

Integrating the Healthcare Enterprise



5

**IHE Pharmacy  
Technical Framework Supplement**

10

**Pharmacy Prescription  
(PRE)**

15

**Rev. 1.7 – Trial Implementation**

20 Date: October 21, 2016

Author: IHE Pharmacy Technical Committee

Email: [pharmacy@ihe.net](mailto:pharmacy@ihe.net)

25

Please verify you have the most recent version of this document. See [here](#) for Trial Implementation and Final Text versions and [here](#) for Public Comment versions.

## Foreword

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

35 This supplement is published on October 21, 2016 for trial implementation and may be available for testing at subsequent IHE Connectathons. The supplement may be amended based on the results of testing. Following successful testing it will be incorporated into the forthcoming Pharmacy Technical Framework. Comments are invited and may be submitted at [http://www.ihe.net/Pharmacy\\_Public\\_Comments](http://www.ihe.net/Pharmacy_Public_Comments).

This supplement describes changes to the existing technical framework documents.

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

40 **Amend Section X.X by the following:**

Where the amendment adds text, make the added text **bold underline**. Where the amendment removes text, make the removed text **bold strikethrough**. When entire new sections are added, introduce with editor’s instructions to “add new text” or similar, which for readability are not bolded or underlined.

45

General information about IHE can be found at: <http://www.ihe.net>.

Information about the IHE Pharmacy domain can be found at: [http://www.ihe.net/IHE\\_Domains](http://www.ihe.net/IHE_Domains).

50

Information about the organization of IHE Technical Frameworks and Supplements and the process used to create them can be found at: [http://www.ihe.net/IHE\\_Process](http://www.ihe.net/IHE_Process) and <http://www.ihe.net/Profiles>.

The current version of the IHE Pharmacy Technical Framework can be found at: [http://www.ihe.net/Technical\\_Frameworks](http://www.ihe.net/Technical_Frameworks).

## CONTENTS

55	Introduction.....	5
	Open Issues and Questions.....	5
	Closed Issues.....	5
	<b>Volume 1 – Profiles .....</b>	<b>6</b>
60	1.n Copyright Permissions .....	6
	2.5 Dependencies of the Pharmacy Integration Profiles .....	6
	2.7 History of Annual Changes .....	6
	3 Pharmacy Prescription Content Profile.....	7
	3.1 Purpose and Scope .....	7
65	3.2 Process Flow .....	8
	3.2.1 Use Case 1: Placing a prescription .....	8
	3.3 Actors/Transactions.....	8
	3.4 Options .....	9
	3.5 Groupings .....	9
70	3.5.1 Community Pharmacy Prescription and Dispense .....	9
	3.6 Security Considerations.....	9
	3.7 Content Modules .....	10
	3.7.1 Structure of a Pharmacy Prescription Document .....	11
	Glossary .....	12
75	<b>Volume 3 – Content Modules.....</b>	<b>15</b>
	5.0 Namespaces and Vocabularies.....	15
	5.1 IHE Format Codes.....	15
	6.0 Pharmacy Content Modules.....	15
	6.3 HL7 Version 3.0 Content Modules .....	15
80	6.3.1 CDA Document Content Modules .....	15
	6.3.1.1 Pharmacy Prescription Specification 1.3.6.1.4.1.19376.1.9.1.1.1 .....	15
	6.3.2 CDA Header Content Modules .....	19
	6.3.3 CDA Section Content Modules .....	19
	6.3.3.1 Prescription Section Content Module (1.3.6.1.4.1.19376.1.9.1.2.1) .....	19
85	6.3.4 CDA Entry Content Modules .....	21
	6.3.4.1 Medicine Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.1) .....	21
	6.3.4.2 Prescription Item Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.2).....	29
	6.3.4.6 Dosage Instructions Content Module (1.3.6.1.4.1.19376.1.9.1.3.6) .....	39
	6.3.4.7 Amount of units of the consumable Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.8) .....	44
90	6.3.4.8 Substitution handling Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.9) .....	45
	6.3.4.9 Reference to Medication Treatment Plan Item Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.10) .....	47
	6.3.4.10 Reference to Prescription Item Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.11) .....	49
95	Appendices.....	52

100	Appendix A Validating CDA Documents using the Framework .....	53
	A.1 Validating Documents.....	53
	A.2 Validating Sections .....	53
	A.3 Phases of Validation and Types of Errors.....	53
	Appendix B Extensions to CDA Release 2.....	54
	B.1 IHE PHARM Extensions .....	54
	B.1.1 Used for Medicine Entry Content Module .....	54

105 **Introduction**

The Pharmacy Prescription Document Profile (PRE) describes the content and format of a prescription document generated during the process in which a health care professional (in most cases, but not necessarily always, a medical specialist or a general practitioner) decides that the patient needs medication. A prescription is an entity that can be seen as an order to anyone entitled to dispense (prepare and hand out) medication to the patient.

110 Documents created according to this profile are intended to be used in the context of the “Community Prescription and Dispense” Integration Profile (CMPD). This supplement also references other documents<sup>1</sup>. The reader should have already read and understood these documents:

- 115
1. [PHARM Common parts document](#)
  2. [PHARM Community Prescription and Dispense Integration Profile \(CMPD\)](#)
  3. [PCC Technical Framework Volume 1](#)
  4. [PCC Technical Framework Volume 2](#)
  5. [IT Infrastructure Technical Framework Volume 1](#)

120

  6. [IT Infrastructure Technical Framework Volume 2](#)
  7. [IT Infrastructure Technical Framework Volume 3](#)
  8. HL7®<sup>2</sup> and other standards documents referenced in this document

**Open Issues and Questions**

- 125
- Prescription of non-medication products": shall they be covered by this profile?
  - Prescription Section Content Module: It is still in discussion, if it's allowed to state the CCD®<sup>3</sup> template as "parent", or if we have to weaken it to "derived from".

**Closed Issues**

- 130
- Substitution Handling: Evaluation if incorporating parts of the HL7 COCT\_RM360000UV structure to provide a semantically better solution than now. -> The structure has been corrected in accordance with HL7 (see CP-PHARM-005\_v2)

---

<sup>1</sup> The first seven documents can be located on the IHE Website at [http://ihe.net/Resources/Technical\\_Frameworks/](http://ihe.net/Resources/Technical_Frameworks/). The remaining documents can be obtained from their respective publishers.

<sup>2</sup> HL7 is the registered trademark of Health Level Seven International.

<sup>3</sup> CCD is the registered trademark of Health Level Seven International.

# Volume 1 – Profiles

*Add the following to Section 1.n*

## 1.n Copyright Permissions

135 Health Level Seven, Inc. has granted permission to the IHE to reproduce tables from the HL7 standard. The HL7 tables in this document are copyrighted by Health Level Seven, Inc. All rights reserved. Material drawn from these documents is credited where used.

*Add the following to Section 2.5*

## 2.5 Dependencies of the Pharmacy Integration Profiles

Pharmacy Prescription (PRE)	PCC	Content definition	This profile includes Section and Entry Content Modules of the Patient Care Coordination (PCC) domain.
--------------------------------	-----	--------------------	--

140

*Add the following to Section 2.7*

## 2.7 History of Annual Changes

*Add Section 3*

145 **3 Pharmacy Prescription Content Profile**

The Pharmacy Prescription Document Profile (PRE) describes the content and format of a prescription document generated during the process in which a health care professional (in most cases, but not necessarily always, a medical specialist or a general practitioner) decides that the patient needs medication. A prescription is an entity that can be seen as an order to anyone entitled to dispense (prepare and hand out) medication to the patient.

150

Documents created according to this profile are intended to be used in the context of the “Community Prescription and Dispense” Integration Profile (CMPD).

### **3.1 Purpose and Scope**

155

The Community Pharmacy Prescription and Dispense workflow starts with the creation of a prescription in case the health care professional decides that the patient needs medication.

160

A prescription document is issued by one ordering healthcare professional for one patient, in the context of zero or one administrative encounter (between the patient and the ordering physician and/or the healthcare institution). A prescription may contain one or more Prescription Items (lines on a paper prescription). Each line relates to one medication. A prescription is the outcome of a clinical decision.

This profile defines the content and format of such a prescription document.

For a detailed overview on the whole Pharmacy domain business processes, please refer to the “Common parts” document, which is accompanying this profile.

---

165    **3.2 Process Flow**

**3.2.1 Use Case 1: Placing a prescription**

During treatment of a patient, physicians or other allowed persons may have to prescribe drugs for the patient. The prescription shall contain one or more positions (Prescription Items) which contain the drug identified by a medication identifier, the dosing as well as other information necessary for correct dispensing and administering by the patient.

Usually the physician uses the prescribing module in the physician information system for preparing the prescription. After the prescription is completely assembled it shall be submitted to the Community Pharmacy Prescription and Dispense system to be validated and dispensed.

Refer to the Community Pharmacy Prescription and Dispense Integration Profile (CMPD) for detailed use case information.

**3.3 Actors/Transactions**

There are two actors in this profile, the Content Creator and the Content Consumer. Content is created by a Content Creator and is to be consumed by a Content Consumer. The sharing or transmission of content from one actor to the other is addressed by the appropriate use of IHE profiles described below, and is out of scope of this profile. A Document Source or a Portable Media Creator may embody the Content Creator. A Document Consumer, a Document Recipient or a Portable Media Importer may embody the Content Consumer. The sharing or transmission of content or updates from one actor to the other is addressed by the use of appropriate IHE profiles described in the section on Content Bindings with XDS, XDM and XDR in PCC TF-2:4.1



**Figure 3.3-1: Actor Diagram**

190

### 3.4 Options

Options that may be selected for this content profile are listed in Table 3.4-1 along with the actors to which they apply. Dependencies between options when applicable are specified in notes.

**Table 3.4-1: Pharmacy Prescription Actors and Options**

Actor	Option	Section
Content Consumer	View Option (See Note 1) Document Import Option (See Note 1) Section Import Option (See Note 1) Discrete Data Import Option (See Note 1)	PCC TF-2: 3.1.1 PCC TF-2: 3.1.2 PCC TF-2: 3.1.3 PCC TF-2: 3.1.4
Content Creator	No options defined	

Note 1: The actor shall support at least one of these options.

### 200 3.5 Groupings

Content profiles may impose additional requirements on the transactions used when grouped with actors from other IHE profiles.

