

Integrating the Healthcare Enterprise



5

IHE Patient Care Coordination (PCC) Technical Framework Supplement

10

CDA Content Modules

15

Rev. 2.6 – Trial Implementation

20 Date: June 17, 2020
 Author: IHE PCC Technical Committee
 Email: pcc@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 IHE Patient Care Coordination Technical Framework V11.0. 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 different than traditional IHE supplements. It serves as the trial implementation staging area for content modules. The content modules (section level, entry level) defined during trial implementation are gathered in this document to provide a central location for readers of supplements from PCC, QRPH and/or other domains that use the dictionary of content modules first defined by PCC. After individual modules are successfully tested and reviewed, they will be moved to final text. At that time, they are removed from this document. Thus, this supplement will continue to exist for some time as new content modules are defined and documented here. Likewise, content modules will be removed as they go to final 40 text.



Note: For some profiles, the domain technical committee has elected to document the content modules in the specific profile supplement; therefore, they are not documented in this supplement.

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

Amend Section X.X by the following:

50 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.

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

55 Information about the IHE PCC domain and the IHE QRPH domain can be found at:
http://www.ihe.net/IHE_Domains.

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 and <http://www.ihe.net/Profiles>.

60 The current version of the IHE PCC Technical Framework and the IHE QRPH Technical Framework can be found at: http://www.ihe.net/Technical_Frameworks.

CONTENTS

	Introduction.....	20
	Profile Abstract	20
65	Open Issues and Questions	20
	Closed Issues.....	20
	Volume 1 – Integration Profiles.....	21
	Glossary	22
	2.5 History of Annual Changes	22
70	Volume 2 – Transactions and Content Modules.....	25
	6.1 Conventions	25
	6.2 Folder Content Modules.....	26
	6.2.1 EDES Folder Specification	26
	6.2.2 APR Folder Specification.....	26
75	6.2.3 LDR Folder Specification	26
	6.3 HL7 Version 3.0 Content Modules	26
	6.3.1 CDA Document Content Modules	26
	6.3.1.x History and Physical Specification 1.3.6.1.4.1.19376.1.5.3.1.1.16.1.4.....	26
	6.3.1.x.1 Format Code	26
80	6.3.1.x.2 LOINC Code.....	26
	6.3.1.x.3 Standards.....	26
	6.3.1.x.4 Specification	27
	6.3.1.x.5 Conformance.....	27
	6.3.2 CDA Header Content Modules	29
85	6.3.2.1 Language Communication 1.3.6.1.4.1.19376.1.5.3.1.2.1.....	29
	6.3.2.2 Employer and School Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.2	29
	6.3.2.3 Healthcare Providers and Pharmacies 1.3.6.1.4.1.19376.1.5.3.1.2.3	29
	6.3.2.4 Patient Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.4	29
90	6.3.2.5 Spouse 1.3.6.1.4.1.19376.1.5.3.1.2.4.1.....	29
	6.3.2.5.1 Parent Template	30
	6.3.2.5.2 Specification	30
	6.3.2.5.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4'><templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.1'>	30
95	6.3.2.5.4 <associatedEntity classCode='PRS'>	30
	6.3.2.5.5 <code code='127848009 184142008' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>	30
	6.3.2.5.6 Completed Example.....	31
100	6.3.2.6 Natural Father of Fetus 1.3.6.1.4.1.19376.1.5.3.1.2.4.2	31
	6.3.2.6.1 Parent Template	31
	6.3.2.6.2 Specification	31
	6.3.2.6.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4'><templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.2'>	32

	6.3.2.6.4 <associatedEntity classCode='PRS'>	32
105	6.3.2.6.5 <code code='xx-fatherofbaby' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />	32
	6.3.2.6.6 Completed Example.....	32
	6.3.2.7 Authorization 1.3.6.1.4.1.19376.1.5.3.1.2.5.....	33
110	6.3.2.7.1 Parent Template	33
	6.3.2.7.2 Specification	33
	6.3.2.7.3 <authorization typeCode='AUTH'>	33
	6.3.2.7.4 <consent classCode='CONS' moodCode='EVN'>.....	33
	6.3.2.7.5 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.5' />	33
115	6.3.2.7.6 <id root=' '/>	33
	6.3.2.7.7 <code code=' ' codeSystem=' ' codeSystemName=' ' displayName=' '/> ... 34	34
	6.3.3 CDA Section Content Modules.....	34
	6.3.3.1 Reasons for Care.....	34
120	6.3.3.1.1 Reason for Referral.....	34
	6.3.3.1.2 Coded Reason for Referral	34
	6.3.3.1.3 Chief Complaint.....	34
	6.3.3.1.4 Hospital Admission Diagnosis	34
	6.3.3.1.5 Proposed Procedure Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.1	34
125	6.3.3.1.6 EBS Estimated Blood Loss Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.2.....	35
	6.3.3.1.7 Proposed Anesthesia Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.3	36
	6.3.3.1.8 Reason for Procedure Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.4.....	37
	6.3.3.1.9 Reason for Visit Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1.1	38
	6.3.3.1.10 Injury Incident Description Section 1.3.6.1.4.1.19376.1.5.3.1.1.19.2.1 ... 38	38
	6.3.3.2 Other Condition Histories.....	39
130	6.3.3.2.1 History of Present Illness.....	39
	6.3.3.2.2 Hospital Course	39
	6.3.3.2.3 Active Problems.....	39
	6.3.3.2.4 Discharge Diagnosis	39
	6.3.3.2.5 History of Past Illness	39
135	6.3.3.2.6 Encounter Histories	39
	6.3.3.2.7 History of Outpatient Visits.....	39
	6.3.3.2.8 History of Inpatient Visits	39
	6.3.3.2.9 List of Surgeries.....	39
	6.3.3.2.10 Coded List of Surgeries	39
140	6.3.3.2.11 Allergies and Other Adverse Reactions.....	39
	6.3.3.2.12 Family medical History	40
	6.3.3.2.13 Coded Family Medical History	40
	6.3.3.2.14 Social History Section	40
	6.3.3.2.15 Functional Status	40
145	6.3.3.2.16 Review of Systems	40
	6.3.3.2.17 Hazardous Working Conditions	40
	6.3.3.2.18 Pregnancy History	40

	6.3.3.2.19 Medical Devices	40
	6.3.3.2.20 Foreign Travel	40
150	6.3.3.2.21 Pre-procedure Family Medical History Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.5 (Deprecated).....	40
	6.3.3.2.22 Coded Functional Status Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.1.....	41
	6.3.3.2.22.1 Standards.....	41
155	6.3.3.2.22.2 Parent Template	41
	6.3.3.2.23 Pain Scale Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2.....	42
	6.3.3.2.24 Braden Score Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3.....	43
	6.3.3.2.25 Geriatric Depression Scale Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4 ..	44
160	6.3.3.2.26 Physical Function Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5.....	44
	6.3.3.2.26.1 Constraints	45
	6.3.3.2.27 Preprocedure Review of Systems Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.13	49
	6.3.3.2.28 Estimated Delivery Date Section 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.1....	50
	6.3.3.2.29 History of Tobacco Use Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.8	51
165	6.3.3.2.30 Current Alcohol/Substance Abuse Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.10	52
	6.3.3.2.31 History of Blood Transfusion Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.12 ..	52
	6.3.3.2.32 Anesthesia Risk Review of Systems Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.14.....	53
170	6.3.3.2.33 Implanted Medical Device Review Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.46.....	54
	6.3.3.2.34 Pregnancy Status Review Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.47	54
	6.3.3.2.35 History of Infection Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1.....	55
	6.3.3.2.36 Coded Social History Section 1.3.6.1.4.1.19376.1.5.3.1.3.16.1	55
175	6.3.3.2.37 Coded History of Infection Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1.1	57
	6.3.3.2.38 Prenatal Events Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.2	57
	6.3.3.2.39 Labor and Delivery Events Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.3 ...	58
180	6.3.3.2.40 Newborn Delivery Information Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.4	60
	6.3.3.2.41 Postpartum Hospitalization Treatment Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.7.....	62
	6.3.3.2.42 Event Outcomes Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9	64
185	6.3.3.2.43 Newborn Status at Maternal Discharge 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.8	64
	6.3.3.2.44 History of Surgical Procedures Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.2	65
	6.3.3.2.45 Operative Note Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.6	65
	6.3.3.2.46 Child Functional Status Assessment 1.3.6.1.4.1.19376.1.7.3.1.1.13.3....	66
190	6.3.3.2.47 Psychomotor Development Section 1.3.6.1.4.1.19376.1.7.3.1.1.13.4.....	66

	6.3.3.2.48 Eating and Sleeping Assessment Section 1.3.6.1.4.1.19376.1.7.3.1.1.13.5	67
195	6.3.3.2.49 Coded Event Outcomes 1.3.6.1.4.1.19376.1.7.3.1.1.13.7	68
	6.3.3.2.50 Intentionally blank	69
	6.3.3.2.51 Intentionally blank	69
	6.3.3.2.52 Intentionally blank	69
	6.3.3.2.53 Notifications, Alerts, and Reminders Section 1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1.x.....	69
200	6.3.3.2.54 Pain Assessment Panel Section 1.3.6.1.4.1.19376.1.5.3.1.1.20.2.4	70
	6.3.3.2.55 History of Cognitive Function Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.11	71
	6.3.3.2.56 Isolation Status Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.8.....	71
	6.3.3.2.57 Restraints Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.10	72
205	6.3.3.2.58 Risk Indicators for Hearing Loss	72
	6.3.3.2.59 Cancer Diagnosis Section 1.3.6.1.4.1.19376.1.7.3.1.3.14.1	73
	6.3.3.3 Medications	74
	6.3.3.3.1 Medications Section.....	74
	6.3.3.3.2 Admission Medication History Section.....	74
210	6.3.3.3.3 Medications Administered Section.....	74
	6.3.3.3.4 Hospital Discharge Medications Section.....	74
	6.3.3.3.5 Immunizations Section	74
	6.3.3.4 Physical Exams.....	74
	6.3.3.4.30 Coded Detailed Physical Examination Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1.....	74
215	6.3.3.4.31 Pelvis Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.10	76
	6.3.3.4.32 Admission Physical Exam Section 1.3.6.1.4.1.19376.1.5.3.1.1.22.1.1.2.1	77
	6.3.3.4.33 Discharge Status 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.12	77
220	6.3.3.5 Relevant Studies	78
	6.3.3.5.1 Results.....	78
	6.3.3.5.2 Coded Results	78
	6.3.3.5.3 Hospital Studies Summary	78
	6.3.3.5.4 Coded Hospital Studies Summary	78
	6.3.3.5.5 Consultations 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.8	78
225	6.3.3.5.6 Antenatal Testing and Surveillance Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5.....	79
	6.3.3.5.7 Coded Antenatal Testing and Surveillance Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5.1.....	79
230	6.3.3.5.8 Intentionally blank	80
	6.3.3.5.9 Intentionally blank	80
	6.3.3.5.10 Intentionally blank	80
	6.3.3.5.11 Hearing Screening Coded Results	80
	6.3.3.5.11.1 Parent Template	81
	6.3.3.6 Plans of Care.....	82

