient procedures

Type of Template	Template	Summary of New Content or Update to Template
		are included in the Common MU Data Set and have associated vocabulary requirements.
Document	Diagnostic Imaging Report (V2) 2.16.840.1.113883.10.20.22.1.5:2014-06-09	Updated to reference a contained template that has been versioned.
Document	Discharge Summary (V2) 2.16.840.1.113883.10.20.22.1.8:2014-06-09	Updated to reference a contained template that has been versioned.
Document	History and Physical (V2) 2.16.840.1.113883.10.20.22.1.3:2014-06-09	Updated to reference a contained template that has been versioned.
Document	Operative Note (V2) 2.16.840.1.113883.10.20.22.1.7:2014-06-09	Updated to reference a contained template that has been versioned.
Document	Procedure Note (V2) 2.16.840.1.113883.10.20.22.1.6:2014-06-09	Updated to reference a contained template that has been versioned.
Document	Progress Note (V2) 2.16.840.1.113883.10.20.22.1.9:2014-06-09	Updated to reference a contained template that has been versioned.
Document	Referral Note (NEW) 2.16.840.1.113883.10.20.22.1.14	The Referral Note type communicates pertinent patient information to the consulting provider from a referring provider
Document	Transfer Summary (NEW) 2.16.840.1.113883.10.20.22.1.13	A Transfer Summary exchanges information between providers when a patient moves between health care settings.
Document	Unstructured Document (V2) .16.840.1.113883.10.20.22.1.10:2014-06-09	Updated to reference a contained template that has been versioned.
Document	US Realm Header - Patient Generated Document (NEW) 2.16.840.1.113883.10.20.29.1	This template can be used in conjunction with the US C-CDA General Header. It includes further constraints to address cases when the patient, a patient-controlled device, or a person who has a personal or legal relationship to the patient, authors the document.
Section	Advance Directives Section (entries optional) (V2) 2.16.840.1.113883.10.20.22.2.21:2014-06-09 Advance Directives Section (entries required) (V2) 2.16.840.1.113883.10.20.22.2.21.1:2014-06-09	Section contains the new Advance Directive Organizer.

Type of Template	Template	Summary of New Content or Update to Template
Section	Allergies Section (entries required) (V2) 2.16.840.1.113883.10.20.22.2.6.1:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Anesthesia Section (V2) 2.16.840.1.113883.10.20.22.2.25:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Assessment and Plan Section (V2) 2.16.840.1.113883.10.20.22.2.9:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Complications Section (V2) 2.16.840.1.113883.10.20.22.2.37:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Encounters Section (entries optional) V2 2.16.840.1.113883.10.20.22.2.22:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Encounters Section (entries required) V2 2.16.840.1.113883.10.20.22.2.22.1:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Functional Status Section (V2) 2.16.840.1.113883.10.20.22.2.14:2014-06-09	Functional Status section focuses on physical observations and assessments of a patient's ability. The cognitive status and pressure ulcer content was moved. Cognitive status content is in the Mental Status Section and Pressure Ulcer observations merged into the Wound Observation, which is located in the Physical Exam Section with additional pressure ulcer content. NOTE: Functional Status Problem Observation was deprecated.
Section	Goals Section (NEW) 2.16.840.1.113883.10.20.22.2.60	This template represents patient Goals. A goal is a defined outcome or condition to be achieved in the process of patient care.
Section	Health Concerns Section (NEW) 2.16.840.1.113883.10.20.22.2.58	The Health Concerns section contains data that describes an interest or worry about a health state or process that has the potential to require attention, intervention, or management.
Section	Health Status Evaluations and Outcomes Section (NEW) 2.16.840.1.113883.10.20.22.2.61	This template is a section that contains Health Status Evaluations and Outcomes.
Section	History of Past Illness Section (V2) 2.16.840.1.113883.10.20.22.2.20:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Hospital Admission Diagnosis Section (V2) 2.16.840.1.113883.10.20.22.4.34:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Admission Medications Section (entries optional)	Updated to reference a contained template that has been versioned.