#### 3.5.1 Community Pharmacy Prescription and Dispense

Actors from the Pharmacy CMPD Profile embody the Content Creator and Content Consumer sharing function of this profile. A Content Creator or Content Consumer may be grouped with appropriate actors from the XDS, XDM or XDR Profiles to exchange the content described therein. The metadata sent in the document sharing or interchange messages has specific relationships or dependencies (which we call bindings) to the content of the clinical document described in the content profile.

210 The Patient Care Coordination Technical Framework defines the bindings to use when grouping the Content Creator of this profile with actors from the Pharmacy CMPD Integration Profiles.

### 3.6 Security Considerations

The PRE Integration Profile assumes that a minimum security and privacy environment has been established across all participants. There must exist security policies regarding the use of training, agreements, risk management, business continuity and network security that need to be already in place prior to the implementation of PRE.

The IHE ITI ATNA Integration Profile is required of the actors involved in the IHE transactions specified in this profile to protect node-to-node communication and to produce an audit trail of the PHI related actions when they exchange messages.

220 In addition, the IHE ITI DSG Integration Profiles can be applied to the actors involved in the transactions specified in this profile to securely identify individuals involved in transactions and verify document integrity and authorizations (DSG).

Interested parties should also read the detailed Security Considerations sections provided for each of the aforementioned profiles in the IHE ITI Technical Framework and its supplements.

- 225     The PRE Profile does have a few security considerations of its own.  
Pharmacy systems should be thoughtfully designed so that providers are able to review and verify information before it is imported into their Pharmacy system, and that positive user acknowledgements are made before import, and audit trails are recorded when imports occur.
- 230     Imported information should be traceable both to the source Pharmacy system, and the receiver that accepted it into the Pharmacy system. XDS Affinity domain policies should support policies and procedures for tracing information flows between Pharmacy systems.
- 235     Because the information being transferred is in XML, it will be common that different Pharmacy systems utilize different transformations to render the contents into human readable form. A Content Creator should make available the transforms used by the sending provider to review the documents, and a Content Consumer must support rendering the information as seen by the sending provider, allowing both providers to see what was sent in its original rendered form.

### **3.7 Content Modules**

- Content modules describe the content of a payload found in an IHE transaction. Content profiles are transaction neutral. They do not have dependencies upon the transaction that they appear in. These dependencies are reflected in the Bindings listed above.
- 240     All Pharmacy Prescriptions shall be structured and coded as required by the Pharmacy Prescription Document Content Module described in this profile. The inclusion of the specific coded attributes explicitly defined as optional, may be supported by specific implementations of Document Sources using an IHE identified coded terminology of their choice. The requirements and manner in which implementations support such capabilities is beyond the scope of this integration profile.

### 3.7.1 Structure of a Pharmacy Prescription Document

**Pharmacy Prescription CDA Document**  
*Pharmacy Prescription Content Module*  
(1.3.6.1.4.1.19376.1.9.1.1.1)

1..1

**Prescription**  
*Prescription Section Content Module*  
1.3.6.1.4.1.19376.1.9.1.2.1

1..n

**Prescription Item**  
*Prescription Item Entry Content Module*  
1.3.6.1.4.1.19376.1.9.1.3.2

:



## Glossary

255 **Add the following terms to the Glossary:**

### **Dispense/Dispensation**

Dispensation is the act of assigning a medication to a patient, normally as indicated in the corresponding prescription. Since prescriptions can span long periods of time, a single prescription may result in medicines dispensed several times.

260 **Dispense Item**

A Dispense Item belongs to one Dispensation and represents one dispensed medication. It contains the dispensed medicinal product including information such as product code, brand name and packaging information.

### **Dosage Instructions**

265 Dosage Instructions are a set of data elements which together represent the dosage instructions to a medication such as duration of treatment, medication frequency, dose quantity and route of administration.

### **ICA**

270 ICAs are Intolerances, Contra-indications and Allergies. An ICA may be considered as a relationship between a Patient and a Medicine. A detected problem in a Pharmaceutical Advice may refer to an ICA.

### **Medication Dispenser**

275 In the domain of community pharmacy a Medication Dispenser is an abstract actor which dispenses prescribed medication to a patient (generally a healthcare professional, usually a pharmacist when the patient enters the pharmacy to get the prescribed medication).

### **Medication**

A medication is part of a Prescription Item and defines the actual prescribed drug. It contains the brand or generic name of the drug, national and/or regional drug codes, unit strength, active ingredients and packaging information.

280 **Medication Brand Name**

The brand name is the name given to a medicine by the pharmaceutical company that makes it. This is also called the "proprietary name".

### **Medication Generic/Scientific Name**

285 The generic or scientific name is the term given to the active ingredient in the medicine that is decided by an expert committee and is understood internationally. This is also called the "non-proprietary name".

### **Medication Treatment Plan**

290 The Medication Treatment Plan (MTP) of a patient is the collection of all medications the patient was planned to take in the past, presently or in the future. The Medication Treatment Plan is the complete set of all Medication Treatment Plan Items of the patient, i.e., not partitioned or grouped by pathology, planner, organization, etc.

### **Medication Treatment Plan Item**

295 A Medication Treatment Plan Item is a single medication the patient was planned to take in the past or is planned to take presently or in the future, including its name, dosage, frequency of intake, etc. as well as other information such as patient- and fulfillment instructions and substitution handling. A Medication Treatment Plan Item triggers prescriptions and/or, dispenses in order to fulfill the medication treatment planned by the item.

### **Pharmaceutical Adviser**

300 A Pharmaceutical Adviser is an abstract actor which validates Prescription Items issued on a prescription (generally a healthcare professional, usually a pharmacist when the patient enters the pharmacy to get the prescribed medication).

### **Pharmaceutical Advice**

305 A Pharmaceutical Advice document is the outcome of the validation or review of one Prescription- or Dispense Item. It contains the overall result of the validation or review which affects the further processing as well as additional information such as Intolerances, Contraindications and Allergies (ICAs) and all other information which was discovered during validation.

310 A Pharmaceutical Advice document is also used to manage Prescription- or Dispensation Items (e.g., change, cancel, etc.) as well as to document Medication Interaction Checking Issues and their resolutions.

### **Pharmaceutical Advice Item**

315 A Pharmaceutical Advice Item belongs to one Pharmaceutical Advice and represents the validation outcome or management command regarding the referenced Prescription- or Dispense Item (e.g., change, cancel, etc.). It may also carry Medication Interaction Checking Issue information regarding the referenced item.

### **Pharmaceutical Advice Concern Item**

A Pharmaceutical Advice Concern Item belongs to one Pharmaceutical Advice Item and represents the information to concerns (e.g., problems, allergies, etc.) which the Prescription- or Dispense Item referenced by the underlying Pharmaceutical Advice Item causes.

### **Pharmacy Medication list**

A Pharmacy Medication list is a collection of Prescription- and Dispense items (and their related Pharmaceutical Advice Items) representing the Medication information of the patient at a certain point of time and according to business rules specified.

### **Prescriber**

- 325 A prescriber is an abstract actor who issues a prescription to a patient (generally a healthcare professional, usually a physician during treatment of a patient).

### **Prescription**

- 330 A prescription is issued by one ordering healthcare professional for one patient, in the context of zero or one administrative encounter (between the patient and the ordering physician and/or the healthcare institution).

### **Prescription Item**

- 335 A Prescription Item belongs to one prescription and represents one prescribed medication. It may be associated with one or more observations. Prescription Item is the atomic entity for logistics, distribution and billing. It contains the prescribed medicine and dosage information as well as other information to the prescribed item such as patient- and fulfillment instructions and substitution handling.

# Volume 3 – Content Modules

## 5.0 Namespaces and Vocabularies

codeSystem	codeSystemName	Description
1.3.6.1.4.1.19376.1.9	IHE Pharmacy Object Identifiers	This is the root OID for all IHE Pharmacy objects
1.3.6.1.4.1.19376.1.5.3.4		Namespace OID used for IHE Extensions to CDA® <sup>4</sup> Release 2.0

- 340 See also the Namespaces and Vocabularies of the IHE PCC Technical Framework ([PCC-TF2/Namespaces and Vocabularies](#)).

## 5.1 IHE Format Codes

Profile	Format Code	Media Type	Template ID
<b>2010 Profiles</b>			
Pharmacy Prescription (PRE)	urn:ihe:pharm:pre:2010	text/xml	1.3.6.1.4.1.19376.1.9.1.1.1

- 345 **6.0 Pharmacy Content Modules**

## 6.3 HL7 Version 3.0 Content Modules

### 6.3.1 CDA Document Content Modules

*Add Section 6.3.1.1*

#### 6.3.1.1 Pharmacy Prescription Specification 1.3.6.1.4.1.19376.1.9.1.1.1

- 350 The Pharmacy Prescription specification includes a Prescription section to capture Prescription Items prescribed to a patient as well as supporting sections containing information related to this prescription (e.g., diagnosis, etc.).

Structure	Pharmacy Prescription
Format Code	urn:ihe:pharm:pre:2010
LOINC Code	57833-6 (Prescriptions)

<sup>4</sup> CDA is the registered trademark of Health Level Seven International.

Structure	Pharmacy Prescription
Document Template ID	1.3.6.1.4.1.19376.1.9.1.1.1
Section name / template ID	Prescription 1.3.6.1.4.1.19376.1.9.1.2.1
Entry name / template ID	Prescription Item 1.3.6.1.4.1.19376.1.9.1.3.2
Medicine Content Entry Module template ID	Medication of Prescription Item 1.3.6.1.4.1.19376.1.9.1.3.1

### 6.3.1.1.1 Format Code

355 The XDSDocumentEntry format code for this content is **urn:ihe:pharm:pre:2010**.

### 6.3.1.1.2 Parent Template

This document is an instance of the [Medical Document](#) template.