	6.3.3.9.2 Pre-procedure Risk Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.44	98
280	6.3.3.9.3 Antepartum Visit Summary Flowsheet Section 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.....	99
	6.3.3.9.4 Progress Note Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.7.....	100
	6.3.3.9.5 ED Diagnosis Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.9.....	101
	6.3.3.9.6 Acuity Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.2.....	101
285	6.3.3.9.7 Assessments Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.4.....	102
	6.3.3.10 Section Content Modules-Non-categorized	103
	6.3.3.10.1 VRDR Death Report Section- Section Content Module (1.3.6.1.4.1.19376.1.7.3.1.3.23.2).....	103
290	6.3.3.10.1.1 Pregnancy Status Entry Condition.....	105
	6.3.3.10.1.2 Autopsy Results Entry Condition	106
	6.3.3.10.1.3 Death Certification Entry Condition.....	106
	6.3.3.10.1.4 Death Causal Information Entry Condition	106
	6.3.3.10.2 Coded Hospital Course Section 1.3.6.1.4.1.19376.1.7.3.1.3.23.1	106
	6.3.3.10.3 Resources to Support Goals Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.1 ...	107
	6.3.3.10.4 Healthy Weight Care Plan Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.2	108
295	6.3.3.10.5 Occupational Data for Health Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.1	109
	6.3.3.10.5.1 Occupational Data for Health Section < 74166-0>	109
	6.3.4 CDA Entry Content Modules	111
300	6.3.4.25 Family History Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.3	112
	6.3.4.25.1 Standards.....	112
	6.3.4.25.2 Parent Template	112
	6.3.4.25.3 Specification	112
	6.3.4.25.4 <templateId root='2.16.840.1.113883.10.20.1.22'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.3'>	112
305	6.3.4.25.5 <code code='' displayName='' codeSystem='' codeSystemName='' />	112
	6.3.4.25.6 <value xsi:type='CD' code='' displayName='' codeSystem='' codeSystemName='' />.....	112
	6.3.4.26 Pregnancy History Organizer 1.3.6.1.4.1.19376.1.5.3.1.4.13.5.1	113
	6.3.4.27 EDD Observation 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1	113
310	6.3.4.27.1 Specification	114
	6.3.4.27.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1'>	114
	6.3.4.27.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>	115
	6.3.4.27.4 <code code='11778-8' codeSystem='2.16.840.1.113883.6.1'>.....	115
	6.3.4.27.5 <value xsi:type='TS' value=' '>	115
315	6.3.4.27.6 <author typeCode='AUT'><assignedAuthor><id root=' ' extension=' ' /></assignedAuthor></author>	115
	6.3.4.27.7 <author typeCode='AUT'><time value=' '/></author>	115
	6.3.4.27.8 <entryRelationship typeCode='SPRT'>	115
	6.3.4.27.9 <observation> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'> : </observation> [1st nesting] 115	115
320	6.3.4.27.10 <code code=' ' codeSystem='2.16.840.1.113883.6.1'> [1st nesting] ...	116

	6.3.4.27.11 <entryRelationship typeCode='DRIV'>.....	116
325	6.3.4.27.12 <observation> <templateId root=' ' /> : </observation> [2st nesting]	116
	6.3.4.27.13 <code code=' ' codeSystem='2.16.840.1.113883.6.1'> [2nd nesting]..	116
	6.3.4.27.14 <repeatNumber value=' '/><interpretationCode code=' ' codeSystem=' '/> <targetSiteCode code=' ' codeSystem=' '/>.....	117
	6.3.4.28 Antepartum Visit Summary Battery 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2	117
330	6.3.4.28.1 Specification	117
	6.3.4.28.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2' />	118
	6.3.4.28.3 <organizer classCode='BATTERY' moodCode='EVN'>	118
	6.3.4.28.4 <id root=' ' extension=' '/>	118
335	6.3.4.28.5 <code code='(57061-4)' codeSystem='2.16.840.1.113883.6.1'>	118
	6.3.4.28.6 <author/><time/><assignedAuthor><id/></assignedAuthor></author>.....	118
	6.3.4.28.7 <statusCode code='completed' />	118
	6.3.4.28.8 <component>	118
	6.3.4.29 Advance Directive Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.7	120
340	6.3.4.29.1 Standards.....	121
	6.3.4.29.2 Specification	121
	6.3.4.29.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' /> <templateId root='2.16.840.1.113883.10.20.1.17' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.7' />	121
345	6.3.4.29.4 <code code=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />.....	121
	6.3.4.29.5 <value xsi:type='BL' value='true false' />.....	122
350	6.3.4.29.6 <reference typeCode='REFR'> <templateId root='2.16.840.1.113883.10.20.1.36' /> <externalDocument classCode='DOC' moodCode='EVN'> <id root=' ' extension=' '/> <text><reference value=' '/></text>.....	122
	6.3.4.30 Blood Type Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.6.....	122
	6.3.4.30.1 Standards.....	122
	6.3.4.30.2 Specification	122
355	6.3.4.30.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.6' /> <templateId root='2.16.840.1.113883.10.20.1.31' />	123
	6.3.4.30.4 <code code='882-1' displayName='ABO+RH GROUP' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />	123
360	6.3.4.30.5 <repeatNumber value=' '/>	123
	6.3.4.30.6 <value xsi:type='CE' code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>	123
	6.3.4.30.7 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/> <methodCode code=' ' codeSystem=' ' codeSystemName=' '/> <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>	123
365	6.3.4.31 Encounters 1.3.6.1.4.1.19376.1.5.3.1.4.14	124

	6.3.4.33.2.12 <entryRelationship typeCode='RSON'>	130
	6.3.4.34 Transport 1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1.....	130
	6.3.4.35 Encounter Disposition 1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2	130
415	6.3.4.35.1 Specification	131
	6.3.4.35.1.1 <act classCode='ACT' moodCode='INT EVN'>.....	131
	6.3.4.35.1.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2'/>.....	131
	6.3.4.35.1.3 <id root="" extension="/".....	131
	6.3.4.35.1.4 <code code="" displayName="" codeSystem="" codeSystemName="" />	132
420	6.3.4.35.1.5 <text><reference value="#xxx"/></text>.....	132
	6.3.4.35.1.6 <effectiveTime><low value="/" /><high value="/" /><effectiveTime/>	132
	6.3.4.35.1.7 <performer typeCode='PRF'>	132
	6.3.4.35.1.8 <assignedEntity>	132
425	6.3.4.35.1.9 <id root="" extension="/".....	132
	6.3.4.35.1.10 <addr></addr>	132
	6.3.4.35.1.11 <telecom value="" use="/" />	132
	6.3.4.35.1.12 <assignedPerson><name/></assignedPerson>	133
430	6.3.4.35.1.13 <participant typeCode='RCV'> <time value="/" /> <participantRole classCode='ROL'> <id root="" extension="/" /> <addr></addr> <telecom value="" use="/" /> <playingEntity><name/></playingEntity>	133
	6.3.4.36 Reserved for Coverage Activity	133
	6.3.4.37 Reserved for Payer Entry.....	133
	6.3.4.38 Pain Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.1	133
435	6.3.4.38.1 Parent Template	133
	6.3.4.38.2 Specification	134
	6.3.4.38.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />	134
	6.3.4.38.4 <code code='38208 5' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'> <translation code='406127006' displayName='Pain intensity' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />	134
440	6.3.4.38.5 <value xsi:type='CO' value=' ' />	134
	6.3.4.38.6 <interpretationCode code='301379001 40196000 76948002 67849003' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />	135
445	6.3.4.38.7 <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>	135
	6.3.4.38.8 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>	135
450	6.3.4.39 Braden Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.2	136
	6.3.4.40 Braden Score Component 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.3	136
	6.3.4.41 Geriatric Depression Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.4 ..	136
	6.3.4.42 Geriatric Depression Score Component 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.5 ..	136
	6.3.4.43 Survey Panel 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7	136
	6.3.4.43.1 Parent Template	136

455	6.3.4.43.2 Uses.....	136
	6.3.4.43.3 Specification	136
	6.3.4.43.3.1<organizer classCode='CLUSTER' moodCode='EVN'>	137
	6.3.4.43.3.2<templateId root='2.16.840.1.113883.10.20.1.32'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7'>	137
460	6.3.4.43.3.3<id root=' ' extension=' '/>	137
	6.3.4.43.3.4<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>.....	137
	6.3.4.43.3.5<statusCode code='completed'>	137
	6.3.4.43.3.6<effectiveTime value=' '/>	137
465	6.3.4.43.3.7<!-- one or more survey observations --> <component typeCode='COMP'>.....	137
	6.3.4.44 Survey Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6	137
	6.3.4.44.1 Parent Template.....	138
	6.3.4.44.2 Uses.....	138
470	6.3.4.44.3 Specification	138
	6.3.4.44.3.1<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'> <templateId root='2.16.840.1.113883.10.20.1.31'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6'>	138
475	6.3.4.44.3.2<code code=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'>	138
	6.3.4.44.3.3<value xsi:type='CO CD INT PQ' .../>	138
	6.3.4.44.3.4<interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/> 138	
	6.3.4.44.3.5<methodCode code=' ' codeSystem=' ' codeSystemName=' '/> <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/> ...	139
480	6.3.4.45 Acuity 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1	139
	6.3.4.45.1 Specification	139
	6.3.4.45.1.1 <observation classCode='OBS' moodCode='EVN'>	139
	6.3.4.45.1.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1'>	139
485	6.3.4.45.1.3 <id root=" extension="/">.....	139
	6.3.4.45.1.4 <code code=" displayName=" codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>	140
	6.3.4.45.1.5 <originalText><reference value='#xxx'><orginalText>	140
	6.3.4.45.1.6 <text><reference value='#text'></text>	140
490	6.3.4.45.1.7 <effectiveTime>.....	140
	6.3.4.45.1.8 <high value="/">	140
	6.3.4.46 Intravenous Fluids 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.2	140
	6.3.4.46.1 Specification	141
	6.3.4.46.1.1 Medication Fields.....	142
495	6.3.4.46.1.2 <substanceAdministration classCode='SBADM' moodCode='INT EVN'>	142

	6.3.4.46.1.3 <templateId root='2.16.840.1.113883.10.20.1.24'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.1'>	142
500	6.3.4.46.1.4 <id root="" extension="/">	142
	6.3.4.46.1.5 <code code="" displayName="" codeSystem="" codeSystemName="">	143
	6.3.4.46.1.6 <text><reference value=""></text>	143
	6.3.4.46.1.7 <statusCode code='completed active'>	143
505	6.3.4.46.1.8 <effectiveTime xsi:type='IVL_TS'>	143
	6.3.4.46.1.9 <low value=""><high value="">	143
	6.3.4.46.1.10 <approachSiteCode code="" codeSystem=""> originalText><reference value="/"></originalText> </approachSiteCode>	143
510	6.3.4.46.1.11 <doseQuantity><low value="" unit=""><high value="" unit=""> </doseQuantity>	144
	6.3.4.46.1.12 <low high value=""> <translation> <originalText><reference value=""></originalText> </translation></low high>	144
515	6.3.4.46.1.13 <rateQuantity><low value="" unit=""><high value="" unit=""></rateQuantity>	144
	6.3.4.46.1.14 <consumable>	144
	6.3.4.47 Nursing Assessments Battery 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4	144
	6.3.4.47.1 Specification	145
	6.3.4.47.1.1 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4'>	145
520	6.3.4.47.1.2 <organizer classCode='BATTERY' moodCode='EVN'>	145
	6.3.4.47.1.3 <id root=' ' extension=' '/>	145
	6.3.4.47.1.4 <code code='X-ASSESS' codeSystem='2.16.840.1.113883.6.1'>	145
	6.3.4.47.1.5 <author/><time/><assignedAuthor><id/></assignedAuthor></author>	146
525	6.3.4.47.1.6 <statusCode code='completed'>	146
	6.3.4.47.1.7 <component>	146
	6.3.4.48 Antenatal Testing and Surveillance Battery 1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10	146
	6.3.4.48.1 Specification	147
530	6.3.4.48.1.1 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10'>	147
	6.3.4.48.1.2 <organizer classCode='BATTERY' moodCode='EVN'>	147
	6.3.4.48.1.3 <id root=' ' extension=' '/>	147
535	6.3.4.48.1.4 <code code='XX- XX-ANTENATALTESTINGBATTERY' codeSystem='2.16.840.1.113883.6.1'>	147
	6.3.4.48.1.5 <author/><time/><assignedAuthor><id/></assignedAuthor></author>	148
540	6.3.4.48.1.6 <statusCode code='completed'>	148
	6.3.4.48.1.7 <component>	148