Type of Template	Template	Summary of New Content or Update to Template
	(V2) 2.16.840.1.113883.10.20.22.2.44:2014-06-09	
Section	Discharge Diagnosis Section (V2) 2.16.840.1.113883.10.20.22.2.24:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Discharge Medications Section (entries optional) (V2) 2.16.840.1.113883.10.20.22.2.11:2014-06-09	Updated to reference a contained template that has been versioned.
Section	Hospital Discharge Medications Section (entries required) (V2)	Updated to reference a contained template that has been versioned.
Section	Immunizations Section (entries optional) (V2)	Updated to reference a contained template that has been versioned.
Section	Immunizations Section (entries required) (V2) 2.16.840.1.113883.10.20.22.2.2.1.2	Updated to reference a contained template that has been versioned.
Section	Instructions Section (V2)	Updated to reference a contained template that has been versioned.
Section	Interventions Section (V2) (2.16.840.1.113883.10.20.21.2.3.2)	Updated description of template to better describe interventions and to clarify that instructions are a subset of interventions. Now includes Intervention Act (NEW).
Section	Medical (General) History Section (V2)	Updated to reference a contained template that has been versioned.
Section	Medical Equipment Section (V2) 2.16.840.1.113883.10.20.22.2.23.2	Updated to include the Medical Equipment Organizer template and the Medical Device applied template.
Section	Medications Administered Section (V2)	Updated to reference a contained template that has been versioned.
Section	Medication Section (V2) (entries required) 2.16.840.1.113883.10.20.22.2.1.1.2 Medication Section (V2) (entries optional) 2.16.840.1.113883.10.20.22.2.1.2	Updated to contain the new Drug Monitoring Act and newer version of Medication Activity (V2).
Section	Mental Status Section (NEW) 2.16.840.1.113883.10.20.22.2.56	Mental Status section contains observation and evaluations related to patient's psychological and mental competency and deficits. NOTE: Cognitive Status Problem Observation was deprecated.
Section	Nutrition Section (NEW) 2.16.840.1.113883.10.20.22.2.57	This template represents diet and nutrition information and patient's nutritional status. Note: Discharge Diet Section was deprecated.

Type of Template	Template	Summary of New Content or Update to Template
Section	Payers Section (V2)	Updated to reference a contained template that has been versioned.
Section	Physical Exam Section (V2)	Updated to include Wound Observation.
Section	Physical Findings of Skin Section (NEW) 2.16.840.1.113883.10.20.22.2.62	This template represents direct observations made by the clinician of the skin.
Section	Plan of Treatment (V2) 2.16.840.1.113883.10.20.22.2.10.2	Previous name: Plan of Care Updated to include Handoff Communication template.
Section	Planned Procedure Section (V2)	Updated to reference a contained template that has been versioned.
Section	Postprocedure Diagnosis Section (V2)	Updated to reference a contained template that has been versioned.
Section	Preoperative Diagnosis Section (V2)	Updated to reference a contained template that has been versioned.
Section	Problem Section (V2)(entries optional) 2.16.840.1.113883.10.20.22.2.5.2 Problem Section (V2)(entries required) 2.16.840.1.113883.10.20.22.2.5.1.2	Updated to contain Health Status Observation, which was in the Problem observation template. This represents a patient's overall general status.
Section	Procedure Findings Section (V2)	Updated to reference a contained template that has been versioned.
Section	Procedure Indications Section (V2)	Updated to reference a contained template that has been versioned.
Section	Procedures Section (entries optional) (V2)	Updated to reference a contained template that has been versioned.
Section	Procedures Section (entries required) (V2)	Updated to reference a contained template that has been versioned.
Section	Reason for Referral (Section) 1.3.6.1.4.1.19376.1.5.3.1.3.1.2	Updated to contain the Patient Referral Activity Act.
Section	Results Section (entries optional) (V2)	Updated to reference a contained template that has been versioned.
Section	Results Section (entries required) (V2) 2.16.840.1.113883.10.20.22.2.3.1.2	Updated to reference a contained template that has been versioned.
Section	Social History Section (V2) 2.16.840.1.113883.10.20.22.2.17.2	Updated to include Caregiver, Cultural and Religious Observation and Characteristics of Home Environment templates.
Section	Vital Signs Section (entries optional) (V2)	Updated to reference a contained template that has been versioned.
Section	Vital Signs Section (entries required) (V2) 2.16.840.1.113883.10.20.22.2.4.1.2	Updated to reference a contained template that has been versioned.

Type of Template	Template	Summary of New Content or Update to Template
Entry	Act Reference (NEW) (2.16.840.1.113883.10.20.22.4.122)	This template represents the act of referencing another entry in the same CDA document instance.
Entry	Admission Medication (V2)	Updated to reference a contained template that has been versioned.
Entry	Advance Directive Observation(V2) 2.16.840.1.113883.10.20.22.4.48.2	Guidance added to allow for documentation of a primary and secondary health care agent or substitute decision makers.
Entry	Advance Directive Organizer (NEW)	This template represents related patient directives.
Entry	Allergy Concern Act (V2) 2.16.840.1.113883.10.20.22.4.30.2	Previous Name: Allergy Problem Act Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119).
Entry	Allergy - Intolerance Observation (V2) 2.16.840.1.113883.10.20.22.4.7.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Clarified that effectiveTime is SHALL, that effectiveTime/low is SHALL, and effectiveTime/high is MAY. Updated value set: In discussion with NLM, FDA, and others, have updated the value set to Substance-Reactant for Intolerance 2.16.840.1.113762.1.4.1010.1. Value set will be housed at NLM's Value Set Authority Center. Deprecated nested Allergy Status Observation
Entry	Characteristics of Home Environment (NEW) 2.16.840.1.113883.10.20.22.4.109	This template represents the patient's home environment including, but not limited to, type of residence, living arrangement, and housing status.
Entry	Cognitive Abilities Observation (NEW) 2.16.840.1.113883.10.20.22.4.126	This template represents a patient's ability to perform specific cognitive tasks.
Entry	Cognitive Status Observation (V2) 2.16.840.1.113883.10.20.22.4.74.2	Previous Name: Cognitive Status Result Observation (V2) Removed conformance to Result Observation. Updated observation code binding.
Entry	Cognitive Status Organizer (V2) 2.16.840.1.113883.10.20.22.4.75.2	Previous Name: Cognitive Status Result Organizer (V2) Removed conformance to Result Organizer.
Entry	Coverage Activity (V2)	Updated to reference a contained template that has been versioned.
Entry	Cultural and Religious Observation (NEW) 2.16.840.1.113883.10.20.22.4.111	This template represents a patient's spiritual, religious, and cultural belief practices.