### 6.3.1.1.3 Standards

<b>HL7V3 NE2009</b>	<a href="#">HL7 V3 2009 Normative Edition</a>
<b>CDAR2</b>	<a href="#">HL7 CDA Release 2.0</a>
<b>CCD</b>	<a href="#">ASTM/HL7 Continuity of Care Document</a>
<b>XMLXSL</b>	<a href="#">Associating Style Sheets with XML documents</a>

### 6.3.1.1.4 Data Element Index

Data Elements	CDA Release 2.0
<b>Patient Information</b>	recordTarget/patientRole
Patient Administrative Identifiers	recordTarget/patientRole/id
Patient Name	recordTarget/patientRole/patient/name
Patient Gender	recordTarget/patientRole/patient/administrativeGenderCode
Patient Birth Date	recordTarget/patientRole/patient/birthTime
Patient Address	recordTarget/patientRole/addr
Patient Telecom	recordTarget/patientRole/telecom
<b>HCP Person Information</b>	author
HCP ID(s)	author/assignedAuthor/id
HCP Profession	author/functionCode
HCP Name	author/assignedAuthor/assignedPerson/name
HCP Telecom	author/assignedAuthor/telecom
HCP Specialty	author/assignedAuthor/code
<b>HCP Represented Organization</b>	author/assignedAuthor/representedOrganization
HCP Organization Name	author/assignedAuthor/representedOrganization/name
HCP Organization Address	author/assignedAuthor/representedOrganization/addr

Data Elements	CDA Release 2.0
HCP Organization Telecom	author/assignedAuthor/representedOrganization/telecom
<b>Service Event</b> <sup>5</sup>	documentationOf/serviceEvent
Date of Service Event	documentationOf/serviceEvent/effectiveTime
Service Event Code	documentationOf/serviceEvent/code
<b>Encounter in the healthcare institution</b> <sup>6</sup>	componentOf/encompassingEncounter
ID of the encounter	componentOf/encompassingEncounter/id
Date of Admission/Encounter start date	componentOf/encompassingEncounter/effectiveTime/low
Date of Discharge/Encounter end date	componentOf/encompassingEncounter/effectiveTime/high
Authorization	authorization/consent
Patient contacts	guardian
Payers	PAYMENT SOURCES
General Medical Information Height, Weight	VITAL SIGNS
Allergies and Drug Sensitivities	ALLERGIES, ADVERSE REACTIONS, ALERTS
Active Problems	PROBLEM LIST
Resolved Problems	HISTORY OF PAST ILLNESS
Immunizations	HISTORY OF IMMUNIZATIONS
Pregnancy History	HISTORY OF PREGNANCIES
Prescription	PRESCRIPTIONS

360

### 6.3.1.1.5 Data Element Specification

Data Element Name	Opt	Template ID
Patient Information Name Personal Identification Gender Date of Birth	R	1.3.6.1.4.1.19376.1.5.3.1.1.1
HCP Person Information Name HCP Identification		
HCP Organization Information Name Address Organization Identifier		

<sup>5</sup> Service Event is optional and may contain service event information of the medical event in which context the prescription has been taken.

<sup>6</sup> Encounter is optional and shall contain encounter information if applicable.

Data Element Name	Opt	Template ID
Contact Information		
<u>Patient Information</u> Address Contact Information	R2	1.3.6.1.4.1.19376.1.5.3.1.1.1
<u>HCP Person Information</u> Profession Specialty		
<u>Patient Information</u> Marital Status Race Ethnicity Religious Affiliation	O	1.3.6.1.4.1.19376.1.5.3.1.1.1
<u>HCP Person Information</u> Contact Information		
Authorization	R2	1.3.6.1.4.1.19376.1.5.3.1.2.5
Patient Contacts	O <sup>7</sup>	1.3.6.1.4.1.19376.1.5.3.1.2.4
Payers	R2	1.3.6.1.4.1.19376.1.5.3.1.1.5.3.7
Coded Vital Signs	O <sup>8</sup>	1.3.6.1.4.1.19376.1.5.3.1.1.5.3.2
Allergies and Other Adverse Reactions	O	1.3.6.1.4.1.19376.1.5.3.1.3.13
Active Problems	O	1.3.6.1.4.1.19376.1.5.3.1.3.6
History of Past Illness	O	1.3.6.1.4.1.19376.1.5.3.1.3.8
Immunizations	O	1.3.6.1.4.1.19376.1.5.3.1.3.23
Pregnancy History	O <sup>9</sup>	1.3.6.1.4.1.19376.1.5.3.1.1.5.3.4
<u>Prescription</u>	R	1.3.6.1.4.1.19376.1.9.1.2.1

Additional explanation:

365 The sections “Coded Vital Signs”, “Allergies and Drug Sensitivities”, “Active Problems”, “Resolved Problems”, “Immunizations”, “Pregnancy History” are considered as sections containing medical information of the patient.

Although real-world projects may require some of these information, no stricter constraints as optional (O) could been applied to these sections in the profile due to the large degree of diversity in business requirements and privacy issues among different current.

---

<sup>7</sup> In case the patient is governed by a guardian, this element is R and shall contain the information about the guardian

<sup>8</sup> The Coded Vital Signs section should contain at least the height and weight of the patient.

<sup>9</sup> In case the patient is currently pregnant, this element is R and shall contain information about the current pregnancy. It shall not be used to document past pregnancies.

370 **6.3.1.1.6 Conformance**

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

```

380 <ClinicalDocument xmlns='urn:hl7-org:v3'>
<typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
385 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.1' />
<templateId root='1.3.6.1.4.1.19376.1.9.1.1.1' />
<id root=' ' extension=' '/>
<code code='57833-6' displayName='Prescription for medication'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<title>Pharmacy Prescription</title>
<effectiveTime value='20100719012005' />
<confidentialityCode code='N' displayName='Normal'
      codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality' />
<languageCode code='en-US' />
390   :
<component>
  <structuredBody>
    :
    </structuredBody>
  </component>
</ClinicalDocument>
395

```

**6.3.2 CDA Header Content Modules****6.3.3 CDA Section Content Modules**400 *Add Section 6.3.3.1***6.3.3.1 Prescription Section Content Module (1.3.6.1.4.1.19376.1.9.1.2.1)**

<b>Template ID</b>	1.3.6.1.4.1.19376.1.9.1.2.1	
<b>Parent Template</b>	CCD 3.9 (2.16.840.1.113883.10.20.1.8) 1.3.6.1.4.1.19376.1.5.3.1.3.19	
<b>General Description</b>	The Prescription Section contains a description of the medications in a given prescription for the patient. It includes entries for Prescription Items as described in the Prescription Item Entry Content Module.	
<b>LOINC Code</b>	<b>Opt</b>	<b>Description</b>
57828-6	R	PRESCRIPTIONS
<b>Entries</b>	<b>Opt</b>	<b>Description</b>
1.3.6.1.4.1.19376.1.9.1.3.2	R	Prescription Item Entry Content Module

```

405 <component>
<section>
<templateId root='2.16.840.1.113883.10.20.1.8' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.19' />
<templateId root='1.3.6.1.4.1.19376.1.9.1.2.1' />
410 <!-- The section ID is the Prescription ID -->
<id root=' ' extension=' ' />
<code code='57828-6' displayName='PRESCRIPTIONS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<title>Prescriptions</title>
415 <text>
      Text as described above
</text>
<author>
  :
</author>
420 <!-- Each entry is a Prescription Item -->
<!-- Prescription Item 1 -->
<entry>
  :
425 <!-- Required element indicating the prescription entry content module -->
  <templateId root='1.3.6.1.4.1.19376.1.9.1.3.2' />
  :
</entry>
<!-- Prescription Item 2 -->
430 <entry>
  :
<!-- Required element indicating the prescription entry content module -->
  <templateId root='1.3.6.1.4.1.19376.1.9.1.3.2' />
  :
</entry>
435 </section>
</component>

```

### 6.3.3.1.1 Parent Templates

The parents of this template are CCD 3.9 and PCC 1.3.6.1.4.1.19376.1.5.3.1.3.19 except the requirement CCD-CONF-301 (“The value for ‘section/code’ SHALL be “10160-0” “History of medication use”).

### 6.3.3.1.2 Prescription ID

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

A Prescription identifier SHALL be represented in the section **<id>** Element. The data type of the ID is II. Although HL7 allows for multiple identifiers, one and only one shall be used.

445

If this section is used in a Prescription document, this identifier shall be set equal to the document ID (ClinicalDocument/id).

### 6.3.3.1.3 Prescriber

450 <author>...</author>

In the case where the prescriber or the timestamp of a prescription is different from the author and timestamp of the prescription-document, the prescriber and timestamp of the prescription shall be represented by the <author> element of the section.

Data element	HL7 V3 Data Type	CDA Body position (relative XPath expression)
Prescriber Profession	CE	<i>author/functionCode</i>
Timestamp of prescribing	TS	<i>author/time</i>
Prescriber ID	II	<i>author/assignedAuthor/id</i>
Prescriber Specialty	CE	<i>author/assignedAuthor/code</i>
Prescriber Name	PN	<i>author/assignedAuthor/assignedPerson/name</i>
Prescriber Organization Identifier	II	<i>author/assignedAuthor/representedOrganization/id</i>
Prescriber Organization Name	ON	<i>author/assignedAuthor/representedOrganization/name</i>
Prescriber Organization Address	AD	<i>author/assignedAuthor/representedOrganization/addr</i>

455

### 6.3.4 CDA Entry Content Modules

Add Section 6.3.4.1

#### 6.3.4.1 Medicine Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.1)

460 A Medicine entry describes a medicine and is used within Prescription- or Dispensation Items. It describes either a medicinal product, a generic/scientific name or a magistral preparation/compound medicine and contains information such as name, medication form, packaging information and active ingredients.

465 This entry uses the structure of the HL7 V3 R\_Medication Universal Common Message Element (CMET), Release 2.

This structure is part of the HL7 V3 2009 Normative Edition (COCT\_RM230100UV). The incorporation of this structure is done according to section **1.4 CDA Extensibility** of the HL7 CDA standard. Such an extension of the base CDA standard is an accepted practice in IHE (e.g., in the XD\* Lab specification).

470 For the purposes of IHE Pharmacy this extension is necessary to satisfy the requirements for Prescription, Pharmaceutical Advice and Dispense data elements to represent required information which is not supported by the base CDA standard, such as packaging information, generic equivalent and ingredients.

475 The rules of section **1.4 CDA Extensibility** require the designation of a new XML namespace for the XML elements in this structure. For the purposes of documentation, the **namespace *urn:ihe:pharm:medication*** shall be used.

The following specification and constraints are applied to the structures of the CMET.

#### **6.3.4.1.1 Standards**

480 This part describes the general structure for a Prescription Item. It is based on the following standards:

<b>HL7V3 NE2009</b>	<a href="#">HL7 V3 2009 Normative Edition</a>
-------------------------	---

#### **6.3.4.1.2 Parent Template**

This entry content module has no parent structure.