	6.3.4.49 Immunization Recommendation	148
	6.3.4.50 Alert Entry	148
	6.3.4.51 Antigen Dose	149
545	6.3.4.52 Intentionally blank	149
	6.3.4.53 Intentionally blank	149
	6.3.4.54 Observation Request 1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1	149
	6.3.4.54.1 Uses	149
	6.3.4.54.2 Specification	149
	6.3.4.54.2.1 <observation classCode='OBS' moodCode='INT PRP GOL'>	150
550	6.3.4.54.2.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1'>	150
	6.3.4.54.2.3 <templateId root='2.16.840.1.113883.10.20.1.25'>	150
	6.3.4.54.2.4 <id root=' ' extension=' '/>	150
	6.3.4.54.2.5 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' ' />	150
555	6.3.4.54.2.6 <text><reference value='#xxx'></text> -OR- <text>text</text> ..	150
	6.3.4.54.2.7 <statusCode code='active'>	151
	6.3.4.54.2.8 <effectiveTime value=' '/>	151
	6.3.4.54.2.9 <value xsi:type=' ' .../>	151
560	6.3.4.54.2.10 <methodCode code=' ' codeSystem=' ' codeSystemName=' ' />....	151
	6.3.4.54.2.11 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />	151
	6.3.4.54.2.12 <author><assignedAuthor	
	classCode='ASSIGNED'>...<assignedAuthor></author> ..	151
	6.3.4.55 Risk Indicators for Hearing Loss Entry 1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1 ...	151
565	6.3.4.55.1 Specification	152
	6.3.4.55.2 <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1'>	152
	6.3.4.55.3 <organizer classCode='BATTERY' moodCode='EVN'>	152
	6.3.4.55.4 <id root=' ' extension=' '/>	153
	6.3.4.55.5 <code code='58232-0' codeSystem='2.16.840.1.113883.6.1'>	153
570	6.3.4.55.6 <author/><time/><assignedAuthor><id/></assignedAuthor></author>	153
	6.3.4.55.7 <statusCode code='completed'>	153
	6.4.4.55.8 <component>	153
	6.3.4.56 Cancer Diagnosis Entry 1.3.6.1.4.1.19376.1.7.3.1.4.14.1	153
	6.3.4.56.1 Parent Template	153
575	6.3.4.56.2 Specification	154
	6.3.4.56.3 <act classCode='ACT' moodCode='EVN'>	156
	6.3.4.56.4 <templateId root='2.16.840.1.113883.10.20.1.27'> <templateId	
	root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2'>	156
580	6.3.4.56.5 <!-- 1..* entry relationships identifying problems of concern -->	
	<entryRelationship type='SUBJ'><observation classCode='OBS'	
	moodCode='EVN'><templateIDroot='1.3.6.1.4.1.19376.1.5.3.1.4.5'>...</	
	observation>	156
	6.3.4.56.6 <observation classCode="OBS" moodCode="EVN">	156
	6.3.4.56.7 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5'> <templateId	
	root='1.3.6.1.4.1.19376.1.7.3.1.4.14.1'>	156

585	6.3.4.56.8 <code code="282291009" codeSystem=" 2.16.840.1.113883.6.96" codeSystemName="SNOMED CT" displayName="Diagnosis"/> 157
	6.3.4.56.9 <statusCode code='completed' /> 157
	6.3.4.56.10 <effectiveTime value="xxx" /> 157
	6.3.4.56.11 <value xsi:type='CD' code=' ' codeSystem=' ' codeSystemName=' ' displayName=' '> 157
590	6.3.4.56.12 <qualifier><name code="31206-6" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName=" Behavior ICD-O-3 Cancer"/><value code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> 157
595	6.3.4.56.13 <qualifier><name code="21861-0" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Dx confirmed by Cancer"/><value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> 157
600	6.3.4.56.14 <targetSiteCode code=" " codeSystem="" codeSystemName=" " displayName=" "> 158
	6.3.4.56.15 <qualifier><name code="20228-3" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Anatomic part Laterality"/><value code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> 158
605	6.3.4.56.16 <entryRelationship typeCode="SUBJ" inversionInd="false"/> 158
	6.3.4.56.17 <observation classCode="OBS" moodCode="EVN"> <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.14.2"/> [1 st nesting] 158
	6.3.4.56.18 <code code="75620-5" displayName="TNM Clinical Stage Information" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> [1 st nesting] 158
610	6.3.4.56.19 <statusCode code="completed"/> [1 st nesting] 158
	6.3.4.56.20 <value xsi:type="CD" code="" codeSystem="" codeSystemName="" displayName=" "> [1 st nesting] 158
615	6.3.4.56.21 <qualifier><name code="21909-7" displayName=" Descriptor.clinical Cancer" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/><value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> [1 st nesting] 159
620	6.3.4.56.22 <qualifier><name code="21917-0" displayName="Version TNM Classification" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/><value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> [1 st nesting] 159
625	6.3.4.56.23 <participant typeCode="PPRF"> <participantRole> <code code="21910-5" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Stager.clinical Cancer"/><playingEntity nullFlavor="NA"> <code xsi:type="CE"

	code="" codeSystem="" codeSystemName=" " displayName=" "/> [1 st nesting]	159
630	6.3.4.56.24 <!-- 0..3 entryRelationships identifying simple observations for TNM Clinic Tumor, TNM Clinical Nodes, and TNM Clinical Metastases--><entryRelationship typeCode="COMP" inversionInd="false"><observation classCode='OBS'moodCode='EVN'><templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>...</observation>[2nd nesting]	159
635	6.3.4.56.25 <code code="" displayName=" " codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> [2 nd nesting].....	160
	6.3.4.56.26 <value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/>.....	160
640	6.3.4.57 Patient Transfer 1.3.6.1.4.1.19376.1.5.3.1.25.1.4.1	160
	6.3.4.57.1 Parent Template	161
	6.3.4.57.2 Specification	161
	6.3.4.57.3 <act classCode='ACT' moodCode='INT EVN'>	161
	6.3.4.57.4 <templateId root='TBD'>	161
645	6.3.4.57.5 <id root=" extension="/">	161
	6.3.4.57.6 <code code=" displayName=" codeSystem=" codeSystemName=" />...	161
	6.3.4.57.7 <text><reference value='#xxx'></text>	161
	6.3.4.57.8 statusCode.....	161
	6.3.4.57.9 <effectiveTime><low value="/"><high value="/"><effectiveTime/>	162
650	6.3.4.57.10 participant	162
	6.3.4.57.11 templateId	162
	6.3.4.57.12 participantRole.....	162
	6.3.4.57.13 <id root=" extension="/">	162
	6.3.4.57.14 <code>	162
655	6.3.4.57.15 <addr></addr>	162
	6.3.4.57.16 <telecom>	162
	6.3.4.57.17 playingEntity.....	162
	6.3.4.57.18 name.....	163
	6.3.4.58 Death Pronouncement Entry Content Module (1.3.6.1.4.1.19376.1.7.3.1.4.23.1)	163
660	6.3.4.59 Death Location Type Entry Content Module	164
	6.3.4.64 History of Employment Status Observation.....	165
	6.3.4.65 Usual Occupation Observation Entry	167
	6.3.4.66 Past or Present Occupation Observation Entry	169
665	6.3.4.67 Work Schedule Observation Entry	172
	6.3.4.68 Weekly Work Hours Observation Entry	174
	6.3.4.69 Usual Occupation Duration Entry	175
	6.3.4.71 Pregnancy Status Review Organizer (1.3.6.1.4.1.19376.1.5.3.1.4.22).....	176
	6.3.4.71.1 Specification	176
670	6.3.4.71.2 <organizer classCode='CLUSTER' moodCode='EVN'>.....	176
	6.3.4.71.3 <templateId root="/">	177

	6.3.4.71.4 <id root=' ' extension=' '/>	177
	6.3.4.71.5 <code code='118185001' displayName='Pregnancy Observations' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED-CT' />	177
675	6.3.4.71.6 <statusCode code='completed' />	177
	6.3.4.71.7 <effectiveTime value=' '/>	177
	6.3.4.71.8 <author typeCode='AUT'><assignedEntity1 typeCode='ASSIGNED'>...</assignedEntity1></author>	177
680	6.3.4.71.9 <component typeCode='COMP' />	177
	6.3.4.72 Pregnancy Status Review Observation (1.3.6.1.4.1.19376.1.5.3.1.4.22.1)	177
	6.3.4.72.1 Parent Template	177
	6.3.4.72.1.1 Uses	177
	6.3.4.72.2 Specification	178
685	6.3.4.72.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.22.1' /> <templateId root=' '/>	178
	6.3.4.72.4 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>	178
690	6.3.4.72.5 <repeatNumber value=' '/>	178
	6.3.4.72.6 <value xsi:type=' ' ... />	178
	6.3.4.72.7 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/> <methodCode code=' ' codeSystem=' ' codeSystemName=' '/> <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>	178
	6.3.4.73 Performer	179
695	6.3.4.73.1 <performer typeCode="PRF" />	179
	6.3.4.73.2 <templateId root="1.3.6.1.4.1.19376.1.5.3.1.1.24.3.5" />	179
	6.3.4.73.3 <assignedEntity classCode="ASSIGNED" />	179
	6.3.4.73.4 <id root="" extension="" />	179
700	6.3.4.73.5 <addr></addr>	180
	6.3.4.73.6 <telecom></telecom>	180
	6.3.4.73.7 <assignedPerson>	180
	6.3.4.73.8 <name></name>	180
	6.3.4.73.9 <representedOrganization>	180
705	6.3.4.73.10 <id root='...' extension='...' />	180
	6.3.4.73.11 <name></name>	181
	6.3.4.73.12 <addr></addr>	181
	6.3.4.73.13 <telecom></telecom>	181
	6.3.4.74 Weekly Work Days Observation Entry	181
	6.4 HL7 Version 2.0 Content Modules	182
710	6.5 PCC Value Sets	182
	6.5.A Antepartum History of Past Illness Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.1.. 183	183
	6.5.C Antepartum Family History and Genetic Screening Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.4	184
	6.5.D Antepartum Review of Systems Menstrual History Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.5	185