Type of Template	Template	Summary of New Content or Update to Template
Entry	Current Smoking Status (V2) 2.16.840.1.113883.10.20.22.4.78.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Change name Smoking Status template and value set to Current Smoking Status to support MU Stage 2 requirements for exchange documents. Remove constraint on effectiveTime/low in conjunction with name change to clarify time to be recorded as the observation time and not the time when the patient is/was/never was smoking. Change observation/code from "ASSERTION" to SNOMED 229819007 Tobacco use and exposure (observable). Added two SNOMED CT codes to Current Smoking Status value set and Tobacco Use value set. The two additions are: "Heavy tobacco smoker" 428071000124103; "Light tobacco smoker" 428061000124105.
Entry	Deceased Observation (V2)	Updated to reference a contained template that has been versioned.
Entry	Discharge Medication (V2)	Updated to reference a contained template that has been versioned.
Entry	Drug Monitoring Act (NEW) 2.16.840.1.113883.10.20.22.4.123	This template represents a drug monitoring therapy and reflect what a clinician intends a patient to be taking.
Entry	Encounter Activity (V2)	Updated to reference a contained template that has been versioned.
Entry	Encounter Diagnosis (V2)	Updated to reference a contained template that has been versioned.
Entry	Functional Status Observation (V2) 2.16.840.1.113883.10.20.22.4.67.2	Previous Name: Functional Status Result Observation (V2) Observation code binding was updated.
Entry	Functional Status Organizer (V2) 2.16.840.1.113883.10.20.22.4.66.2	Previous Name: Functional Status Result Organizer (V2) An author was added and an entryRelationship for Self -Care Activities. Removed conformance to Result Organizer.
Entry	Goal Observation (NEW) (2.16.840.1.113883.10.20.22.4.121)	This template represents a patient care goal.
Entry	Handoff Communication (NEW) 2.16.840.1.113883.10.20.22.4.141	This template represents provider hand-off communication such as in a transfer of care situation.
Entry	Health Concern Act (NEW) (2.16.840.1.113883.10.20.22.4.132)	This template represents a health concern.
Entry	Health Status Observation (V2)	Updated to reference a contained template that has been versioned.

Type of Template	Template	Summary of New Content or Update to Template
Entry	Hospital Admission Diagnosis (V2)	Updated to reference a contained template that has been versioned.
Entry	Hospital Discharge Diagnosis (V2)	Updated to reference a contained template that has been versioned.
Entry	Immunization Activity (V2) 2.16.840.1.113883.10.20.22.4.52.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Updated value set: In discussion with NLM, FDA, and others, have updated the preferred value set to be primarily CVX, and MAY use RxNorm as a translation. Value set will be housed at NLM's Value Set Authority Center. Created a new Substance Administered Act (templateId:2.16.840.1.113883.10.20.22.4.118) to better represent the order of a medication or immunization within a series.
Entry	Immunization Medication Information (V2)	Updated to reference a contained template that has been versioned.
Entry	Indication (V2) 2.16.840.1.113883.10.20.22.4.19.2	The id element was updated to reference a problem recorded elsewhere. The id must equal an entry/id in the same document to relate them to each other.
Entry	Instruction (V2)	Updated to reference a contained template that has been versioned.
Entry	Intervention Act (NEW) (2.16.840.1.113883.10.20.21.2.3.2)	This template represents an Intervention Act, and is a wrapper for intervention-type activities considered to be parts of the same intervention.
Entry	Medical Device Applied (NEW) 2.16.840.1.113883.10.20.22.4.115	This template represents devices in or applied to a patient's body.
Entry	Medical Equipment Organizer (NEW) 2.16.840.1.113883.10.20.22.4.135	This clinical statement represents a set of current or historical medical devices/equipment in use or ordered.
Entry	Medication Activity (V2) 2.16.840.1.113883.10.20.22.4.16.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Updated value set: In discussion with NLM, FDA, and others, have updated the value set to Medication Clinical Drug (2.16.840.1.113762.1.4.1010.4). Value set will be housed at NLM's Value Set Authority Center. Created a new Substance Administered Act (templateId:2.16.840.1.113883.10.20.22.4.118) to better represent the order of a medication or immunization within a series. Created a new Drug Monitoring Act (templateId:2.16.840.1.113883.10.20.22.4.123) to reflect medication monitoring activities. Clarified relationship between actStatus and effectiveTime in Volume 1, and