485

### 6.3.4.1.3 Specification

```

<manufacturedProduct xmlns:pharm="urn:ihe:pharm:medication" classCode="MANU">
    <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.7.2"/>
    <templateId root="2.16.840.1.113883.10.20.1.53"/>
    <manufacturedMaterial classCode="MMAT" determinerCode="KIND">
        <templateId root="1.3.6.1.4.1.19376.1.9.1.3.1"/>
        <!-- National medicinal product code (brand-level) -->
        <code code="" displayName="" codeSystem="" codeSystemName="" />
        <!-- Brand name -->
        <name>... </name>
        <!-- Pharmaceutical dose form -->
        <pharm:formCode code="" displayName="" codeSystem="" codeSystemName="" />
        <lotNumberText>...</lotNumberText>
        <pharm:expirationTime value=' '/>
        <!-- Container information -->
        <pharm:asContent classCode="CONT">
            <pharm:containerPackagedMedicine classCode="CONT" determinerCode="INSTANCE">
                <!-- Medicinal product code (package-level) -->
                <pharm:code code="" displayName="" codeSystem="" codeSystemName="" />
                <!-- Brand name (package) -->
                <pharm:name>...</pharm:name>
                <pharm:formCode code="" displayName="" codeSystem="" codeSystemName="" />
                <pharm:capacityQuantity value="" unit="" />
                <pharm:asSuperContent>
                    <pharm:containerPackagedMedicine classCode='CONT'
                        determinerCode='INSTANCE'>
                        <pharm:capacityQuantity value=' ' unit=' '/>
                    </pharm:containerPackagedMedicine>
                </pharm:asSuperContent>
                </pharm:containerPackagedMedicine>
            </pharm:asContent>
            <!-- These are optional generic equivalents -->
            <pharm:asSpecializedKind classCode="GRIC">
                <pharm:generalizedMedicineClass classCode="MMAT">
                    <pharm:code code="" displayName="Generic Equivalent"
                        codeSystem="" codeSystemName="" />
                    <pharm:name>...</pharm:name>
                </pharm:generalizedMedicineClass>
            </pharm:asSpecializedKind>
            <!-- This is the list of active ingredients -->
            <pharm:ingredient classCode="ACTI">
                <!-- strength of ingredient -->
                <pharm:quantity>
                    <numerator xsi:type="PQ" value="" unit="" />
                    <denominator xsi:type="PQ" value="" unit="" />
                </pharm:quantity>
                <pharm:ingredient classCode="MMAT" determinerCode="KIND">
                    <pharm:code code="" displayName="Active Ingredient 1"
                        codeSystem="" codeSystemName="" />
                    <pharm:name>Active Ingredient 1</pharm:name>
                </pharm:ingredient>
            </pharm:ingredient>
            <pharm:ingredient classCode="ACTI">
                <!-- strength of ingredient -->
                <pharm:quantity>
                    <numerator xsi:type="PQ" value="" unit="" />
                    <denominator xsi:type="PQ" value="" unit="" />
                </pharm:quantity>
                <pharm:ingredient classCode="MMAT" determinerCode="KIND">
                    <pharm:code code="" displayName="Active Ingredient 2"
                        codeSystem="" codeSystemName="" />
                </pharm:ingredient>
            </pharm:ingredient>
        </pharm:ingredient>
    </manufacturedMaterial>
</manufacturedProduct>

```

550           

```
<pharm:name>Active Ingredient 2</pharm:name>
  </pharm:ingredient>
</pharm:ingredient>
</manufacturedMaterial>
</manufacturedProduct>
```

555   **6.3.4.1.3.1 Medicine Entry General Specification**

```
<manufacturedProduct xmlns:pharm="urn:ihe:pharm:medication"
  classCode="MANU">
  <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.7.2"/>
  <templateId root="2.16.840.1.113883.10.20.1.53"/>
  <manufacturedMaterial classCode="MMAT" determinerCode="KIND">
    ...
  </manufacturedMaterial>
</manufacturedProduct>
```

565   The `<manufacturedProduct>` element of the Medicine Entry SHALL contain a XML namespace “pharm” having the value “urn:ihe:pharm:medication”.

See PCC TF2, Product Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7.2) specification.

**6.3.4.1.3.2 Medicine Entry Template ID**

```
<templateId root='1.3.6.1.4.1.19376.1.9.1.3.1'/>
```

The Template ID for a Medicine Entry SHALL be provided.

570   **6.3.4.1.3.3 Code**

```
<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '>
  <originalText>
    <reference value=' '/>
  </originalText>
</code>
```

575   The `<code>` element of the `<manufacturedMaterial>` element SHALL be present and describes the code of the medication.

The medication may be either

- a brand/product or
- described as a generic/scientific name or
- a descriptor of a magistral preparation/compound medicine

The <originalText> shall contain a <reference> whose URI value points to the name and strength of the medication, or just the name alone if strength is not relevant.

585 If the medicine is uncoded (e.g., magistral preparations, compound medicine, ...) nullFlavor="NA" SHALL be used.

#### **6.3.4.1.3.4 Name**

< name>...</ name>

The element SHALL contain the name of the medication (e.g., "Adol 500mg Caplet").

The medication may be either

- 590
- a brand/product or
  - described as a generic/scientific name or
  - a descriptor of a magistral preparation/compound medicine

If the medicine has no brand name (e.g., magistral preparations, compound medicine, ...) nullFlavor="NA" SHALL be used.

595 **6.3.4.1.3.5 Form Code**

<pharm:formCode code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>

This code represents the pharmaceutical dose form (e.g., tablet, capsule, liquid) and SHOULD be present, if not implied by the product. It MAY be present if implied by the product. The value of this code affects the units used in the substance administration quantity element.

600 **6.3.4.1.3.6 Lot Number**

<lotNumberText>...</lotNumberText>

The <lotNumberText> element MAY be present and is a string representation of a lot number of this specific instance of the product.

605 The provided lot number SHALL refer to the primary packaged item described in the Packaging element.

#### **6.3.4.1.3.7 Expiration Date**

<pharm:expirationTime value=' '/>

610 The <pharm:expirationTime> element MAY be present and SHALL contain a value attribute containing the date (e.g., specific date, specific date including time) of expiration of this specific instance of the product.

The value given in the <pharm:expirationTime> element SHALL refer to the primary packaged item described in the Medicine Packaging element.

### 6.3.4.1.3.8 Packaging

```

<pharm:asContent classCode='CONT'>
  <pharm:containerPackagedMedicine classCode='CONT'
    determinerCode='INSTANCE'>
    <!-- Medicinal product code (package-level) -->
    <pharm:code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>
    <!-- Brand name (package) -->
    <pharm:name>...</pharm:name>
    <pharm:formCode code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>
    <pharm:capacityQuantity value=' ' unit=' '/>
    <pharm:asSuperContent>
      <pharm:containerPackagedMedicine classCode='CONT'
        determinerCode='INSTANCE'>
        <pharm:capacityQuantity value=' ' unit=' '/>
      </pharm:containerPackagedMedicine>
    </pharm:asSuperContent>
  </pharm:containerPackagedMedicine>
</pharm:asContent>

```

This structure describes the packaging of the medication and MAY be present. It represents the primary description of the packaging of the medicine (e.g., the medicine is packaged in ampoules of 50ml volume each) and may include additional packaging information of how many of the primary packaged items are within an outer package (e.g., 5 ampoules are packaged in a box).

The primary description of the package SHOULD be consistent with the given pharmaceutical dose form (<pharm:formCode> of the medication, see chapter “Medication Form Code”).  
Example: a consistent pharmaceutical dose form to the package form “Ampoules” would be e.g., “Solution for injection”.

In case the package describes a product, the <pharm:code> element provides the code for the product and SHOULD be present.

In case the package describes a product, and the package has a brand name, it SHOULD be described in the <pharm:name> element (e.g., Xylocaine 1% with Adrenaline Inj, 5 injections package).

The <pharm:formCode> element represents the form of the package (e.g., tablet container, bottle, ...) and SHOULD be present, if not implied by the product. It MAY be present if implied by the product. It SHALL be present if the <asSuperContent> element is present.

The <pharm:capacityQuantity> element SHALL be present and describes the capacity of the packaging.

If the capacityQuantity is given in countable units, the unit attribute SHALL NOT be present. If the capacityQuantity is given in non-countable units, the unit attribute SHALL be present and the value SHALL be out of the UCUM code system.

655 A <pharm:asSuperContent> element MAY be present to represent the quantity of the given primary packaged item (e.g., ampoule of 50ml) within an outer package (e.g., 5 ampoules in a box). It SHALL contain a <pharm:containerPackagedMedicine> element which SHALL contain a <pharm:capacityQuantity> element describing the quantity of the given primary packaged item present in the outer package. If the capacityQuantity is given in countable units, the unit attribute SHALL NOT be present. If the capacityQuantity is given in non-countable units, the unit attribute SHALL be present and the value SHALL be out of the UCUM code system.

Examples:

660 For example, to represent a medicinal product with pharmaceutical dose form "Tablets", available as a "Tablet container" with 30 tablets the <asContent> data elements would be set to:

```
<pharm:asContent classCode='CONT'>
    <pharm:containerPackagedMedicine classCode='CONT'
        determinerCode='INSTANCE'>
        :
        <pharm:formCode code=' ' displayName='Tablet container' codeSystem=' '
            codeSystemName=' '/>
        <pharm:capacityQuantity value='30' /> <!-- 30 tablets in the package -->
        </pharm:containerPackagedMedicine>
    </pharm:asContent>
```

665 670 For example, to represent a medicinal product with pharmaceutical dose form "Solution for injection", available as "Ampoules" with 50ml volume, packaged as 5 ampoules per box, the <asContent> data elements would be set to:

```
<pharm:asContent classCode='CONT'>
    <pharm:containerPackagedMedicine classCode='CONT'
        determinerCode='INSTANCE'>
        :
        <pharm:formCode code=' ' displayName='Ampoules' codeSystem=' '
            codeSystemName=' '/>
    <pharm:capacityQuantity value='50' unit='ml' /> <!-- 50ml per ampoule -->
    <pharm:asSuperContent>
        <pharm:containerPackagedMedicine classCode='CONT'
            determinerCode='INSTANCE'>
            <pharm:capacityQuantity value='5' /> <!-- 5 ampoules in a box -->
        </pharm:containerPackagedMedicine>
    </pharm:asSuperContent>
</pharm:containerPackagedMedicine>
</pharm:asContent>
```

690 Note: The "Ampoule" can be seen as a container representing a volume and as the "unit of use"/"unit of presentation". In this example it only represents container packaging.