	6.5.E Antepartum History of Infection Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.6	185
	6.5.F Antepartum Laboratory Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.7	186
	6.5.G Antepartum Education Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.8	189
	6.5.H JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set	191
720	6.5.H.1 Metadata	191
	6.5.H.2 JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set	191
	6.5.I JCIH-EHDI Risk Indicators for Hearing Loss Codes	193
	6.5.I.1 Metadata	193
725	6.5.I.2 JCIH-EHDI Risk Indicators for Hearing Loss Value Set	194
	6.5.I.3 Pending Codes for SNOMED-CT Findings/Situation to support Risk Indicators for Hearing Loss	194
	6.5.J JCIH-EHDI Risk Indicators for Hearing Loss - Procedures Codes	195
	6.5.J.1 Metadata	195
730	6.5.J.2 JCIH-EHDI Risk Indicators for Hearing Loss - Procedures Value	195
	6.5.K Newborn Hearing Procedure Codes	196
	6.5.K.1 Metadata	196
	6.5.K.2 JCIH-EHDI Newborn Hearing Procedure Value Set	197
	6.5.L JCIH-EHDI Newborn Hearing Screening Method Codes	197
	6.5.L.1 Metadata	197
	6.5.L.2 JCIH-EHDI Newborn Hearing Screening Method Value Set	198
	6.5.M JCIH-EHDI Hearing Screen Right Codes– Right	198
	6.5.M.1 Metadata	198
	6.5.M.2 JCIH-EHDI Hearing Screen Right Value Set	199
740	6.5.N JCIH-EHDI Hearing Screen Left Codes	199
	6.5.N.1 Metadata	199
	6.5.N.2 JCIH-EHDI Hearing Screen Left Value Set	200
	6.5.O JCIH-EHDI Reason for no Hearing Loss Diagnosis or Screening Codes(SNOMED)	200
745	6.5.O.1 Metadata	200
	6.5.O.2 JCIH-EHDI Reason for no Hearing Loss Diagnosis or Screening Value Set	201
	6.5.P JCIH-EHDI Newborn Hearing Loss Referrals Codes	202
	6.5.P.1 Metadata	202
	6.5.P.2 JCIH-EHDI Newborn Hearing Loss Referrals Value Set	203
750	6.5.Q JCIH-EHDI Newborn Hearing Loss Reason for no Follow-up – Patient Reason Codes	203
	6.5.Q.1 Metadata	203
	6.5.Q2 JCIH-EHDI Newborn Hearing Loss Reason for no Follow-up – Patient Reason Value Set	204
	6.5.R Joint Commission Medical Reason Codes	206
	6.5.R.1 Metadata	206
	6.5.R.2 Joint Commission Medical Reason Value Set	206
	6.5.S JCIH-EHDI Inpatient Screening Results not Performed Codes	207
	6.5.S.1 Metadata	207

760	6.5.S.2 JCIH-EHDI Inpatient Screening Results not Performed Value Set	208
	6.5.T JCIH-EHDI Evidence of Hearing Screening Performed Codes	208
	6.5.T.1 Metadata	208
	6.5.T.2 JCIH-EHDI Evidence of Hearing Screening Performed Value Set	209
765	6.5.U JCIH-EHDI Procedure Declined Value Set Codes	209
	6.5.U.1 Metadata	209
	6.5.U.2 JCIH-EHDI Procedure Declined Value Set Value Set	210
770	6.5.V JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set Codes	210
	6.5.V.1 Metadata	210
	6.5.V.2 JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set	211
775	6.5.W Primary Site Value Set	212
	6.5.X Histologic Type Value Set	212
	6.5.Y Derived AJCC Descriptor (T,N,M) Value Set	212
	6.5.Z TNM Edition Value Set	212
	6.5.AA TNM Stage Group Value Set	213
780	6.5.BB TNM Stage Descriptor Value Set	214
	6.5.CC TNM Tumor Value Set	214
	6.5.DD TNM Node Value Set	215
	6.5.EE TNM Metastasis Value Set	216
	6.5.FF QRPH VRDR Autopsy Procedure Performed Codes	216
	6.5.FF.1 Metadata	216
	6.5.FF.2 VRDR Autopsy Procedure Performed Value Set	217
785	6.5.GG QRPH VRDR Autopsy Not Performed Codes	219
	6.5.GG.1 Metadata	219
	6.5.GG.2 VRDR Autopsy Not Performed Value Set	220
790	6.5.HH VRDR Discharge Death Codes	220
	6.5.HH.1 Metadata	220
	6.5.HH.2 VRDR Discharge DeathValue Set	221
	6.5.II VRDR Death Location Type Codes	221
	6.5.II.1 Metadata	221
	6.5.II.2 VRDR Death Location Type Value Set	222
795	6.5.JJ VRDR Death Certification Procedure Codes	223
	6.5.JJ.1 Metadata	223
	6.5.JJ.2 VRDR Death Certification Procedure Performed Value Set	224
	6.5.KK VRDR Death Pronouncement Procedure Codes	224
	6.5.KK.1 Metadata	224
	6.5.KK.2 VRDR Death Pronouncement Procedure Performed Value Set	225
	6.6 Concept Domains	225

Introduction

800 This supplement is written for trial implementation. It is written as changes to the latest revision of the documents listed below. The reader should have already read and understood these documents:

1. [PCC Technical Framework Volume 1](#)
2. [PCC Technical Framework Volume 2](#)

805 This supplement also references other documents¹. The reader should have already read and understood these documents:

1. [IT Infrastructure Technical Framework Volume 1](#)
2. [IT Infrastructure Technical Framework Volume 2](#)
3. [IT Infrastructure Technical Framework Volume 3](#)
4. HL7®² and other standards documents referenced in Volume 1 and Volume 2

This supplement defines a number of PCC and QRPH content modules that are shared between various content documents. These are provided for trial implementation and will be published in the same format for trial implementation. Upon completion, some content modules will be moved to final text; others may remain in trial implementation.

815 Profile Abstract

This supplement does not describe a profile.

Open Issues and Questions

None

Closed Issues

820 None

¹ The first three documents can be located on the IHE Website at http://ihe.net/Technical_Frameworks/#IT. The remaining document can be obtained from its respective publisher.

² HL7 is the registered trademark of Health Level Seven International.

Volume 1 – Integration Profiles

None

Glossary

Add the following terms to the Glossary:

825 None

2.5 History of Annual Changes

Add the following bullet to the end of the bullet list in Section 2.5

- 830
- Added a set of CDA®³ Content Modules shared across several Integration Profiles for the 2010-2011 documentation cycle.
 - In the 2011-2012 documentation cycle, the following CDA Section Content Modules were added as well as various Entry Content Modules and Value Sets:
 - PCC Transport Summary Profiles supplement
 - Sending Facility
 - Receiving Facility
 - Mass Causality Incident
 - Unit Response Level
 - Protocols Used
 - Extra Attendants Information
 - Invasive Airway
 - Isolation Status
 - Restraints
 - Ventilator Usage
 - Provider Level
- 835
- QRPH EHCP Profile
 - Risk Indicators for Hearing Loss
 - Hearing Screening Coded Results
- 840
- QRPH PRPH-Ca Profile
 - Cancer Diagnosis
- 845

³ CDA is the registered trademark of Health Level Seven International.

- 850 • In the 2012-2013 documentation cycle, edits were made based on CPs. In addition, the following content modules were added:
- QRPH VRDR Section Content Modules
 - 6.3.3.10.1 VRDR Death Report Section
 - 6.3.3.10.2 Coded Hospital Course Section
- 855 • QRPH VRDR Entry Content Modules were added
 - 6.3.3.4.58 Death Pronouncement Entry Content Module
 - 6.3.3.4.59 Death Location Type Entry Content Module
- 860 • Some QRPH VRDR value sets were added
- 865 • QRPH HW Section Content Modules
 - 6.3.3.10.3 Resources to Support Goals Section
 - 6.3.3.10.4 Healthy Weight Care Plan Section
 - 6.3.3.10.5 Occupational Data for Health Section
- 870 • QRPH HW Entry Content Modules
 - 6.3.4.60 Occupational Data For Health Organizer
 - 6.3.4.61 Employment Status Organizer
 - 6.3.4.62 Usual Occupation and Industry Organizer
 - 6.3.4.63 History of Occupation Organizer
 - 6.3.4.64 Employment Status Observation
 - 6.3.4.65 Usual Occupation and Industry Observation Entry
 - 6.3.4.66 Occupation Observation Entry
 - 6.3.4.67 Work Schedule Observation Entry
 - 6.3.4.68 Weekly Work Hours Observation Entry
 - 6.3.4.69 Usual Occupation Duration Entry
 - 6.3.4.70 Usual Industry Duration Entry
- 875 • In the 2013-2014 documentation cycle, edits were made based on CPs.
- In the 2015-2016 documentation cycle, edits were made based on CPs. In addition, the following Volume 2 section was added to this document:
- 6.6 Concept Domains

880

- In the June 2020 publication, the Occupational Data for Health structures were updated to align with changes made in the Healthy Weight supplement (Rev. 2.3)

Volume 2 – Transactions and Content Modules

885 Please note that in December of 2012, a new supplement template was released. The new template separates *Transactions* (Volume 2) and *Content Modules* (Volume 3). As a result, in newer supplements you will find content module definitions in volume 3. The section numbering scheme; however, remains the same.

Add Section 6.1

6.1 Conventions

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

R

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

R2

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

905 O

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

C

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

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

<i>Add Section 6.2</i>

6.2 Folder Content Modules

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

6.2.1 EDES Folder Specification

This section intentionally left blank.

925 **6.2.2 APR Folder Specification**

This section intentionally left blank.

6.2.3 LDR Folder Specification

This section intentionally left blank.

6.3 HL7 Version 3.0 Content Modules

- 930 This section contains content modules based upon the HL7 CDA Release 3.0 Standard, and related standards and/or implementation guides.

6.3.1 CDA Document Content Modules

<i>Add Section 6.3.1.x</i>

6.3.1.x History and Physical Specification 1.3.6.1.4.1.19376.1.5.3.1.1.16.1.4

- 935 The History and Physical document content module is a Medical Summary and inherits all header constraints from Medical Summary (1.3.6.1.4.1.19376.1.5.3.1.1.2). The intention of this document content module is to provide a base from which other document content modules may be derived. Future work may also result in a content profile for History and Physical.