Type of Template	Template	Summary of New Content or Update to Template
		made effectiveTime/low a MAY. Updated to include the updated Instructions (V2) template.
Entry	Medication Dispense (V2)	Updated to reference a contained template that has been versioned.
Entry	Medication Information (V2)	Updated to reference a contained template that has been versioned.
Entry	Medication Supply Order (V2)	Updated to reference a contained template that has been versioned.
Entry	Mental Status Observation (NEW) 2.16.840.1.113883.10.20.22.4.125	This template represents observations relating intellectual, mental capabilities and state of mind.
Entry	Non-Medicinal Supply Activity (V2)	Updated to reference a contained template that has been versioned.
Entry	Nutrition Assessment (NEW) 2.16.840.1.113883.10.20.22.4.138	The template represents the patient's nutrition habits including intake, diet requirements or diet followed.
Entry	Nutrition Recommendation (NEW) 2.16.840.1.113883.10.20.22.4.130	This template represents nutrition regimens, interventions, and procedures.
Entry	Nutrition Status Observation (NEW) 2.16.840.1.113883.10.20.22.4.124	This template describes the overall nutritional status of the patient and findings related to nutritional status.
Entry	Outcome Observation (NEW) (2.16.840.1.113883.10.20.22.4.144)	This template represents an Outcome Observation. An Outcome Observation evaluates a goal and is the actual outcome of an intervention(s).
Entry	Patient Priority Preference (NEW)	Updated to reference a contained template that has been versioned.
Entry	Patient Referral Act (NEW) 2.16.840.1.113883.10.20.22.4.140	This template represents the type of referral whether full care or shared care.
Entry	Physician of Record Participant (V2)	Updated to reference a contained template that has been versioned.
Entry	Planned Act (V2) 2.16.840.1.113883.10.20.22.4.39.2	Previous name: Plan of Care Activity Act Revised to align more closely with their similar non-plan templates where applicable. Updated description and template with additional guidance constraints around effective time, performer, participants, and priority.
Entry	Planned Coverage (NEW) 2.16.840.1.113883.10.20.22.4.129	This template represents the amount of insurance coverage that is projected to cover an act or procedure.
Entry	Planned Encounter (V2) 2.16.840.1.113883.10.20.22.4.40.2	Previous name: Plan of Care Activity Encounter Revised to align more closely with their similar non-plan templates where applicable. Updated description and added additional guidance constraints

Type of Template	Template	Summary of New Content or Update to Template
		around effective time, performer, participants, and priority.
Entry	Planned Immunization Activity (NEW) 2.16.840.1.113883.10.20.22.4.120	The template describes planned immunizations.
Entry	Planned Observation (V2) 2.16.840.1.113883.10.20.22.4.44.2	Previous name: Plan of Care Activity Observation Revised to align more closely with an observation template where applicable. Updated description and added additional guidance constraints around code, value, effective time, performer, participants, and priority.
Entry	Planned Procedure (V2) 2.16.840.1.113883.10.20.22.4.41.2	Previous name: Plan of Care Activity Procedure Revised to align more closely with their similar non-plan template where applicable. Updated description and added additional guidance constraints around effective time, performer, participants, and priority.
Entry	Planned Medication Activity (V2) 2.16.840.1.113883.10.20.22.4.42.2	Previous name: Plan of Care Activity Substance Administration Revised to align more closely with their similar non-plan templates where applicable. Updated description and added additional guidance constraints around effective time, performer, participants, and priority.
Entry	Planned Supply (V2) 2.16.840.1.113883.10.20.22.4.43.2	Previous name: "Plan of Care Activity Supply" Revised to align more closely with their similar non-plan templates where applicable. Updated description and added additional guidance constraints around effective time, performer, participants, and priority.
Entry	Policy Activity (V2)	Updated to reference a contained template that has been versioned.
Entry	Postprocedure Diagnosis (V2)	Updated to reference a contained template that has been versioned.
Entry	Preoperative Diagnosis (V2)	Updated to reference a contained template that has been versioned.
Entry	Problem Concern Act (V2) 2.16.840.1.113883.10.20.22.4.3.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Added new MAY 0* component template "Provider Priority Preference" to assert the importance of the concern to a provider.
Entry	Problem Observation (V2) 2.16.840.1.113883.10.20.22.4.4.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Changed effectiveTime from SHOULD 01 to SHALL 11 Deprecated nested Problem Status Added a new Prognosis Observation template (templateId 2.16.840.1.113883.10.20.22.4.113)