### 6.3.4.1.3.9 Generic Equivalent

695    <pharm:asSpecializedKind classCode='GRIC'>  
      <pharm:generalizedMedicineClass classCode='MMAT'>  
         <pharm:code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>  
         <pharm:name>...</pharm:name>  
      </pharm:generalizedMedicineClass>  
   </pharm:asSpecializedKind>

700    The classCode of "GRIC" identifies this structure as the representation of a generic equivalent of  
      the medication described in the current Medicine entry.

One or more elements MAY be present.

The <pharm:code> element contains the coded representation of the generic medicine, and the  
<pharm:name> element may be used for the plain text representation.

### 6.3.4.1.3.10 Active Ingredient List

705    <pharm:ingredient classCode='ACTI'>  
      <pharm:quantity>  
         <nominator xsi:type='PQ' value=' ' unit=' '/>  
         <denominator xsi:type='PQ' value=' ' unit=' '/>  
      </pharm:quantity>  
    <pharm:ingredient classCode='MMAT' determinerCode='KIND'>  
      <pharm:code code=' ' displayName=' '  
             codeSystem='2.16.840.1.113883.6.73' codeSystemName='ATC WHO'>  
      <pharm:name>...</pharm:name>  
    </pharm:ingredient>  
  </pharm:ingredient>

715    One or more active ingredients SHOULD be represented with this structure. The classCode of  
      "ACTI" indicates that this is an active ingredient.

720    The <pharm:code> element SHOULD be present and contains the coded representation of the  
      active ingredient. The <pharm:name> element SHALL be present and is used for the plain text  
      representation. The WHO ATC terminology SHOULD be used to code the active ingredients,  
      where applicable.

725    The medication strength is represented as the ratio of the active ingredient(s) to a unit of  
      medication. The <pharm:quantity> element SHOULD be present and represents the strength of  
      the active ingredient(s) as the ratio of the active ingredient(s) to a unit of medication. The  
      <pharm:quantity> element contains the numerator and denominator of the strength ratio.

The following example shows the strength of 10 mg of the ingredient per ml of the medication:

730

```
<pharm:quantity>
  <numerator xsi:type="PQ" value="10" unit="mg"/>
  <denominator xsi:type="PQ" value="1" unit="ml"/>
</pharm:quantity>
```

The following examples show the strength of the ingredient in 1 unit of the given medication.

2% of the ingredient (e.g., XYLOPHIL 2% GEL):

735

```
<pharm:quantity>
  <numerator xsi:type="PQ" value="2" unit="%"/>
  <denominator xsi:type="PQ" value="1"/>
</pharm:quantity>
```

740

5mg of the ingredient (e.g., XYZAL 5MG F.C.TABLET):

```
<pharm:quantity>
  <numerator xsi:type="PQ" value="5" unit="mg"/>
  <denominator xsi:type="PQ" value="1"/>
</pharm:quantity>
```

745

*Add Section 6.3.4.2*

### 6.3.4.2 Prescription Item Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.2)

750

A Prescription Item belongs to one prescription and represents one prescribed medication. It may be associated with one or more observations. Prescription Item is the atomic entity for logistics, distribution and billing. It contains the prescribed medicine and dosage information as well as other information to the prescribed item such as patient- and fulfillment instructions and substitution handling.

#### 6.3.4.2.1 Standards

755

This part describes the general structure for a Prescription Item. It is based on the following standards:

HL7V3 NE2009	<a href="#">HL7 V3 2009 Normative Edition</a>
CCD	<a href="#">ASTM/HL7 Continuity of Care Document</a>
IHE PCC	<a href="#">Medications Entry (1.3.6.1.4.1.19376.1.5.3.1.4.7)</a>

#### 6.3.4.2.2 Parent Template

760

This entry content module is based on the HL7 CCD template medication activity 2.16.840.1.113883.10.20.1.24 and inherits the structure of the Medication Entry Content Module, 1.3.6.1.4.1.19376.1.5.3.1.4.7.

### 6.3.4.2.3 Specification

This section makes use of the medicine and instruction entry content modules.

765

This specification relies on the PCC Medication Entry Content Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification and only describes additional constraints.

770 The chapters below identify and describe these fields, and indicate the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

```

775 <substanceAdministration classCode='SBADM' moodCode='INT'>
    <templateId root='2.16.840.1.113883.10.20.1.24' />
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7' />
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.2' />
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.6' />
    <id root=' ' extension=' ' />
    <code code=' ' codeSystem=' ' displayName=' ' codeSystemName=' ' />
    <text><reference value="#med-1"/></text>
    <statusCode code='completed' />
    <effectiveTime xsi:type='IVL_TS'>
        <low value=' '/>
        <high value=' '/>
    </effectiveTime>
    <effectiveTime operator='A' xsi:type='TS|PIVL_TS|EIVL_TS|PIVL_PPD_TS|SXPRTS'>
        :
    </effectiveTime>
    <repeatNumber value=' ' />
    <routeCode code=' ' codeSystem=' ' displayName=' ' codeSystemName=' ' />
    <approachSiteCode code=' ' codeSystem=' ' displayName=' ' codeSystemName=' ' />
    <doseQuantity value=' ' unit=' ' />
    <rateQuantity value=' ' unit=' ' />
    <consumable>
        :
        <!-- Medicine Entry Module (1.3.6.1.4.1.19376.1.9.1.3.1) -->
        :
    </consumable>
    <!--
        Author of the prescription in case of usage elsewhere as in a PRE document
-->
    <author>
        :
    </author>
    <!-- 0..* entries describing the components -->
    <entryRelationship typeCode='COMP'>
        <sequenceNumber value=' ' />
        :
    </entryRelationship>
    <!-- An optional entry relationship that indicates the reason for use -->
    <entryRelationship typeCode='RSON'>
        <act classCode='ACT' moodCode='EVN'>
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4.1' />
            <id root=' ' extension=' ' />
        </act>
    </entryRelationship>
    <!-- Reference to a related prescription activity (supply) -->
    <entryRelationship typeCode='REFR'>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3' />
    <!--
        Optional instructions for the patient -->
    <entryRelationship typeCode='SUBJ' inversionInd='true'>
        <act classCode='ACT' moodCode='INT'>
            <templateId root='2.16.840.1.113883.10.20.1.49' />
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3' />
            <code code='PINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'
                codeSystemName='IHEActCode' />
            ...
    </entryRelationship>
    <!-- Optional instructions for Pharmacist -->
    <entryRelationship typeCode='SUBJ' inversionInd='true'>
        <act classCode='ACT' moodCode='INT'>
            <templateId root='2.16.840.1.113883.10.20.1.43' />
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3.1' />
            <code code='FINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'
                codeSystemName='IHEActCode' />
            ...
    </entryRelationship>

```

```

840     </entryRelationship>
841     <!-- Amount of units of the consumable to dispense -->
842     <entryRelationship typeCode='COMP'>
843         <templateId root='1.3.6.1.4.1.19376.1.9.1.3.8' /> <!-- PHARM -->
844         <supply classCode='SPLY' moodCode='RQO'>
845             <independentInd value='false' />
846             <quantity value=' ' unit=' '/>
847         </supply>
848     </entryRelationship>

849     <!-- Substitution handling -->
850     <entryRelationship typeCode='COMP'>
851         <templateId root='1.3.6.1.4.1.19376.1.9.1.3.9' /> <!-- PHARM -->
852         <supply classCode="SPLY" moodCode="RQO">
853             <independentInd value="false" />
854             <pharm:subjectOf4>
855                 <pharm:substitutionPermission classCode="SUBST" moodCode="PERM">
856                     <pharm:code code='E' displayName='equivalent' codeSystem='2.16.840.1.113883.5.1070' codeSystemName='HL7 Substance Admin Substitution' />
857                 </pharm:substitutionPermission>
858             </pharm:subjectOf4>
859         </supply>
860     </entryRelationship>
861     <
862         <precondition>
863             <criterion>
864                 <text><reference value=' '></text>
865             </criterion>
866         </precondition>
867     </substanceAdministration>

```

### 6.3.4.2.3.1 Prescription Item Entry General Specification

870 <substanceAdministration classCode='SBADM' moodCode='INT'>  
871 ...  
872 </substanceAdministration>

The moodCode SHALL be set to INT.

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

875 **6.3.4.2.3.2 Prescription Item Entry TemplateID**

<templateId root='2.16.840.1.113883.10.20.1.24' />	<!-- CCD -->
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7' />	<!-- PCC -->
<templateId root='1.3.6.1.4.1.19376.1.9.1.3.2' />	<!-- PHARM -->

880 A templateId of '1.3.6.1.4.1.19376.1.9.1.3.2' SHALL be present to indicate that this entry is conforming to the Prescription Item Entry Content Module.

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

### 6.3.4.2.3.3 Prescription Item Entry Additional Template ID

<templateId root=' '/>

885 The <templateId> element identifies this <entry> as a particular type of medication event, allowing for validation of the content.

The templateId must use one of the values in the table below for the root attribute.

<b>Root</b>	<b>Description</b>
1.3.6.1.4.1.19376.1.5.3.1.4.7.1	A "normal" <substanceAdministration> act that may not contain any subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.8	A <substanceAdministration> act that records tapered dose information in subordinate <substanceAdministration> act.
1.3.6.1.4.1.19376.1.5.3.1.4.9	A <substanceAdministration> act that records split dose information in subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.10	A <substanceAdministration> act that records conditional dose information in subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.11	A <substanceAdministration> act that records combination medication component information in subordinate <substanceAdministration> acts.

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

#### 890 **6.3.4.2.3.4 Prescription Item ID**

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

This ID represents the Prescription Item ID and SHALL be present.

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

#### **6.3.4.2.3.5 Code**

895 <**code code=' '** displayName=' ' codeSystem=' ' codeSystemName=' '/>

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

#### **6.3.4.2.3.6 Narrative Text**

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

900 This element SHALL be present. The URI given in the value attribute of the <reference> element points to an element in the narrative content that contains the complete text describing the medication prescribed.

See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

#### **6.3.4.2.3.7 Status Code**

<**statusCode code='completed'>**

905 See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

Please note that this element does NOT represent the status of the Prescription Item (e.g., still open, already dispensed, etc.). There is no dedicated data element to record such a status, please

refer to the Community Medication Prescription and Dispense (CMPD) Profile for more information.