6.3.1.x.1 Format Code

- 940 The XDSDocumentEntry format code for this content is **urn:ihe:pcc:hp:2008**

6.3.1.x.2 LOINC Code

The LOINC code for this document is **34117-2 HISTORY AND PHYSICAL**

6.3.1.x.3 Standards

CDAR2	HL7 CDA Release 2.0
CDTHP	CDA for Common Document Types History and Physical Notes (DSTU)

6.3.1.x.4 Specification

945 This section references content modules using Template ID as the key identifier. Definitions of the modules are found in either:

- IHE Patient Care Coordination Volume 2: Final Text
- IHE PCC CDA Content Modules Supplement (this document, for Trial Implementation)

Table 6.3.1.x.4-1: History and Physical Data Elements

Data Element Name	Opt	Template ID
Chief Complaint	R	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1
History of Present Illness	R	1.3.6.1.4.1.19376.1.5.3.1.3.4
History of Past Illness	R	1.3.6.1.4.1.19376.1.5.3.1.3.8
Medications	R	1.3.6.1.4.1.19376.1.5.3.1.3.19
Allergies and Other Adverse Reactions Section	R	1.3.6.1.4.1.19376.1.5.3.1.3.13
Social History	R	1.3.6.1.4.1.19376.1.5.3.1.3.16
Family History	R	1.3.6.1.4.1.19376.1.5.3.1.3.14
Review of Systems	R	1.3.6.1.4.1.19376.1.5.3.1.3.18
Detailed Physical Examination This section SHALL include Vital Signs (1.3.6.1.4.1.19376.1.5.3.1.3.25) as a subsection.	R	1.3.6.1.4.1.19376.1.5.3.1.1.9.15
Results Diagnostic Findings; use this OR Coded Results	R	1.3.6.1.4.1.19376.1.5.3.1.3.27
Coded Results Diagnostic Findings; use this OR Results	R	1.3.6.1.4.1.19376.1.5.3.1.3.28
Assessment and Plan	R	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5

950

6.3.1.x.5 Conformance

955 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 Summaries](#) content module, and so must conform to the requirements of that template as well, thus all <templateId> elements shown in the example below shall be included.

```

<ClinicalDocument xmlns='urn:hl7-org:v3'>
  <typeId extension="POCD_HD000040" root="2.16.840.1.113883.1.3"/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.2'/>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.16.1.4'/>
  <id root=' ' extension=' ' />
  <code code='34117-2' displayName='HISTORY AND PHYSICAL'
    codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
  <title>History and Physical</title>
  <effectiveTime value='20080601012005' />
  <confidentialityCode code='N' displayName='Normal'
    codeSystem='2.16.840.1.113883.5.25' codeSystemName='Confidentiality' />
  <languageCode code='en-US' />
  :
  <component><structuredBody>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1' />
        <!-- Required Chief Complaint Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.4' />
        <!-- Required History of Present Illness Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.8' />
        <!-- Required History of Past Illness Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.19' />
        <!-- Required Medications Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.13' />
        <!-- Required Allergies and Other Adverse Reactions Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.16' />
        <!-- Required Social History Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.14' />
        <!-- Required Family History Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18' />
        <!-- Required Review of Systems Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.15' />
        <!-- Required Detailed Physical Examination Section content -->
      </section>
    </component>
  </structuredBody>
</component>

```

```

1025   </component>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.27' />
    <!-- Required Results Section content -->
  </section>
</component>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.28' />
    <!-- Required Coded Results Section content -->
  </section>
</component>
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.5' />
    <!-- Required Assessment and Plan Section content -->
  </section>
</component>
</structuredBody></component>
</ClinicalDocument>

```

Figure 6.3.1.x.5-1: Sample History and Physical Document*Add Section 6.3.2***6.3.2 CDA Header Content Modules***Add Section 6.3.2.1***1050 6.3.2.1 Language Communication 1.3.6.1.4.1.19376.1.5.3.1.2.1***Add Section 6.3.2.2***6.3.2.2 Employer and School Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.2***Add Section 6.3.2.3***6.3.2.3 Healthcare Providers and Pharmacies 1.3.6.1.4.1.19376.1.5.3.1.2.3***Add Section 6.3.2.4***6.3.2.4 Patient Contacts 1.3.6.1.4.1.19376.1.5.3.1.2.4***Add Section 6.3.2.5***6.3.2.5 Spouse 1.3.6.1.4.1.19376.1.5.3.1.2.4.1**

The spouse header element records the spouse of a patient, and inherits other constraints from the [Patient Contacts](#) entry. Items in bold in the example below show the additional constraints on this element.

This element SHALL be included as a participant in the header of the CDA document in the event of the pregnancy. If this does not apply to the patient this element SHALL use a null flavor.

1065 **6.3.2.5.1 Parent Template**

The parent of this template is [Patient Contacts](#).

6.3.2.5.2 Specification

```
<participant typeCode='IND'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4' />
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.1' />
  <time value='20070213' />
  <associatedEntity classCode='PRS'>
    <code code='xx-spouse|184142008' displayName=' ' codeSystem='2.16.840.1.113883.6.96'
codeSystemName='SNOMED CT' />
    <addr></addr>
    <telecom value=' ' use=' ' />
    <assignedPerson><name></name></assignedPerson>
  </associatedEntity>
</participant>
```

1070

1075

1080

6.3.2.5.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4' /><templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.1' />

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

1085

```
<rule context='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4.1"]'>
  <assert test='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4"'>
    A participant using template 1.3.6.1.4.1.19376.1.5.3.1.2.4.1 must also use template
    1.3.6.1.4.1.19376.1.5.3.1.2.4.
  </assert>
</rule>
```

1090

6.3.2.5.4 <associatedEntity classCode='PRS'>

The classCode attribute of the <associatedEntity> element shall be PRS.

1095

```
<rule context='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4.1"]'>
  <assert test='..../h17:associatedEntity/@classCode = "PRS"'>
    The classCode attribute of the associated entity shall be PRS.
  </assert>
</rule>
```

6.3.2.5.5 <code code='127848009|184142008' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />

1100

This element SHALL use 127848009 to represent the patient's spouse or 184142008 to represent the patient's next of kin. The code system name is SNOMED CT.

6.3.2.5.6 Completed Example

```

<!-- Husband/Domestic Partner -->
<participant typeCode="IND">
    <associatedEntity classCode="NOK">
        <code code="184142008" displayName="patient's next of kin"
            codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"/>
        <addr>
            <streetAddressLine>45 Chunn Dr.</streetAddressLine>
            <city>Spring Hill</city>
            <state>TN</state>
            <postalCode>37174</postalCode>
            <country>USA</country>
        </addr>
        <telecom value="tel: (999) 555-1212" use="WP"/>
        <associatedPerson>
            <name>
                <prefix>Mr.</prefix>
                <given>John</given>
                <family>Youngston</family>
            </name>
        </associatedPerson>
    </associatedEntity>
</participant>

```

1125 Add Section 6.3.2.6

6.3.2.6 Natural Father of Fetus 1.3.6.1.4.1.19376.1.5.3.1.2.4.2

This header element records the natural father of the fetus, and inherits other constraints from the Patient Contacts (1.3.6.1.4.1.19376.1.5.3.1.2.4) entry. Items in bold in the example below show the additional constraints on this element.

1130 This element SHALL be included as a participant in the header of the CDA document in the event of the pregnancy. If the father of the baby is unknown this element SHALL use a null flavor.

6.3.2.6.1 Parent Template

The parent of this template is Patient Contacts (1.3.6.1.4.1.19376.1.5.3.1.2.4).

1135 **6.3.2.6.2 Specification**

```

<participant typeCode='IND'>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.1' />
<time value='20070213' />
<associatedEntity classCode='PRS'>
    <code code='xx-fatherofbaby' displayName=' ' codeSystem='2.16.840.1.113883.6.96'
        codeSystemName='SNOMED CT' />
    <addr></addr>
    <telecom value=' ' use=' ' />
    <assignedPerson><name></name></assignedPerson>
</associatedEntity>
</participant>

```

6.3.2.6.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4'><templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.4.2'>

1150 The <templateId> element identifies this person as the natural father and must be recorded exactly as shown above.

```
<rule context='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4.2"]'>
  <assert test='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4"'>
    A participant using template 1.3.6.1.4.1.19376.1.5.3.1.2.4.2 must also use template
    1.3.6.1.4.1.19376.1.5.3.1.2.4.
  </assert>
</rule>
```

6.3.2.6.4 <associatedEntity classCode='PRS'>

1160 The classCode attribute of the <associatedEntity> element SHALL be PRS.

```
<rule context='h17:templateId/@root="1.3.6.1.4.1.19376.1.5.3.1.2.4.2"]'>
  <assert test='..../h17:associatedEntity/@classCode = "PRS"'>
    The classCode attribute of the associated entity shall be PRS.
  </assert>
</rule>
```

6.3.2.6.5 <code code='xx-fatherofbaby' displayName=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>

For father of baby the code SHALL be xx-fatherofbaby (requested). The code system name is SNOMED CT.

1170 **6.3.2.6.6 Completed Example**

```
<!-- Father of baby -->
<participant typeCode="IND">
  <associatedEntity classCode="NOK">
    <code code="xx-fatherofbaby" displayName="Father of Baby"
      codeSystem="2.16.840.1.113883.6.96" codeSystemName="SNOMED CT"/>
    <addr>
      <streetAddressLine>18 Oak Valley Dr.</streetAddressLine>
      <city>Monteagle</city>
      <state>TN</state>
      <postalCode>37205</postalCode>
      <country>USA</country>
    </addr>
    <telecom value="tel: (999) 555-1212" use="WP"/>
    <associatedPerson>
      <name>
        <prefix>Mr.</prefix>
        <given>Thomas</given>
        <family>Caster</family>
      </name>
    </associatedPerson>
  </associatedEntity>
</participant>
```

<i>Add Section 6.3.2.7</i>

1195 **6.3.2.7 Authorization 1.3.6.1.4.1.19376.1.5.3.1.2.5**

1200 Each `<authorization>` element in the CDA Header represents an informed consent. When the document being shared represents the informed consent to a policy expressed by the XDS Affinity Domain within the document, it shall do so in an `<authorization>` element. More than one `<authorization>` element may be present. The consent to share information shall have a unique identifier contained in the `<id>` element, representing the patient consent to that policy. The policy being consented to shall be represented in the `<code>` element. Note that other `<authorization>` elements may be present representing other sorts of consents associated with the document.

6.3.2.7.1 Parent Template

1205 None

6.3.2.7.2 Specification

```

<authorization typeCode='AUTH'>
  <consent classCode='CONS' moodCode='EVN'>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.5' />
    <id root=''/>
    <code code='' codeSystem='' codeSystemName='' displayName='' />
    <statusCode code='completed' />
  </consent>
</authorization>

```

6.3.2.7.3 <authorization typeCode='AUTH'>

1220 At least one `<authorization>` element must be present in a consent medical document in documents shared by Document Source Actors that implement the privacy option. The `typeCode` attribute shall be present and be valued with AUTH, indicating that this is an authorization act related to the document.

6.3.2.7.4 <consent classCode='CONS' moodCode='EVN'>

1225 Each authorization element shall have one `<consent>` element. The `classCode` shall be present and be valued with CONS, indicating that the related act is an informed consent. The `moodCode` shall be EVN, indicating that this element represents an act that has occurred.

6.3.2.7.5 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.2.5' />

The `<templateId>` element shall be recorded as shown above and identifies this consent as an authorization entry.

6.3.2.7.6 <id root=' '/>

1230 The `<consent>` element shall have one identifier that is used to uniquely identify the consent act. This identifier shall contain a root attribute, and shall not contain an extension attribute.

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

The <consent> element shall have one <code> element that is used to identify the consent policy that was agreed to by the patient.