Type of Template	Template	Summary of New Content or Update to Template
		Removed the nested Health Status Observation from within Problem Observation, to now be a top-level clinical statement within Problem Section. Provide guidance on the use of observation/value/qualifier. Provide guidance on the representation of "no known problems". Added new MAY 0* component template "Provider Priority Preference" to assert the importance of the problem to a provider. Added new MAY 0* component template "Patient Priority Preference" to assert the importance of the problem to a patient.
Entry	Procedure Activity Act (V2)	Updated to reference a contained template that has been versioned.
Entry	Procedure Activity Observation (V2)	Updated to reference a contained template that has been versioned.
Entry	Procedure Activity Procedure (V2)	Updated to reference a contained template that has been versioned.
Entry	Prognosis Observation (NEW) 2.16.840.1.113883.10.20.22.4.113	This template represents a prognosis that is associated with a problem or concern.
Entry	Progress Toward Goal Observation (NEW) (2.16.840.1.113883.10.20.22.4.110)	This template represents a patient's progress toward a goal.
Entry	Provider Priority Preference (NEW)	Updated to reference a contained template that has been versioned.
Entry	Reaction Observation (V2)	Updated to reference a contained template that has been versioned.
Entry	Result Observation (V2) 2.16.840.1.113883.10.20.22.4.2.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Add conformance statement for Result Observation value to require units of measure expressed using UCUM for physical quantity data types. Added guidance for representing the original results as obtained from the lab in a <translation></translation>
Entry	Result Organizer (V2) 2.16.840.1.113883.10.20.22.4.1.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Added a MAY constraint, to optionally include an effectiveTime
Entry	Self-Care Activities (ADLs and IADLs) (NEW) 2.16.840.1.113883.10.20.22.4.128	This template represents the adult patient's daily self-care ability
Entry	Sensory and Speech Status (NEW) 2.16.840.1.113883.10.20.22.4.127	This Template represents patient's sensory and speech ability.

Type of Template	Template	Summary of New Content or Update to Template
Entry	Severity Observation (V2)	Updated to reference a contained template that has been versioned.
Entry	Social History Observation (V2) 2.16.840.1.113883.10.20.22.4.38.2	Value Set Social History Type Value Set 2.16.840.1.113883.3.88.12.80.60 STATIC changed to DYNAMIC and expanded.
Entry	Substance Administered Act (NEW)	Updated to reference a contained template that has been versioned.
Entry	Substance or Device Allergy - Intolerance Observation (V2)	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119).
	2.16.840.1.113883.10.20.24.3.90.2	Clarified that effectiveTime is SHALL, that effectiveTime/low is SHALL, and effectiveTime/high is MAY.
		Moved requirement for original text referencing into the Allergy-Intolerance Observation.
		Updated value set: In discussion with NLM, FDA, and others, have updated the value set to Substance-Reactant for Intolerance 2.16.840.1.113762.1.4.1010.1. Value set will be housed at NLM's Value Set Authority Center.
Entry	Tobacco Use (V2) 2.16.840.1.113883.10.20.22.4.85.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119).
Entry	Vital Sign Observation (V2) 2.16.840.1.113883.10.20.22.4.27.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119). Added a constraint, that a PQ @unit be selected from UCUM.
Entry	Vital Signs Organizer (V2) 2.16.840.1.113883.10.20.22.4.26.2	Added new Author Participation template (templateId 2.16.840.1.113883.10.20.22.4.119).
Entry	Wound Characteristics (NEW) 2.16.840.1.113883.10.20.22.4.134	This template represents characteristics of a wound (e.g., integrity of suture line, odor, and erythema).
Entry	Wound Measurement (NEW) 2.16.840.1.113883.10.20.22.4.133	This template represents the Wound Measurement Observations of wound width, depth, and length.
Entry	Wound Observation (NEW) 2.16.840.1.113883.10.20.22.4.114	This template represents acquired or surgical wounds types, wound measurement and characteristics. Pressure ulcer observation content was merged into this template. Note: Pressure Ulcer Observation was deprecated.
Other	Author Participation (NEW) (2.16.840.1.113883.10.20.22.4.119)	Constrains the Author participant.

APPENDIX C — EXTENSIONS TO CDA R2

Where there is a need to communicate information for which there is no suitable representation in CDA R2, extensions to CDA R2 have been developed. These extensions are described above in the context of the section where they are used. This section serves to summarize the extensions and provide implementation guidance.

Extensions created for this guide include:

- sdtc:raceCode The raceCode extension allows for multiple races to be reported for a patient.
- sdtc:id The id extension in the family history organizer on the related subject allows for unique identification of the family member(s).
- sdtc:deceasedInd The deceasedIndextension (= "true" or "false") in the family history organizer on the related subject is used inside to indicate if a family member is deceased.
- sdtc:deceasedTime The deceasedTime extension in the family history organizer on the related subject allows for reporting the date and time a family member died.
- sdtc:birthTime The sdtc:birthTime element allows for the birth date of any person to be recorded. The purpose of this extension is to allow the recording of the subscriber or member of a health plan in cases where the health plan eligibility system has different information on file than the provider does for the patient.
- sdtc:dischargeDispositionCode The sdtc:dischargeDispositionCode element allows the provider to record a discharge disposition in an encounter activity.