910      **6.3.4.2.3.8 Dosage Instructions**

The Prescription Item SHALL contain dosage instructions according to the specification of the dosage instructions in the Dosage Instructions Content Module (1.3.6.1.4.1.19376.1.9.1.3.6).

Note: The following elements part of the Dosage Instructions:

- Prescription Item Entry Additional Template ID
- Effective Time (Duration of Treatment)
- Medication Frequency
- Route of Administration
- Approach Site Code
- Dose Quantity
- Rate Quantity
- Related Components

920      **6.3.4.2.3.9 Number of repeats/refills**

<repeatNumber value=' '/>

925      The <repeatNumber> element SHALL be present, and SHALL contain the number of allowed repeats/refills of the Prescription Item.

If the <repeatNumber> element has a value of zero no repeat/refill is allowed (single dispense). The total number of dispenses will be thus equal to the repeat number plus one.

930      **6.3.4.2.3.10 Consumable**

<consumable>

930      :

      <!-- Medicine Entry Module (1.3.6.1.4.1.19376.1.9.1.3.1) -->

      :

</consumable>

935      The <consumable> element SHALL be present, and shall contain a medication entry, conforming to the Medicine Entry template (1.3.6.1.4.1.19376.1.9.1.3.1).

See PHARM-TF3, Medicine Entry Module (1.3.6.1.4.1.19376.1.9.1.3.1) specification.

930      **6.3.4.2.3.11 Prescriber**

<author>...</author>

940 In the case that the Prescription Item is used within a Prescription document according to the “Pharmacy Prescription” (PRE) Profile this element SHALL NOT be present.

945 In all other cases (e.g., when used in a “Pharmacy Medication List” document according to the PML Profile or in medication sections of other documents as for example Discharge Summaries, etc.) this element SHOULD be present and represent the prescriber of the Prescription Item. It SHALL contain the author and the timestamp of the Prescription document in which the Prescription Item was described (or, if given, the author of the Prescription section within the Prescription document).

Data element	HL7 V3 Data Type	CDA Body position (relative XPath expression)
Prescriber Profession	CE	<i>author/functionCode</i>
Timestamp of prescribing	TS	<i>author/time</i>
Prescriber ID	II	<i>author/assignedAuthor/id</i>
Prescriber Specialty	CE	<i>author/assignedAuthor/code</i>
Prescriber Name	PN	<i>author/assignedAuthor/assignedPerson/name</i>
Prescriber Organization Identifier	II	<i>author/assignedAuthor/representedOrganization/id</i>
Prescriber Organization Name	ON	<i>author/assignedAuthor/representedOrganization/name</i>
Prescriber Organization Address	AD	<i>author/assignedAuthor/representedOrganization/addr</i>

#### 6.3.4.2.3.12 Reason

950 <entryRelationship typeCode='RSON'>  
     <act classCode='ACT' moodCode='EVN'>  
         <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4.1'>  
             <id root=' ' extension=' '/>  
         </act>  
     </entryRelationship>

955 See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

#### 6.3.4.2.3.13 Reference to Medication Treatment Plan Item

<entryRelationship typeCode='REFR'>  
     <templateId root='1.3.6.1.4.1.19376.1.9.1.3.10'> <!-- PHARM -->

```
960   <substanceAdministration classCode='SBADM' moodCode='INT'>
961     ...
962     <consumable>
963       <manufacturedProduct>
964         <manufacturedMaterial nullFlavor='NA' />
965       </manufacturedProduct>
966     </consumable>
967   </substanceAdministration>
968 </entryRelationship>
```

If the prescription is related to a Medication Treatment Plan Item, the reference to it SHALL be present and shall contain a Reference to Medication Treatment Plan Item Entry, conforming to the Reference to Medication Treatment Plan Item Entry template (1.3.6.1.4.1.19376.1.9.1.3.10). See PHARM-TF3, Reference to Medication Treatment Plan Item Entry Module (1.3.6.1.4.1.19376.1.9.1.3.10) specification.

970 Prescription not related to Medication Treatment Plan: This element SHALL NOT be present if the prescription was performed without referring to a Medication Treatment Plan Item.

#### 6.3.4.2.3.14 Reference to a related prescription activity (supply)

```
<entryRelationship typeCode='REFR'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3' />
  ...
</entryRelationship>
```

975 See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.  
This element SHALL NOT be present.

#### 6.3.4.2.3.15 Patient Medication Instructions

```
<entryRelationship typeCode='SUBJ' inversionInd='true'>
  980    <act classCode='ACT' moodCode='INT'>
    <templateId root='2.16.840.1.113883.10.20.1.49' />
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3' />
    <code code='PINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'
      codeSystemName='IHEActCode' />
  990    ...
</entryRelationship>
```

At most one instruction MAY be provided for each <substanceAdministration> entry. When present, this entry relationship SHALL contain a [Patient Medication Instructions](#) (1.3.6.1.4.1.19376.1.5.3.1.4.3) entry.

- 995 Patient Medication Instructions (used in a Prescription Item) are comments from “prescriber to patient” and may contain the following information:
- Human readable dosage instructions (e.g., a representation of the structured dosage instructions as narrative text, any special dosage instructions which could not have been represented in structured way, etc.)
  - General comments by the prescriber to the patient (e.g., “take with food”, etc.)
- 1000

#### 6.3.4.2.3.16 Fulfillment Instructions

```
<entryRelationship typeCode='SUBJ' inversionInd='true'>  
  <act classCode='ACT' moodCode='INT'>  
    <templateId root='2.16.840.1.113883.10.20.1.43'/>  
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.3.1'/>  
    <code code='FINSTRUCT' codeSystem='1.3.6.1.4.1.19376.1.5.3.2'  
         codeSystemName='IHEActCode' />  
    ...  
</entryRelationship>
```

- 1005 1010 At most one instruction MAY be provided for each <substanceAdministration> entry. When present, this entry relationship SHALL contain a [Medication Fulfillment Instructions](#) (1.3.6.1.4.1.19376.1.5.3.1.4.3.1) entry.

Fulfillment Instructions (used in a Prescription Item) are comments from “prescriber to dispenser” and may contain the following information:

- 1015 1020
- A proposal of a product/brand including information about substitution in case of prescribing Generic/Scientific names (e.g., Prescriber prescribes the generic “Paracetamol” but proposes the product “Adol 500mg Caplet” to be dispensed because the patient is used to that medicine)
  - Information to the preparation of compound medicine (e.g., “20 capsules of phenytoin, 20 ml glycerin, 2ml alcohol, Q.S. Syrup to 200ml”)
  - General comments by the prescriber to the dispenser (e.g., if the patient is very old: “Patient is instructed about the dosing, but please repeat the instruction to ensure that the patient understood how to intake the medicine”)

#### 6.3.4.2.3.17 Amount of units of the consumable to dispense

- 1025 <entryRelationship typeCode='COMP'>

```
<supply classCode='SPLY' moodCode='RQO'>
```

```
...
```

```
</supply>
```

```
</entryRelationship>
```

- 1030 This element MAY be present and describes the amount of units to be dispensed. If present, it shall contain a quantity conforming to the Amount of units of the consumable Entry template (1.3.6.1.4.1.19376.1.9.1.3.8). See PHARM-TF3, Amount of units of the consumable Entry Module (1.3.6.1.4.1.19376.1.9.1.3.8) specification.

#### 1035 6.3.4.2.3.18 Substitution handling

```
<entryRelationship typeCode='COMP'>
```

```
<supply classCode="SPLY"
```

```
    moodCode="RQO">
```

```
        <independentInd value="false" />
```

1040 ...

```
</supply>
```

```
</entryRelationship>
```

- One or more `<entryRelationship>` elements, each containing a Substitution handling entry, MAY be present and describe the substitution handling. If present, it shall contain a Substitution handling entry conforming to the Substitution handling Entry template (1.3.6.1.4.1.19376.1.9.1.3.9). See PHARM-TF3, Substitution handling Entry Module (1.3.6.1.4.1.19376.1.9.1.3.9) specification.

#### 6.3.4.2.3.19 ID of parent container (Prescription document)

```
<reference typeCode='XCRPT'>
```

1050 

```
<externalDocument>
```

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

```
</externalDocument>
```

```
</reference>
```

- In the case that the Prescription Item is used within a Prescription document according to the “Pharmacy Prescription” (PRE) Profile this element SHALL NOT be present.

In all other cases (e.g., when used in a “Pharmacy Medication List” document according to the PML Profile or in medication sections of other documents as for example Discharge Summaries,

etc.) this element SHOULD be present and contain the identifier of the Prescription document, the Prescription Item initially has been created.

1060 **6.3.4.2.3.20 Precondition Criterion**

<precondition>

<criterion>

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

</criterion>

1065 </precondition>

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

This element MAY be present.

1070 See PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification.

### **6.3.4.6 Dosage Instructions Content Module (1.3.6.1.4.1.19376.1.9.1.3.6)**

Dosage Instructions are a set of data elements which together represent the dosage instructions to a medication such as duration of treatment, medication frequency, dose quantity, route of administration, etc.

1075 **Note:** Dosage Instructions may be provided structured and/or narrative. This chapter describes *structured* dosage instructions.

If dosage instructions are provided *narrative only*, the Dosage Instructions data elements SHALL NOT be present as a whole—are either null-flavored or omitted.

#### **6.3.4.6.1 Standards**

1080 This part describes the general structure for a Dosage Instructions. It is based on the following standards:

HL7V3 NE2009	<a href="#">HL7 V3 2009 Normative Edition</a>
CDAR2	<a href="#">HL7 CDA Release 2.0</a>

#### **6.3.4.6.2 Parent Template**

This content module has no parent structure.