1235

Add Section 6.3.3

6.3.3 CDA Section Content Modules

Add Section 6.3.3.1

6.3.3.1 Reasons for Care

Add Section 6.3.3.1.1

1240

6.3.3.1.1 Reason for Referral

Add Section 6.3.3.1.2

6.3.3.1.2 Coded Reason for Referral

Add Section 6.3.3.1.3

6.3.3.1.3 Chief Complaint

1245

Add Section 6.3.3.1.4

6.3.3.1.4 Hospital Admission Diagnosis

Add Section 6.3.3.1.5

6.3.3.1.5 Proposed Procedure Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.1	
General Description	The proposed procedure section shall contain a description of the procedures for which a risk assessment is required including procedure names and codes, patient position, dates, and names of surgeons. It shall include entries for procedures as described in the Entry Content Modules and the required and optional subsections.	
LOINC Code	Opt	Description
29554-3	R	PROCEDURE
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.19	R	Procedure Entry
Subsections	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.9.4	R	Reason for Procedure
1.3.6.1.4.1.19376.1.5.3.1.1.9.3	R	Proposed Anesthesia
1.3.6.1.4.1.19376.1.5.3.1.1.9.2	R	Estimated Blood Loss

1.3.6.1.4.1.19376.1.5.3.1.1.9.40	R	Procedure Care Plan
----------------------------------	---	---------------------

```

1250 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.1' />
        <id root=' ' extension=' ' />
        <code code='29554-3' displayName='PROCEDURE'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
        <entry>
            :
            <!-- Required Procedure Entry element -->
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19' />
            :
        </entry>
        <component>
            <section>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.4' />
                <!-- Required Reason for Procedure Section content -->
            </section>
        </component>
        <component>
            <section>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.3' />
                <!-- Required Proposed Anesthesia Section content -->
            </section>
        </component>
        <component>
            <section>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.2' />
                <!-- Required if known Estimated Blood Loss Section content -->
            </section>
        </component>
        <component>
            <section>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.40' />
                <!-- Required if known Procedure Care Plan Section content -->
            </section>
        </component>
    </section>
</component>

```

Figure 6.3.3.1.5-1: Specification for Proposed Procedure Section**Add Section 6.3.3.1.6****6.3.3.1.6 EBS Estimated Blood Loss Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.2**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.2	
General Description	The estimated blood loss section shall contain a description of the blood loss for the procedure.	
LOINC Code	Opt	Description
8717-1	R	OPERATIVE NOTE ESTIMATED BLOOD LOSS

Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.13	R	Simple Observation

1295

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.2'>
    <id root=' ' extension=' '/>
    <code code='8717-1' displayName='OPERATIVE NOTE ESTIMATED BLOOD LOSS'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required Simple Observation element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>
      :
    </entry>
  </section>
</component>

```

1300

1305

1310

Figure 6.3.3.1.6-1: EBS Specification for Estimated Blood Loss Section

1315

*Add Section 6.3.3.1.7***6.3.3.1.7 Proposed Anesthesia Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.3**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.3	
General Description	The proposed anesthesia section shall contain a description of the anesthetic techniques for which a risk assessment is required. It shall include entries for anesthetic procedures as described in the Entry Content Modules.	
LOINC Code	Opt	Description
10213-7	R	Surgical operation note anesthesia
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.19	R	Procedure Entry The procedure entries shall be in INT mood.

```

1320 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.3' />
<id root=' ' extension=' ' />
<code code='10213-7' displayName='Surgical operation note anesthesia'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
<!-- Required Procedure Entry element -->
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19' />
  :
</entry>
</section>
</component>

```

Figure 6.3.3.1.7-1: Specification for Anesthesia Administered Section

<i>Add Section 6.3.3.1.8</i>

6.3.3.1.8 Reason for Procedure Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.4

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.4	
General Description	The reason for procedure section shall contain a description of the reason that the patient is receiving the procedure. It shall include entries for conditions as described in the Entry Content Module.	
LOINC Code	Opt	Description
10217-8	R	OPERATIVE NOTE INDICATIONS
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.5	R2	Problem Entry

```

1340 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.4' />
<id root=' ' extension=' ' />
<code code='10217-8' displayName='OPERATIVE NOTE INDICATIONS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
<!-- Required if known Problem Entry element -->
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5' />
  :
</entry>
</section>
</component>

```

Figure 6.3.3.1.8-1: Specification for Reason for Procedure Section

<i>Add Section 6.3.3.1.9</i>

6.3.3.1.9 Reason for Visit Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1.1	
General Description	This contains a narrative description of the patient's reason for visit.	
LOINC Code	Opt	Description
29299-5	R	REASON FOR VISIT

1360

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.1.1' />
    <id root=' ' extension=' ' />
    <code code='29299-5' displayName='REASON FOR VISIT'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

1365

1370

Figure 6.3.3.1.9-1: Specification for Reason for Visit Section

1375

*Add Section 6.3.3.1.10***6.3.3.1.10 Injury Incident Description Section 1.3.6.1.4.1.19376.1.5.3.1.1.19.2.1**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.19.2.1	
General Description	This section shall include a description of the incident leading to the injury, including status of relevant safety equipment in use (e.g., safety belts, air bag, helmet).	
LOINC Code	Opt	Description
11374-6	R	Injury incident description

1380

1385

1390

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.19.2.1' />
    <id root=' ' extension=' ' />
    <code code='11374-6' displayName='Injury incident description'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

Figure 6.3.3.1.10-1: Sample Injury Incident Description Section*Add Section 6.3.3.2*

6.3.3.2 Other Condition Histories

Add Section 6.3.3.2.1

1395 **6.3.3.2.1 History of Present Illness**

Add Section 6.3.3.2.2

6.3.3.2.2 Hospital Course

Add Section 6.3.3.2.3

6.3.3.2.3 Active Problems

1400 *Add Section 6.3.3.2.4*

6.3.3.2.4 Discharge Diagnosis

Add Section 6.3.3.2.5

6.3.3.2.5 History of Past Illness

Add Section 6.3.3.2.6

1405 **6.3.3.2.6 Encounter Histories**

Add Section 6.3.3.2.7

6.3.3.2.7 History of Outpatient Visits

Add Section 6.3.3.2.8

6.3.3.2.8 History of Inpatient Visits

1410 *Add Section 6.3.3.2.9*

6.3.3.2.9 List of Surgeries

Add Section 6.3.3.2.10

6.3.3.2.10 Coded List of Surgeries

Add Section 6.3.3.2.11

1415 **6.3.3.2.11 Allergies and Other Adverse Reactions**

Add Section 6.3.3.2.12

6.3.3.2.12 Family medical History

Add Section 6.3.3.2.13

6.3.3.2.13 Coded Family Medical History

1420 *Add Section 6.3.3.2.14*

6.3.3.2.14 Social History Section

Add Section 6.3.3.2.15

6.3.3.2.15 Functional Status

Add Section 6.3.3.2.16

1425 **6.3.3.2.16 Review of Systems**

Add Section 6.3.3.2.17

6.3.3.2.17 Hazardous Working Conditions

Add Section 6.3.3.2.18

6.3.3.2.18 Pregnancy History

1430 *Add Section 6.3.3.2.19*

6.3.3.2.19 Medical Devices

Add Section 6.3.3.2.20

6.3.3.2.20 Foreign Travel

Add Section 6.3.3.2.21

1435 **6.3.3.2.21 Pre-procedure Family Medical History Section**

1.3.6.1.4.1.19376.1.5.3.1.1.9.5 (Deprecated)

Add Section 6.3.3.2.22

6.3.3.2.22 Coded Functional Status Assessment Section

1.3.6.1.4.1.19376.1.5.3.1.1.12.2.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.12.2.1	
Parent Template	Functional Status (1.3.6.1.4.1.19376.1.5.3.1.3.17, see PCC TF-2: 6.3.3.2.15)	
General Description	<p>The coded functional status assessment section provides a machine readable and narrative description of the patient's status of normal functioning at the time the document was created.</p> <p>Functional status includes information concerning:</p> <ul style="list-style-type: none"> Ambulatory ability Mental status or competency Activities of Daily Living (ADL's) including bathing, dressing, feeding, grooming Home/living situation having an effect on the health status of the patient Ability to care for self Social activity, including issues with social cognition, participation with friends and acquaintances other than family members Occupation activity, including activities partly or directly related to working, housework or volunteering, family and home responsibilities or activities related to home and family Communication ability, including issues with speech, writing or cognition required for communication Perception, including sight, hearing, taste, skin sensation, kinesthetic sense, proprioception, or balance 	
LOINC Code	Opt	Description
47420-5	R	Functional Status Assessment
Subsections	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2	R	Pain Scale Assessment
1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3	O Note 1	Braden Score Assessment
1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4	O Note 1	Geriatric Depression Scale
1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5	O Note 1	Minimum Data Set

1440

Note 1: At least one of the above optional subsections shall be present

6.3.3.2.22.1 Standards

CDAR2 [HL7 CDA Release 2.0](#)

CRS [HL7 Care Record Summary](#)

CCD [ASTM/HL7 Continuity of Care Document](#)

LOINC [Logical Observation Identifier Names and Codes](#)

SNOMED [Systematized Nomenclature of Medicine Clinical Terminology](#)

6.3.3.2.22.2 Parent Template

The parent of this template is Functional Status (see PCC TF-2: 6.3.3.2.15).

```

1445 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.17' />
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.1' />
        <id root=' ' extension=' ' />
        <code code='47420-5' displayName='Functional Status Assessment'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
    </section>
</component>
1450 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2' />
        <!-- Required Pain Scale Assessment Section content -->
    </section>
</component>
1455 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3' />
        <!-- Optional Braden Score Assessment Section content -->
    </section>
</component>
1460 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4' />
        <!-- Optional Geriatric Depression Scale Section content -->
    </section>
</component>
1465 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5' />
        <!-- Optional Minimum Data Set Section content -->
    </section>
</component>
1470 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.6' />
        <!-- Optional Functional Status Assessment Section content -->
    </section>
</component>
1475 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.7' />
        <!-- Optional Braden Score Assessment Section content -->
    </section>
</component>
1480 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.8' />
        <!-- Optional Geriatric Depression Scale Section content -->
    </section>
</component>
1485 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.9' />
        <!-- Optional Minimum Data Set Section content -->
    </section>
</component>

```

Figure 6.3.3.2.22.2-1: Specification for Coded Functional Status Assessment Section*Add Section 6.3.3.2.23***6.3.3.2.23 Pain Scale Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2	
General Description	The Pain Scale Assessment contains a coded observation reflecting the patient's reported intensity of pain on a scale from 0 to 10.	
LOINC Code	Opt	Description
38208-5	R	Pain severity
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.12.3.1	R	Pain Score Observation

```

1490 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.2' />
<id root=' ' extension=' ' />
<code code='38208-5' displayName='Pain severity'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
  <!-- Required Pain Score Observation element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.1' />
  :
</entry>
</section>
</component>

```

Figure 6.3.3.2.23-1: Specification for Pain Scale Assessment Section

1510 *Add Section 6.3.3.2.24*

6.3.3.2.24 Braden Score Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3	
General Description	This section reports the Braden score and its related assessments in machine and human readable form.	
LOINC Code	Opt	Description
38228-3	R	BRADEN SCALE SKIN ASSESSMENT PANEL
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.12.3.2	R	Braden Score Observation

```

1515 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.3' />
<id root=' ' extension=' ' />
<code code='38228-3' displayName='BRADEN SCALE SKIN ASSESSMENT PANEL'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
  <!-- Required Braden Score Observation element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.2' />
  :
</entry>
</section>
</component>

```

Figure 6.3.3.2.24-1: Specification for Braden Score Section

Add Section 6.3.3.2.25

1535 **6.3.3.2.25 Geriatric Depression Scale Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4	
General Description	This section reports the Geriatric Depression Scale score and its related assessments in machine and human readable form.	
LOINC Code	Opt	Description
48542-5	R	Geriatric Depression Scale (GDS) Panel
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.12.3.4	R	Geriatric Depression Score Observation

1540 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.4' />
 <id root=' ' extension=' ' />
 <code code='48542-5' displayName='Geriatric Depression Scale (GDS) Panel'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 <entry>
 :
 <!-- Required Geriatric Depression Score Observation element -->
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.4' />
 :
 </entry>
 </section>
 </component>

1545 **Figure 6.3.3.2.25-1: Specification for Geriatric Depression Scale Section**

Add Section 6.3.3.2.26

6.3.3.2.26 Physical Function Section 1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5	
General Description	This section reports scores from section G of the Minimum Data Set.	
LOINC Code	Opt	Description
46006-3	R	Physical functioning and structural problems
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7	O	Survey Panel At least one Survey Panel or Survey Observation shall be present.

1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6	O	<u>Survey Observation</u> At least one Survey Panel or Survey Observation shall be present.
------------------------------------	---	--

1560

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.2.5' />
    <id root=' ' extension=' ' />
    <code code='46006-3' displayName='Physical functioning and structural problems'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Optional Survey Panel element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Survey Observation element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6' />
      :
    </entry>
  </section>
</component>

```

1565

1570

1575

1580

1585

Figure 6.3.3.2.26-1: Specification for Physical Function Section

6.3.3.2.26.1 Constraints

[Survey Panel](#) found in this section SHOULD be identified using the panel codes found in the table below, and SHOULD contain one or more survey observations from that panel.