To resolve issues that need to be addressed by extension, the developers of this guide chose to approach extensions as follows:

- An extension is a collection of element or attribute declarations and rules for their application to the CDA Release 2.0.
- All extensions are optional. An extension may be used, but need not be under this guide.
- A single namespace for all extension elements or attributes that may be used by this guide will be defined.
- The namespace for extensions created by the HL7 Structured Documents Working Group (formerly Structured Documents Technical Committee) shall be urn:hl7-org:sdtc.

- This namespace shall be used as the namespace for any extension elements or attributes that are defined by this implementation guide.
- Each extension element shall use the same HL7 vocabularies and data types used by CDA Release 2.0.
- Each extension element shall use the same conventions for order and naming as is used by the current HL7 tooling.
- An extension element shall appear in the XML where the expected RIM element of the same name would have appeared had that element not been otherwise constrained from appearing in the CDA XML schema.

APPENDIX D — MIME MULTIPART/RELATED MESSAGES

The following text is taken from the Claims Attachments Implementation Guide (AIS00000) in Section 2.4.15 For up-to-date guidance, refer to the latest edition of that specification.

MIME Multipart/Related Messages

An attachment is comprised of the CDA document, including any supporting files necessary to render the attested content of the document. Two Internet request for comments (RFCs) are needed to properly construct the mime multipart message. When supporting files are needed, the collection of information shall be organized using a MIME multipart/related package constructed according to RFC 2557. Within the MIME package, supporting files must be encoded using Base-64. RFC-4648 should be used when encoding the contents of the MIME package using Base-64. Finally, RFC-2392 may be used to reference other content that appears in the same X12 transaction to use the same content to answer multiple questions for a single claim. Internet RFCs can be downloaded from the RFC editor page at http://www.rfc-editor.org.

RFC-2557 MIME Encapsulation of Aggregate Documents, Such as HTML (MHTML)

This RFC describes how to construct a MIME multipart/related package, and how URLs are resolved within content items of that package. RFC-2557 can be obtained at: http://www.rfc-editor.org/rfc/rfc2557.txt

A MIME multipart/related package is made up of individual content items. Each content item has a MIME header identifying the item. Each content item is delimited from other content items using a string of application specified text. In addition, there must be an ending boundary. The actual content is recorded between these delimiter strings using a BASE-64 encoding of the content item. There is also a MIME header for the entire package.

The first content item of a multipart/related message supporting attachments is the CDA document, containing the header and structured or non-structured body. Subsequent content items included in this package will contain additional content that appears within the body of the document. The CDA document will reference these additional content items by their URLs.

Referencing Supporting Files in Multipart/Related Messages

Because the CDA document and its supporting files may have already existed in a clinical information system, references may already exist within the CDA document to URLs that are not accessible outside of the clinical information system that created the document. When the CDA document is sent via attachments, these URLs may no longer be accessible by the receiving information system. Therefore, each content item that is referenced by a URL within the CDA document must be included as a content item in

http://www.hl7.org/documentcenter/public/wg/ca/CDAR2AIS0000R030_ImplementationGuideDraft.pdf

¹⁵

the MIME package. Each content item may specify the URL by which it is known using the Content-Location header. The receiver of this MIME package shall translate URL references according the RFC-2557. This will ensure resolution of the original URL to the correct content item within the MIME package. Thus, URL references contained within an original document need not be rewritten when the CDA package is transmitted. Instead, these URLs are simply supplied as the value of the Content-Location header in the MIME package.

This capability allows for the same content item to be referred to more than once in a MIME multipart/related package without requiring the content item to be supplied twice. However, it does not allow a separate MIME multipart/related package to contain references to information sent in a previously recorded package.

Referencing Documents from Other Multiparts within the Same X12 Transactions

RFC-2392 is used when referencing content across MIME package boundaries, but still contained within the same X12 transaction (ST to SE). This can occur when the same document answers multiple questions for a single claim. Each component of a MIME package may be assigned a content identifier using the Content-ID header for the content item. For example, this header would appear as:

Content-ID: <07EE4DAC-76C4-4a98-967E-F6EF9667DED1>

This content identifier is a unique identifier for the content item, which means it must never be used to refer to any other content item. RFC-2392 defines the cid: URL scheme (http: and ftp: are two other URL schemes). This URL scheme allows for references by the Content-ID header to be resolved. The URL for the content item identified above would be:

cid:07EE4DAC-76C4-4a98-967E-F6EF9667DED1

Receivers of the MIME multipart message must be able to resolve a cid: URL to the content item that it identifies. Senders must ensure that they only refer to items that have already been transmitted to the receiver by their cid: URL. Thus, this implementation guide prohibits forward URL references using the cid: URL scheme.