1085 **6.3.4.6.3 Specification**

The chapters below identify and describe these fields, and indicate the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

```

1090 <substanceAdministration classCode='SBADM' moodCode='INT|EVN'>
    :
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.6' />
    :
1095    <effectiveTime xsi:type='IVL_TS'>
        <low value=' '/>
        <high value=' '/>
    </effectiveTime>
    <effectiveTime operator='A' xsi:type='TS|PIVL_TS|EIVL_TS|PIVL_PPD_TS|SXPR_TS'>
        ...
    </effectiveTime>
    :
    <routeCode code=' ' codeSystem=' ' displayName=' ' codeSystemName=' ' />
    <approachSiteCode code=' ' codeSystem=' ' displayName=' ' codeSystemName=' ' />
    <doseQuantity value=' ' unit=' '/>
    <rateQuantity value=' ' unit=' '/>
    :
    <!-- 0..* entries describing the components -->
    <entryRelationship typeCode='COMP'>
        <sequenceNumber value=' '/>
        :
        <consumable>
            <manufacturedProduct>
                <manufacturedMaterial nullFlavor='NA' />
            </manufacturedProduct>
        </consumable>
        :
    </entryRelationship>
    :
</substanceAdministration>
```

1120

**6.3.4.6.3.1 Dosage Instructions General Specification**

```
<substanceAdministration classCode='SBADM' moodCode='INT|EVN'>
    ...
</substanceAdministration>
```

1125 The Dosage Instructions are part of a &lt;substanceAdministration&gt; element.

The <substanceAdministration> element itself as well as additional constraints applying to this <substanceAdministration> element (e.g., other applying content module constraints such as IHE PHARM ‘Prescription Item’) are specified in the parent specification using this content module.

**6.3.4.6.3.2 Dosage Instructions TemplateID**

1130 &lt;templateId root='1.3.6.1.4.1.19376.1.9.1.3.6' /&gt;                   &lt;!-- PHARM --&gt;

The <templateId> element identifies this <entry> as a particular type of event, allowing for validation of the content.

The templateId SHALL be set to the value 1.3.6.1.4.1.19376.1.9.1.3.6.

#### **6.3.4.6.3.3 Dosage Instructions Additional Template ID**

1135 **<templateId root=' '/>**

This templateId declares which type of dosage instructions (“normal dosing”, “split dosing”, etc.) is used.

The templateId SHALL be present.

1140 In case (structured) dosage instructions are not provided (e.g., because they are only provided as narrative text), this element SHALL be set to ‘Normal Dosing’.

The templateId SHALL use one of the values in the table below for the root attribute.

<b>Root</b>	<b>Description</b>
1.3.6.1.4.1.19376.1.5.3.1.4.7.1	A "normal" <substanceAdministration> act that may not contain any subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.8	A <substanceAdministration> act that records tapered dose information in subordinate <substanceAdministration> act.
1.3.6.1.4.1.19376.1.5.3.1.4.9	A <substanceAdministration> act that records split dose information in subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.10	A <substanceAdministration> act that records conditional dose information in subordinate <substanceAdministration> acts.
1.3.6.1.4.1.19376.1.5.3.1.4.11	A <substanceAdministration> act that records combination medication component information in subordinate <substanceAdministration> acts.

1145 The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

#### **6.3.4.6.3.4 Effective Time (Duration of Treatment)**

**<effectiveTime xsi:type='IVL\_TS'>**

1150 In case the (structured) dosage instructions include a dose regime<sup>10</sup> this element SHALL be present and specify the entire duration of the medication treatment. In case the Duration of Treatment is unknown the <low> and <high> sub-elements of this element SHALL be set to null flavor “UNK”.

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

In all other cases this element MAY be present.

1155 The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

---

<sup>10</sup> For the scope of this Content Module, “dose regime” groups the data elements Medication Frequency, Dose Quantity and Rate Quantity, provided in a certain type of dosing (“Normal dosing”, “Split dosing”, etc.)

**6.3.4.6.3.5 Medication frequency**

<effectiveTime operator='A'

xsi:type='TS|PIVL\_TS|EIVL\_TS|PIVL\_PPD\_TS|SXPR\_TS' />

1160 In case the (structured) dosage instructions include a dose regime using “Normal Dosing”, this element SHALL be present and SHALL NOT be null flavor.

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

In all other cases this element MAY be present.

The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

1165 **6.3.4.6.3.6 Route of Administration**

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

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

In all other cases this element MAY be present.

1170 The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

**6.3.4.6.3.7 Approach Site Code**

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

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

In all other cases this element MAY be present.

1175 The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

**6.3.4.6.3.8 Dose Quantity**

<doseQuantity value=' ' unit=' ' />

1180 In case the (structured) dosage instructions include a dose regime using “Normal Dosing”, this element SHALL be present and SHALL NOT be null flavor.

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

In all other cases this element MAY be present.

The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

1185 **6.3.4.6.3.9 Rate Quantity**

<rateQuantity value=' ' unit=' ' />

In case the (structured) dosage instructions include a dose regime using “Normal Dosing”, this element SHALL be present and SHALL NOT be null flavor.

If dosage instructions are provided *narrative only*, this element SHALL NOT be present.

1190 In all other cases this element MAY be present.

The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

#### 6.3.4.6.3.10 Related components

<entryRelationship typeCode='COMP'>

1195 <sequenceNumber value=''/>

:

<consumable>

<manufacturedProduct>

<manufacturedMaterial nullFlavor='NA'/'>

1200 </manufacturedProduct>

</consumable>

:

</entryRelationship>

1205 In case the (structured) dosage instructions include a dose regime using “Normal Dosing”, subordinate <substanceAdministration> entries SHALL NOT be present.

If dosage instructions are provided *narrative only*, subordinate <substanceAdministration> entries SHALL NOT be present.

In case the (structured) dosage instructions include a dose regime using other types than “Normal Dosing”, one or more subordinate <substanceAdministration> entries SHALL be present.

1210 In case subordinate <substanceAdministration> entries are present, the <consumable> element of subordinate <substanceAdministration> entries SHALL NOT contain a medicine, but the <manufacturedProduct/manufacturedMaterial> element SHALL be set to nullFlavor='NA'.

The constraints defined for this element in PCC TF2, Medication Entry Module (1.3.6.1.4.1.19376.1.5.3.1.4.7) specification apply to this element.

1215

Add new Section 6.3.4.7

### **6.3.4.7 Amount of units of the consumable Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.8)**

1220 The amount of units of the consumable entry content module is part of the <consumable> element.

The amount of units of the consumable entry content module describes the quantity of the enclosing element.

#### **6.3.4.7.1 Standards**

1225 This part describes the general structure for a Prescription Item. It is based on the following standards:

<b>HL7V3 NE2009</b>	<a href="#">HL7 V3 2009 Normative Edition</a>
<b>CCD</b>	<a href="#">ASTM/HL7 Continuity of Care Document</a>
<b>IHE PCC</b>	<a href="#">Medications Entry (1.3.6.1.4.1.19376.1.5.3.1.4.7)</a>

#### **6.3.4.7.2 Parent Template**

This entry content module has no parent structure.

1230 **6.3.4.7.3 Specification**

The chapters below identify and describe these fields, and indicate the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

1235 

```
<entryRelationship typeCode='COMP'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.8' /> <!-- PHARM -->
    <supply classCode='SPLY' moodCode='RQO'>
        <independentInd value='false' />
        <quantity value=' ' unit=' '/>
    </supply>
</entryRelationship>
```

1240

#### **6.3.4.7.3.1 Amount of units of the consumable Entry General Specification**

```
<entryRelationship typeCode='COMP'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.8' /> <!-- PHARM -->
    <supply classCode='SPLY' moodCode='RQO'>
        <independentInd value='false' />
        <quantity value=' ' unit=' '/>
```

</supply>

1250 </entryRelationship>

#### **6.3.4.7.3.2 Amount of units of the consumable Entry Template ID**

<templateId root='1.3.6.1.4.1.19376.1.9.1.3.8'>

The Template ID for an Amount of units of the consumable Entry SHALL be provided.

#### **6.3.4.7.3.3 Amount of units of the consumable Entry Quantity**

1255 The amount of units of the consumable entry content module describes the quantity of the enclosing <consumable> element.

- If the enclosing <consumable> - element also contains package information, the <quantity> element SHALL contain the amount of primary packaged items of the medication. The value shall refer to the primary layer of the package information given in the <pharm:asContent> element of the consumable (e.g., if the value is 2 and the <pharm:asContent> element describes a blister containing 30 tablets, this means that 2 blisters (with each 30 tablets in it) have been prescribed). Eventually present sub- or super layers of packaging (subContent, asSuperContent elements below the asContent element) are not affected. In this case the unit attribute SHALL NOT be present.
- If the enclosing <consumable> - element does not contain package information, the <quantity> element SHALL contain the amount of consumable units of the medication. In this case the unit attribute MAY be present, if the quantity is in non-countable units. The value of the unit SHALL be out of the UCUM code system.

1270 *Add new Section 6.3.4.8*

#### **6.3.4.8 Substitution handling Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.9)**

An <entryRelationship> element, containing one and only one <pharm:subjectOf4> element describing the substitution handling. The Substitution handling entry content module is part of the <consumable> element.

1275 Technical note: A part of an HL7 Medication Order (PORX\_MT010120UV) is used within the entryRelationship/supply element to express Substitution Handling.

#### **6.3.4.8.1 Standards**

This part describes the general structure for a Prescription Item. It is based on the following standards:

1280

HL7V3 NE2009	<a href="#">HL7 V3 2009 Normative Edition</a>
CCD	<a href="#">ASTM/HL7 Continuity of Care Document</a>

IHE PCC	<a href="#">Medications Entry (1.3.6.1.4.1.19376.1.5.3.1.4.7)</a>
---------	---

### 6.3.4.8.2 Parent Template

This entry content module has no parent structure.

### 6.3.4.8.3 Specification

1285 The chapters below identify and describe these fields, and indicate the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

```

1290 <entryRelationship typeCode='COMP'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.9' /> <!-- PHARM -->
    <supply classCode="SPLY"
        moodCode="RQO">
        <independentInd value="false" />
        <pharm:subjectOf4>
            <pharm:substitutionPermission
                classCode="SUBST"
                moodCode="PERM">
                <pharm:code code=' '
                    displayName=' '
                    codeSystem='2.16.840.1.113883.5.1070'
                    codeSystemName='HL7 Substance Admin Substitution' />
            </pharm:substitutionPermission>
        </pharm:subjectOf4>
    </supply>
</entryRelationship>
```

#### 6.3.4.8.3.1 Substitution handling Entry General Specification

```

<entryRelationship typeCode='COMP'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.9' /> <!-- PHARM -->
    <supply classCode="SPLY"
        moodCode="RQO">
        <independentInd value="false" />
        ...
    </supply>
</entryRelationship>
```

#### 6.3.4.8.3.2 Substitution handling Entry Template ID

```
<templateId root='1.3.6.1.4.1.19376.1.9.1.3.9' />
```

The Template ID for Substitution handling Entry SHALL be provided.