1590 [Survey Observation](#) found in this section SHOULD use the LOINC codes from Table 6.3.3.2.26.1 to express the answer to one or more questions from the Minimum Data Set Section G. The Survey Observations shall not contain a <methodCode> or <targetSiteCode> element, as these are not appropriate to the MDS Survey instrument.

Table 6.3.3.2.26.1-1: Panel Codes

Panel Code	Observation Code	Description	Data Type	Value Set
46007-1	Panel	ADL self-performance or support		
	45588-1	Bed mobility - self-performance	CO	2.16.840.1.113883.6.257.755
	45589-9	Bed mobility - support provided	CO	2.16.840.1.113883.6.257.768
	45590-7	Transfer - self-performance	CO	2.16.840.1.113883.6.257.755
	45591-5	Transfer - support provided	CO	2.16.840.1.113883.6.257.768
	45592-3	Walk in room - self-performance	CO	2.16.840.1.113883.6.257.755
	45593-1	Walk in room - support provided	CO	2.16.840.1.113883.6.257.768
	45594-9	Walk in corridor - self-performance	CO	2.16.840.1.113883.6.257.755
	45595-6	Walk in corridor - support provided	CO	2.16.840.1.113883.6.257.768

Panel Code	Observation Code	Description	Data Type	Value Set
	45596-4	Locomotion on unit - self-performance	CO	2.16.840.1.113883.6.257.755
	45597-2	Locomotion on unit - support provided	CO	2.16.840.1.113883.6.257.768
	45598-0	Locomotion off unit - self-performance	CO	2.16.840.1.113883.6.257.755
	45599-8	Locomotion off unit - support provided	CO	2.16.840.1.113883.6.257.768
	45600-4	Dressing - self-performance	CO	2.16.840.1.113883.6.257.755
	45601-2	Dressing - support provided	CO	2.16.840.1.113883.6.257.768
	45602-0	Eating - self-performance	CO	2.16.840.1.113883.6.257.755
	45603-8	Eating - support provided	CO	2.16.840.1.113883.6.257.768
	45604-6	Toilet use - self-performance	CO	2.16.840.1.113883.6.257.755
	45605-3	Toilet use - support provided	CO	2.16.840.1.113883.6.257.768
	45606-1	Personal hygiene - self-performance	CO	2.16.840.1.113883.6.257.755
	45607-9	Personal hygiene - support provided	CO	2.16.840.1.113883.6.257.768
46008-9	Panel	Bathing		
	45608-7	Bathing - self-performance	CO	2.16.840.1.113883.6.257.860
	45609-5	Bathing - support provided	CO	2.16.840.1.113883.6.257.768
46009-7	Panel	Test for balance		
	45610-3	Balance while standing	CO	2.16.840.1.113883.6.257.876
	45523-8	Balance while sitting	CO	2.16.840.1.113883.6.257.876
46010-5	Panel	Functional limitation in range of motion		
	45524-6	Range of motion^Neck	CO	2.16.840.1.113883.6.257.889
	45525-3	Voluntary movement^Neck	CO	2.16.840.1.113883.6.257.898
	45526-1	Range of motion^Upper Extremity	CO	2.16.840.1.113883.6.257.889
	45527-9	Voluntary movement^Upper Extremity	CO	2.16.840.1.113883.6.257.898
	45528-7	Range of motion^Hand	CO	2.16.840.1.113883.6.257.889
	45529-5	Voluntary movement^Hand	CO	2.16.840.1.113883.6.257.898
	45530-3	Range of motion^Lower Extremity	CO	2.16.840.1.113883.6.257.889
	45531-1	Voluntary movement^Lower Extremity	CO	2.16.840.1.113883.6.257.898
	45532-9	Range of motion^Foot	CO	2.16.840.1.113883.6.257.889
	45533-7	Voluntary movement^Foot	CO	2.16.840.1.113883.6.257.898
	45534-5	Other - range of motion	CO	2.16.840.1.113883.6.257.889
	45535-2	Other - voluntary movement	CO	2.16.840.1.113883.6.257.898
46011-3	Panel	Modes of locomotion		
	45536-0	Uses cane, walker or crutch	CO	2.16.840.1.113883.6.257.117

Panel Code	Observation Code	Description	Data Type	Value Set
	45537-8	Wheeled self	CO	2.16.840.1.113883.6.257.117
	45538-6	Other person wheeled	CO	2.16.840.1.113883.6.257.117
	45539-4	Uses wheelchair for primary locomotion	CO	2.16.840.1.113883.6.257.117
	45540-2	No modes of locomotion	CO	2.16.840.1.113883.6.257.117
46012-1	Panel	Modes of transfer		
	45541-0	Bedfast all or most of the time	CO	2.16.840.1.113883.6.257.117
	45542-8	Bed rails for bed mobility or transfer	CO	2.16.840.1.113883.6.257.117
	45543-6	Lifted manually	CO	2.16.840.1.113883.6.257.117
	45544-4	Lifted mechanically	CO	2.16.840.1.113883.6.257.117
	45545-1	Transfer aid	CO	2.16.840.1.113883.6.257.117
	45546-9	No mode of transfer	CO	2.16.840.1.113883.6.257.117
No Panel	45611-1	Task segmentation	CO	2.16.840.1.113883.6.257.117
46013-9	Panel	ADL functional rehabilitation potential		
	45612-9	Resident sees increased independence capability	CO	2.16.840.1.113883.6.257.117
	45613-7	Staff sees increased independence capability	CO	2.16.840.1.113883.6.257.117
	45614-5	Resident slow performing tasks or activity	CO	2.16.840.1.113883.6.257.117
	45615-2	Difference in morning to evening activities of daily living	CO	2.16.840.1.113883.6.257.117
	45616-0	Activities of daily living rehabilitation potential - none of above	CO	2.16.840.1.113883.6.257.117
	45617-8	Change in activities of daily living function	CO	2.16.840.1.113883.6.257.464

- 1595 The coded original values used in the observations above are described in more detail in the table below.

Explanation	Coded Value
2.16.840.1.113883.6.257.755	
INDEPENDENT-No help or oversight -OR- Help/oversight provided only 1 or 2 times during last 7 days	0
SUPERVISION-Oversight, encouragement or cueing provided 3 or more times during last7 days -OR- Supervision (3 or more times) plus physical assistance provided only 1 or 2 times during last 7 days	1

Explanation	Coded Value
LIMITED ASSISTANCE-Resident highly involved in activity; received physical help in guided maneuvering of limbs or other nonweight-bearing assistance 3 or more times - OR-More help provided only 1 or 2 times during last 7 days	2
EXTENSIVE ASSISTANCE-While resident performed part of activity, over last 7-day period, help of following type(s) provided 3 or more times: - Weight-bearing support - Full staff performance during part (but not all) of last 7 days	3
TOTAL DEPENDENCE-Full staff performance of activity during entire 7 days	4
ACTIVITY DID NOT OCCUR during entire 7 days	8
2.16.840.1.113883.6.257.768	
No setup or physical help from staff	0
Setup help only	1
One person physical assist	2
ADL activity itself did not occur during entire 7 days	8
2.16.840.1.113883.6.257.860	
Independent-No help provided	0
Supervision-Oversight help only	1
Physical help limited to transfer only	2
Physical help in part of bathing activity	3
Total dependence	4
Activity itself did not occur during entire 7 days	8
2.16.840.1.113883.6.257.876	
Maintained position as required in test	0
Unsteady, but able to rebalance self without physical support	1
Partial physical support during test; or stands (sits) but does not follow directions for test	2
Not able to attempt test without physical help	3
2.16.840.1.113883.6.257.889	
No limitation	0
Limitation on one side	1
Limitation on both sides	2
2.16.840.1.113883.6.257.898	
No loss	0
Partial loss	1
Full loss	2
2.16.840.1.113883.6.257.117	
No	0
Yes	1
UTD	-
2.16.840.1.113883.6.257.464	
No change	0
Improved	1

Explanation	Coded Value
Deteriorated	2

*Add Section 6.3.3.2.27*1600 **6.3.3.2.27 Preprocedure Review of Systems Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.13**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.13	
Parent Template	Review of Systems (1.3.6.1.4.1.19376.1.5.3.1.3.18)	
General Description	The pre-procedure review of systems section shall contain only required and optional subsections dealing with the responses the patient gave to a set of routine questions on body systems in general and specific risks of anesthesia not covered in general review of systems.	
LOINC Code	Opt	Description
10187-3	R	REVIEW OF SYSTEMS
Subsections	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.9.46	R	Implanted Medical Device Review
1.3.6.1.4.1.19376.1.5.3.1.1.9.47	R2	Pregnancy Status Review
1.3.6.1.4.1.19376.1.5.3.1.1.9.14	R	Anesthesia Risk Review of Systems

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18' />
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.13' />
    <id root=' ' extension=' ' />
    <code code='10187-3' displayName='REVIEW OF SYSTEMS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.46' />
        <!-- Required Implanted Medical Device Review Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.47' />
        <!-- Required if known Pregnancy Status Review Section content -->
      </section>
    </component>
    <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.14' />
        <!-- Required Anesthesia Risk Review of Systems Section content -->
      </section>
    </component>
  </section>
</component>

```

Figure 6.3.3.2.27-1: Specification for Preprocedure Review of Systems Section

1635

*Add Section 6.3.3.2.28***6.3.3.2.28 Estimated Delivery Date Section 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.1**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.1	
General Description	This section contains the physician's best estimate of the patients due date. This is generally done both on an initial evaluation, and later confirmed at 18-20 weeks. The date is supported by evidence such as the patient's history of last menstrual period, a physical examination, or ultrasound measurements.	
LOINC Code	Opt	Description
57060-6	R	Estimated date of delivery
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1	R	<p>Estimated Delivery Date Observation</p> <p>This is a simple observation to represent the estimated due date with a supporting observation or observations that state the method used and date implied by that method. If one observation is present, then it is to be interpreted as the initial EDD. If the initial observation dates indicate the EDD is within the 18 to 20 weeks completed gestation, that observation will also populate the 18-20 week update. If the initial observation indicates an EDD of more than 20 weeks EGA, then no value will be placed in the 18-20 week update field.</p>