Content items shall not be referenced across X12 transactions using the cid: URL scheme. For example, if the payer previously requested information using a 277, and the provider returned that information in a MIME multipart/related package in a 275, and then the payer requested additional information in another 277, the provider may not refer to the content item previously returned in the prior 275 transaction.

APPENDIX E - CARE PLAN RELATIONSHIPS

Care Plan Relationships and HL7 RIM Terms

The HL7 RIM and CDA Refined Message Information Model (RMIM) together with actRelationshipType codes and act moodCodes very effectively express the relationship between Care Plan components. The following tables highlight the main moodCodes and relationshipCodes used in the CDA Care Plan.

Table 6: moodCodes

Code	Display Name	Meaning
EVN	Event	An actual occurrence of an event – something happened
INT	Intent	An intended or planned event
RQO	Request	A request or order to perform something
GOL	Goal	A goal or objective

Table 7: actRelationship TypeCodes

Code	Display Name	Meaning
COMP	Has component	Shows that one component is part of another (overall goal has components consisting of one or more goals)
REFR	Refers to	Shows a general relationship between components (a goal refers to a health concern)
SPRT	Has support	Show that the one component is supporting evidence for another (Health Concern has support from a problem observation)
RSON	Has reason	Show the reason or rationale for a something (intervention has reason of achieving a goal)
GEVL	Evaluates goal	Used to link an observation (intent or actual) to a goal to indicate that the observation evaluates the goal

The following diagram shows the components of a Care Plan and the flow between them, expressed by HL7 moodCodes and relationshipCodes.

HAS COMPONENT HAS COMPONENT Health Goal [mood **EVALUATIONS/OUTCOMES** Concern REFERS TO GOL₁ (code= **Progress** CONCERN or Toward Goal RISK) Observation [mood EVN] [mood EVN] REFERS TO REFERS TO Intervention [mood: INT/ RQO/ etc.] Outcome Observation Observation [mood EVN] HAS REASON [mood EVN]

Figure 22: Care Plan Relationship Diagram

Care Plan Relationships Story Board Example

Joe is a 24 year-old male quadriplegic with diabetes mellitus admitted to an inpatient unit from his home. During the admission assessment, the nurse notes that he has no sensation from the shoulders down. He is confined to a wheelchair and requires two-person assist. His skin is occasionally moist. Joe reports that he is a "good eater" and is on a diabetic diet. The nurse completes the Braden Skin Scale. The score is 13. Further assessment by the nurse reveals skin is intact with no pressure ulcers.

The following diagram shows the components of a Care Plan and the flow between them expressed using HL7 moodCodes and relationshipCodes with components of the Care Plan relationship storyboard applied to the diagram.

HAS COMPONENT HAS COMPONENT **EVALUATIONS/OUTCOMES** Health Concerns: REFERS TO Goal: Intact skin Impaired mobility At risk for impaired skin **Progress Toward** Goal: Met integrity REFERS TO REFERS TO Interventions: Requested: Turn Q 4 hours Assess skin Q 8 Quadriplegia Performed: Turned Outcome History of pressure 0600, 0800, 1200, Observation: Intact Skin 1600. HAS REASON Diabetes mellitus Skin assessment Braden scale = 13 0800, 1600

Figure 23: Care Plan Relationship Diagram - Instantiated

APPENDIX F — UNIQUE DEVICE IDENTIFICATION (UDI) ISSUING AGENCY FORMATS

Each issuing agency has its own specified format for representing the two main components of a UDI – the Device Identifier (DI) and Production Identifiers (PI). The device identifier is a mandatory, fixed portion of a UDI that identifies the specific version or model of a device and the labeler of that device. The production identifier is the conditional, variable portions of a UDI that identifies one or more of the following when included on the label of the device and include: lot or batch within which a device was manufactured; serial number of a specific device; expiration date of a specific device; date a specific device was manufactured; and for an HCT/P regulated as a device, the distinct identification code required by 21 CFR 1271.290(c). The format of each issuing agency's UDI is outlined in the tables below. These issuing agencies maintain responsibility for the uniqueness of their device identifiers.