1320    **6.3.4.8.3.3 Substitution handling Entry value**

```

<pharm:subjectOf4>
  <pharm:substitutionPermission>
    classCode="SUBST"
    moodCode="PERM">
<pharm:code code='
  displayName=' '
  codeSystem='2.16.840.1.113883.5.1070'
  codeSystemName='HL7 Substance Admin Substitution'>
</pharm:substitutionPermission>
</pharm:subjectOf4>

```

1330

The <value> element identifies what sort of change is permitted between the therapy that was ordered and the therapy that will be provided. It shall be coded in HL7 terminology for substance substitution.

1335

<i>Add Section 6.3.4.9</i>
----------------------------

### **6.3.4.9 Reference to Medication Treatment Plan Item Entry Content Module (1.3.6.1.4.1.19376.1.9.1.3.10)**

1340 An <entryRelationship> element, containing one and only one reference to a Medication Treatment Plan (MTP) Item.

#### **6.3.4.9.1 Standards**

This part describes the general structure for a Reference to a Medication Treatment Plan Item. It is based on the following standards:

<b>HL7V3 NE2009</b>	<a href="#">HL7 V3 2009 Normative Edition</a>
<b>CCD</b>	<a href="#">ASTM/HL7 Continuity of Care Document</a>
<b>IHE PCC</b>	<a href="#">Medications Entry (1.3.6.1.4.1.19376.1.5.3.1.4.7)</a>

1345

### 6.3.4.9.2 Parent Template

This entry content module has no parent structure.

### 6.3.4.9.3 Specification

The chapters below identify and describe these fields, and indicate the constraints on whether or  
1350 not they are required to be sent. The fields are listed in the order that they appear in the CDA  
XML content.

```
1355 <entryRelationship typeCode='REFR'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.10' />
    <substanceAdministration classCode='SBADM' moodCode='INT'>
        <id root=' ' extension=' '/>      <!-- Medication Treatment Plan Item Id -->
        <code code='MTPItem'
            codeSystem='1.3.6.1.4.1.19376.1.9.2.2'
            displayName='Medication Treatment Plan Item'
            codeSystemName='IHE Pharmacy Item Type List'/>
        <consumable>
            <manufacturedProduct>
                <manufacturedMaterial nullFlavor='NA' />
            </manufacturedProduct>
        </consumable>
    </substanceAdministration>
</entryRelationship>
```

#### 6.3.4.9.3.1 Reference to Medication Treatment Plan Item Entry General 1370 Specification

```
<entryRelationship typeCode='REFR'>
    <templateId root='1.3.6.1.4.1.19376.1.9.1.3.10' />
    <substanceAdministration classCode='SBADM' moodCode='INT'>
        ...
    <consumable>
        <manufacturedProduct>
            <manufacturedMaterial nullFlavor='NA' />
        </manufacturedProduct>
    </consumable>
    </substanceAdministration>
</entryRelationship>
```

#### 6.3.4.9.3.2 Reference to Medication Treatment Plan Item Entry Template ID

```
<templateId root='1.3.6.1.4.1.19376.1.9.1.3.10' />
```

1385 The Template ID for Reference to Medication Treatment Plan Item Entry SHALL be provided.

#### **6.3.4.9.3.3 Reference to Medication Treatment Plan Item Entry ID**

<id root=' ' extension=' '/> <!-- Medication Treatment Plan Item Id -->

This ID represents the referenced Medication Treatment Plan Item Id and SHALL be present.

1390

#### **6.3.4.9.3.4 Reference to Medication Treatment Plan Item Entry CODE**

<code code='MTPItem'  
codeSystem='1.3.6.1.4.1.19376.1.9.2.2'  
displayName=' Medication Treatment Plan Item'  
codeSystemName='IHE Pharmacy Item Type List' />

1395

This code indicates that the referenced item is a Medication Treatment Plan Item. It SHALL be present.

**Add Section 6.3.4.10**

#### **6.3.4.10 Reference to Prescription Item Entry Content Module**

##### **(1.3.6.1.4.1.19376.1.9.1.3.11)**

1400

An <entryRelationship> element, containing one and only one reference to a Prescription (PRE) Item.

#### **6.3.4.10.1 Standards**

1405

This part describes the general structure for a reference to a Prescription Item. It is based on the following standards:

<b>HL7V3 NE2009</b>	<a href="#">HL7 V3 2009 Normative Edition</a>
<b>CCD</b>	<a href="#">ASTM/HL7 Continuity of Care Document</a>
<b>IHE PCC</b>	<a href="#">Medications Entry (1.3.6.1.4.1.19376.1.5.3.1.4.7)</a>

#### **6.3.4.10.2 Parent Template**

This entry content module has no parent structure.

**1410 6.3.4.10.3 Specification**

The chapters below identify and describe these fields, and indicate the constraints on whether or not they are required to be sent. The fields are listed in the order that they appear in the CDA XML content.

1415     <entryRelationship typeCode='REFR'>  
      <templateId root='1.3.6.1.4.1.19376.1.9.1.3.11' />  
      <substanceAdministration classCode='SBADM' moodCode='INT'>  
        <id root=' ' extension=' ' />    <!-- Prescription Item Id -->  
        <code code='PREItem'  
          codeSystem='1.3.6.1.4.1.19376.1.9.2.2'  
          displayName='Prescription Item'  
          codeSystemName='IHE Pharmacy Item Type List' />  
        <consumable>  
          <manufacturedProduct>  
            <manufacturedMaterial nullFlavor='NA' />  
          </manufacturedProduct>  
        </consumable>  
      </substanceAdministration>  
   </entryRelationship>

1430

**6.3.4.10.3.1 Reference to Prescription Item Entry General Specification**

<entryRelationship typeCode='REFR'>  
  <templateId root='1.3.6.1.4.1.19376.1.9.1.3.11' />  
  <substanceAdministration classCode='SBADM' moodCode='INT'>  
    ...  
    <consumable>  
      <manufacturedProduct>  
        <manufacturedMaterial nullFlavor='NA' />  
      </manufacturedProduct>  
    </consumable>  
  </substanceAdministration>  
</entryRelationship>

**6.3.4.10.3.2 Reference to Prescription Item Entry Template ID**

1445     <templateId root='1.3.6.1.4.1.19376.1.9.1.3.11' />

The Template ID for Reference to Prescription Item Entry SHALL be provided.

#### **6.3.4.10.3.3 Reference to Prescription Item Entry ID**

```
<id root=' ' extension=' '/> <!-- Prescription Item Id -->
```

This ID represents the referenced Prescription Item Id and SHALL be present.

1450    **6.3.4.10.3.4 Reference to Prescription Entry CODE**

```
<code code='PREItem'  
      codeSystem='1.3.6.1.4.1.19376.1.9.2.2'  
      displayName='Prescription Item'  
      codeSystemName='IHE Pharmacy Item Type List' />
```

1455    This code indicates that the referenced item is a Prescription Item. It SHALL be present.

# Appendices

## **Appendix A Validating CDA Documents using the Framework**

### **1460 A.1 Validating Documents**

For validation of document content modules please refer to PCC-TF2, Appendix A.1.

### **A.2 Validating Sections**

For validation of section content modules please refer to PCC-TF2, Appendix A.2.

### **A.3 Phases of Validation and Types of Errors**

### **1465 For the phases of validation and types of errors please refer to PCC-TF2, Appendix A.3.**

## Appendix B Extensions to CDA Release 2

### B.1 IHE PHARM Extensions

All Extensions to CDA Release 2.0 created by the IHE PHARM Technical Committee are in the namespace `urn:ihe:pharm:medication`.

- 1470 The approach used to create extension elements created for the PHARM Technical Framework is the same as was used for the PCC Technical Framework, the HL7 Care Record Summary (see Appendix E) and the ASTM/HL7 Continuity of Care Document (see Section 7.2).

#### B.1.1 Used for Medicine Entry Content Module

- 1475 The extensions of CDA Release 2 used for the Medicine Entry Content Module are derived of a medication structure based on a standard HL7 V3 Common Message Element Type (CMET) created by the HL7 Pharmacy group. The used CMET is “**R\_Medication Universal**” (**COCT\_MT230100UV**), **Release 2** and fits within the overall entry by extending the “Manufactured Product” structure of the CDA “substanceAdministration” branch.

- 1480 This structure is part of the HL7 V3 2009 Normative Edition (COCT\_RM230100UV). The incorporation of this structure is done according to Section **1.4 CDA Extensibility** of the HL7 CDA standard. Such an extension of the base CDA standard is an accepted practice in IHE (e.g., in the XD\* Lab specification).

- 1485 The rules of Section **1.4 CDA Extensibility** require the designation of a new XML namespace for the XML elements in this structure. For the purposes of documentation, the **namespace** `urn:ihe:pharm:medication` shall be used.

```
1490 <manufacturedProduct xmlns:pharm="urn:ihe:pharm:medication" classCode="MANU">
    <manufacturedMaterial classCode="MMAT" determinerCode="KIND">
        <templateId root="1.3.6.1.4.1.19376.1.9.1.3.1"/>
        <!-- National medicinal product code (brand-level) -->
        <code code="" displayName="" codeSystem="" codeSystemName="" />
        <!-- Brand name -->
        <name>... </name>
        <!-- Pharmaceutical dose form -->
        <pharm:formCode code="" displayName=""
            codeSystem="" codeSystemName="" />
        <lotNumberText>...</lotNumberText>
        <pharm:expirationTime value=' '/>
        <!-- Container information -->
        <pharm:asContent classCode="CONT">
            <pharm:containerPackagedMedicine classCode="CONT" determinerCode="INSTANCE">
                <!-- Medicinal product code (package-level) -->
                <pharm:code code="" displayName="" codeSystem="" codeSystemName="" />
                <!-- Brand name (package) -->
                <pharm:name>...</pharm:name>
                <pharm:formCode code="" displayName=""
                    codeSystem="" codeSystemName="" />
                <pharm:capacityQuantity value="" unit="" />
                <pharm:asSuperContent>
                    <pharm:containerPackagedMedicine classCode='CONT'
                        determinerCode='INSTANCE'>
                        <pharm:capacityQuantity value=' ' unit=' '/>
                    </pharm:containerPackagedMedicine>
                </pharm:asSuperContent>
            </pharm:containerPackagedMedicine>
        </pharm:asContent>
    <!-- ... -->

```

1500 The detailed usage is shown in the specification of the Medicine Entry Content Module  
1505 (1.3.6.1.4.1.19376.1.9.1.3.1) within the PHARM Technical Framework.