Table 8: GS1 UDI Format16

Issuing Agency	Data Delimiters	Identifier	Data type	Human Readable Field Size	Database Field Size	
GS1	(01)	DI	numeric	16	14	
GS1	(11)	Manufacturing/ Production Date	numeric [YYMMDD]	8	6	
GS1	(17)	Expiration Date	numeric [YYMMDD]	8	6	
GS1	(10)	Batch/Lot Number	alphanumeric	22	20	
GS1	(21)	Serial Number	alphanumeric	22	20	
GS1		Maximum Base UDI	alphanumeric	76	66	
ex: (01) 51022222233336(11)141231(17)150707(10)A213B1(21)1234						

¹⁶ http://www.gs1.org/

Table 9: Health Industry Business Communications Council (HIBCC) UDI Format 17

Issuing Agency	Data Delimiters	Identifier	Data Type	Human Readable Field Size	Database Field Size
HIBCC	+	DI	alphanumeric	7 to 24	6 to 23
HIBCC	\$	Lot Number Only	alphanumeric	19	18
HIBCC	\$\$7	Lot Number Only (alternative option)	alphanumeric	21	18
HIBCC	\$\$	Expiration Date followed by Lot	Expiration Date: numeric [MMYY]	6	4
		Number	Lot Number: alphanumeric	18	18
HIBCC	\$\$2	2 Expiration Date followed by Lot	Expiration Date: numeric [MMDDYY]	9	6
		Number	Lot Number: alphanumeric	18	18
HIBCC	\$\$3	Expiration Date followed by Lot Number	Expiration Date: numeric [YYMMDD]	9	6
			Lot Number: alphanumeric	18	18
HIBCC \$\$4	\$\$4	\$\$4 Expiration Date followed by Lot Number	Expiration Date: numeric [YYMMDDHH]	11	8
			Lot Number: alphanumeric	18	18
HIBCC \$\$5	\$\$5	\$5 Expiration Date followed by Lot Number	Expiration Date: numeric [YYJJJ] – Julian Date format	8	5
			Lot Number: alphanumeric	18	18
HIBCC \$\$6	\$\$6	\$6 Expiration Date followed by Lot Number	Expiration Date: numeric [YYJJJHH] – Julian Date format with Hour option	10	7
			Lot Number: alphanumeric	18	18
HIBCC	\$+	Serial Number only	alphanumeric	20	18
HIBCC	\$\$+7	Serial Number only (alternative option)	alphanumeric	22	18
HIBCC	\$\$+	Expiration Date followed by Serial Number	Expiration Date: numeric [MMYY]	7	4
			Serial Number: alphanumeric	18	18
HIBCC	\$\$+2	Expiration Date followed by Serial	Expiration Date: numeric [MMDDYY]	10	6

¹⁷ http://www.hibcc.org/

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Issuing Agency	Data Delimiters	Identifier	Data Type	Human Readable Field Size	Database Field Size
		Number	Serial Number: alphanumeric	18	18
HIBCC	\$\$+3	Expiration Date followed by Serial	Expiration Date: numeric [YYMMDD]	10	6
		Number	Serial Number: alphanumeric	18	18
HIBCC	\$\$+4	Expiration Date followed by Serial	Expiration Date: numeric [YYMMDDHH]	12	8
		Number	Serial Number: alphanumeric	18	18
HIBCC	\$\$+5	Expiration Date followed by Serial Number	Expiration Date: numeric [YYJJJ]	9	5
			Serial Number: alphanumeric	18	18
HIBCC \$\$+6	\$\$+6	Expiration Date followed by Serial Number	Expiration Date: numeric [YYJJJHH]	11	7
			Serial Number: alphanumeric	18	18
HIBCC	/S	Supplemental Serial Number, where lot number also required and included in main secondary data string	alphanumeric	20	18
HIBCC	/16D	Manufacturing Date (supplemental to secondary barcode)	numeric [YYYYMMDD]	12	8
HIBCC		Maximum Base UDI	alphanumeric	70 to 87	58 to 75

Ex of Human Readable Barcode:

⁺ H123 PARTNO 1234567890120 / \$\$420020216 LOT 123456789012345 / SXYZ456789012345678/16D20130202C

Table 10: International Council for Commonality in Blood Banking Automation, Inc. (ICCBBA) UDI Format¹⁸

Issuing Agency	Data Delimiters	Identifier	Data type	Human Readable Barcode Field Size	Database Field Size
ICCBBA	=/	DI	alphanumeric	18	16
ICCBBA	=,	Serial Number	alphanumeric	8	6
ICCBBA	=	Donation Identification Number	alphanumeric	16	15
ICCBBA	=>	Expiration Date	numeric [YYYJJJ]	8	6
ICCBBA	=}	Manufacturing Date	numeric [YYYJJJ]	8	6
ICCBBA	&,1	MPHO Lot Number	alphanumeric	21	18
ICCBBA		Maximum Base UDI for HCT/Ps	alphanumeric	79	67

Ex of Human Readable

Table 11: ICCBBA UDI Format for Blood Bags Only

Issuing Agency	Identifying Symbol	Identifier	Data type	Eye Readable Barcode Field Size	Database Field Size	
ICCBBA	=)	DI for blood containers (bags)	alphanumeric	12	10	
ICCBBA	&)	Lot Number for blood containers (bags)	alphanumeric	12	10	
ICCBBA		Maximum Base UDI for Blood Bags	alphanumeric	24	20	
Ex of Human Readable Barcode: =)1TE123456A&)RZ12345678						

¹⁸ http://iccbba.org/