```

1640 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.1' />
        <id root=' ' extension=' ' />
        <code code='57060-6' displayName='Estimated date of delivery'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
        <entry>
            :
            <!-- Required Estimated Due Date Observation element -->
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1' />
            :
        </entry>
    </section>
</component>

```

Figure 6.3.3.2.28-1: Specification for Estimated Delivery Dates Section*Add Section 6.3.3.2.29***6.3.3.2.29 History of Tobacco Use Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.8**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.8	
General Description	The history of tobacco use section shall contain a description of the responses the patient gave to a set of routine questions on the history of tobacco use.	
LOINC Code	Opt	Description
11366-2	R	HISTORY OF TOBACCO USE

```

1660 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.8' />
        <id root=' ' extension=' ' />
        <code code='11366-2' displayName='HISTORY OF TOBACCO USE'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
    </section>
</component>

```

Figure 6.3.3.2.29-1: Specification for History of Tobacco Use Section*Add Section 6.3.3.2.30*

6.3.3.2.30 Current Alcohol/Substance Abuse Section**1.3.6.1.4.1.19376.1.5.3.1.1.9.10**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.10	
General Description	The history of alcohol/substance abuse section shall contain a description of the responses the patient gave to a set of routine questions on the current abuse of alcohol or other substances.	
LOINC Code	Opt	Description
18663-5	R	HISTORY OF PRESENT ALCOHOL AND/OR SUBSTANCE ABUSE

1680

1685

1690

1695

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.10' />
    <id root=' ' extension=' ' />
    <code code='18663-5' displayName='HISTORY OF PRESENT ALCOHOL AND/OR SUBSTANCE ABUSE'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

Figure 6.3.3.2.30-1: Specification for Current Alcohol/Substance Abuse Section*Add Section 6.3.3.2.31***6.3.3.2.31 History of Blood Transfusion Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.12**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.12	
General Description	The History of Blood Transfusion section shall contain a narrative description of the blood products the patient has received in the past, including any reactions to blood products.	
LOINC Code	Opt	Description
56836-0	R	History of blood transfusion

1700

```

1705 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.12'>
<id root=' ' extension=' ' />
<code code='56836-0' displayName='History of blood transfusion'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
</section>
1710 </component>
1715

```

Figure 6.3.3.2.31-1: Specification for History of Blood Transfusion Section*Add Section 6.3.3.2.32***6.3.3.2.32 Anesthesia Risk Review of Systems Section****1.3.6.1.4.1.19376.1.5.3.1.1.9.14**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.14	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.3.18	
General Description	The anesthesia review of systems section shall contain a description of the responses the patient gave to a set of routine questions on specific risks of anesthesia not covered in general review of systems such as broken teeth, airway limitations, positioning limitations, recent infections, and history of personal anesthesia problems..	
LOINC Code	Opt	Description
57081-2	R	Anesthesia Risk Review of Systems

```

1725 <component>
<section>   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.18'>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.14'>
<id root=' ' extension=' ' />
<code code='57081-2' displayName='Anesthesia Risk Review of Systems'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
</section>
1730 </component>
1735

```

Figure 6.3.3.2.32-1: Specification for Anesthesia Risk Review of Systems Section*Add Section 6.3.3.2.33*

1740

6.3.3.2.33 Implanted Medical Device Review Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.46

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.46	
General Description	The implanted medical device review section shall contain a description of the medical devices that are inserted into the patient, whether internal or partially external.	
LOINC Code	Opt	Description
57080-4	R	Implanted medical device

1745

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.46' />
    <id root=' ' extension=' '/>
    <code code='57080-4' displayName='Implanted medical device'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

1750

1755

Figure 6.3.3.2.33-1: Specification for Implanted Medical Device Review Section

1760

*Add Section 6.3.3.2.34***6.3.3.2.34 Pregnancy Status Review Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.47**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.47	
General Description	The pregnancy status review section shall contain a description of the responses the patient gave to a set of routine questions regarding potential pregnancy in females of child-bearing-age. It shall include a Pregnancy Status Organizer.	
LOINC Code	Opt	Description
11449-6	R	Pregnancy Status-Reported
Entries	Opt	Description
TBD	O	Pregnancy Status Review Organizer

1765 <component>

1770 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.47' />
 <id root=' ' extension=' ' />
 <code code='11449-6' displayName='Pregnancy Status-Reported'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>

1775 </section>
 </component>

Figure 6.3.3.2.34-1: Specification for Pregnancy Status Review Section*Add Section 6.3.3.2.35*

1780

6.3.3.2.35 History of Infection Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1	
General Description	The History of Infection section shall contain a narrative description of any infections the patient may have contracted prior to the patient's current visit or admission.	
LOINC Code	Opt	Description
56838-6	R	History of infectious disease

1785 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1' />
 <id root=' ' extension=' ' />
 <code code='56838-6' displayName='History of infectious disease'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>

1790 </section>
 </component>

Figure 6.3.3.2.35-1: Specification for History of Infection Section*Add Section 6.3.3.2.36***6.3.3.2.36 Coded Social History Section 1.3.6.1.4.1.19376.1.5.3.1.3.16.1**

1800

Table 6.3.3.2.36-1: Coded Social History Section

Template Name		Coded Social History Section			
Template ID		1.3.6.1.4.1.19376.1.5.3.1.3.16.1			
Parent Template		IHE Social History Section 1.3.6.1.4.1.19376.1.5.3.1.3.16			
General Description		The social history section shall contain a narrative description of the person's beliefs, home life, community life, work life, hobbies, and risky habits. It shall include Social History Observations.			
Section Code		29762-2, LOINC, "Social History"			
Author		If not the author from the encompassing context, include author. Role and entity must be specified if not inherited.			
Informant		If not the informant from the encompassing context, include informant. Role and entity must be specified if not inherited.			
Subject		If not the subject from the encompassing context, include subject. Role and entity must be specified if not inherited.			
Opt and Card	Condition	Data Element or Section Name	Template ID	Specification Document	Vocabulary Constraint
Subsections					
O [0..1]		Occupational Data for Health Section	1.3.6.1.4.1.19376.1.7.3.1.3.24 .3		
Entries					
R [1..*]	PCC TF-2 6.3.4.24	Social History Observation	1.3.6.1.4.1.19376.1.5.3.1.4.13 .4		

1805

```

<component>
  <section>
    <templateId root='11.3.6.1.4.1.19376.1.5.3.1.3.16' />
    <templateId root='11.3.6.1.4.1.19376.1.5.3.1.3.16.1' />
    <id root=' ' extension=' ' />
    <code code='29762-2' displayName='SOCIAL HISTORY'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

1810

Figure 6.3.3.2.36-1: Specification for Coded Social History Section

1815

Add Section 6.3.3.2.37

6.3.3.2.37 Coded History of Infection Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1.1	
Parent Template	History of Infection (1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1)	
General Description	The History of Infection section shall contain a narrative description of any infections the patient may have contracted prior to the patient's current condition. It shall include entries for problems as described in the Entry Content Modules.	
LOINC Code	Opt	Description
56838-6	R	History of infectious disease
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.5.2	R	Problem Concern Entry

1820

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1.1.1'>
    <id root=' ' extension=' '/>
    <code code='56838-6' displayName='History of infectious disease'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'>
    <text>
      Text as described above
    </text>
  </section>
</component>
```

1825

1830

Figure 6.3.3.2.37-1: Specification for Coded History of Infection Section*Add Section 6.3.3.2.38***6.3.3.2.38 Prenatal Events Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.2**

Template ID	1.3.6.1.4.1.19376.1.5.3..1.1.21.2.2	
General Description	The Prenatal Events Section shall include narrative text describing pertinent prenatal information that has a direct impact on the process of labor and delivery. It shall also include subsections if known.	
LOINC Code	Opt	Description
57073-9	R	Prenatal events
Subsections	Opt	Description
Coded Results This section SHOULD contain laboratory results and procedures as pertaining to the pregnancy , e.g., amniocentesis, cordocentesis, chorionic villus sampling.	R2	1.3.6.1.4.1.19376.1.5.3.1.3.28
Procedures and Interventions This section SHOULD contain procedures that took place during the prenatal period (i.e., prenatal care, prenatal complications, prenatal surgeries)	R2	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11

Event Outcomes This section contains event outcomes related to prenatal events e.g., miscarriage, infection.	R2	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9
--	----	------------------------------------

1835

```
<component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.2' />
```

1840

```
<id root=' ' extension=' '/>
<code code='57073-9' displayName='Prenatal events'
codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
```

1845

```
<text>
Text as described above
</text>
```

```
<component>
```

```
<section>
```

```
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.28' />
```

```
<!-- Required if known Coded Results Section -->
```

1850

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

```
<component>
```

```
<section>
```

```
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11' />
```

```
<!-- Required if known Procedures and Interventions Section -->
```

1855

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

```
<component>
```

```
<section>
```

```
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9' />
```

```
<!-- Required if known Event Outcomes Section -->
```

1860

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

1865

Figure 6.3.3.2.38-1: Specification for Prenatal Events Section*Add Section 6.3.3.2.39***6.3.3.2.39 Labor and Delivery Events Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.3**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.3	
Parent Template		
General Description	The Labor and Delivery Events Section SHALL include a narrative text containing relevant information collected during the labor and delivery process.	
LOINC Code	Opt	Description
57074-7	R	Labor and delivery process

Subsections	Opt	Description
Procedures and Interventions The subsection SHALL contain procedures and interventions specific to labor and delivery events. These may include induction, the delivery type (e.g., vaginal, vaginal birth after cesarean section or cesarean section along with incision type), electronic fetal monitoring, etc.	R2	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11
Coded Event Outcomes This section SHOULD contain outcomes related to the labor and delivery process such as live birth or stillborn. The subsection shall include coded event outcomes such as live birth or stillborn and also including maternal death with date/time. Furthermore, Coded Event Outcomes section shall contain a simple Observation using LOINC Code 11636-8 that reports the number of births live or dead that occurred during the delivery event.	R2	1.3.6.1.4.1.19376.1.7.3.1.1.13.7

1870 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.3'/>
 <id root=' ' extension=' '/>
 <code code='57074-7' displayName='Labor and delivery process'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11' />
 <!-- Required if known Procedures and Interventions Section -->
 </section>
 </component>
 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.7' />
 <!-- Required if known Coded Event Outcomes Section -->
 </section>
 </component>
 </section>
 </component>

Figure 6.3.3.2.39-1: Specification for Labor and Delivery Process Section

1895 *Add Section 6.3.3.2.40*

6.3.3.2.40 Newborn Delivery Information Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.4

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.4	
General Description	The Newborn Delivery Information Section SHALL include a narrative text containing information collected at the birth and up to the transfer of the infant from the birthing room to a post-natal unit.	
LOINC Code	Opt	Description
57075-4	R	Newborn delivery information from newborn
Subsections	Opt	Description
Coded Detailed Physical Examination Section This section SHALL include information about the newborn genitalia; weight; length; head circumference, size (AGA, SGA or LGA); Apgar score assessment ; vital signs, physical exam findings	R	1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1
Active Problems This section SHALL describe problems that the newborn might have had during or immediately prior to delivery.	R2	1.3.6.1.4.1.19376.1.5.3.1.3.6
Procedures and Interventions This section SHALL include the procedures and interventions received by the newborn such as suction or resuscitation.	R2	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11
Medications Administered This section SHALL include the medication that was administered to the newborn while in the birthing suite such as: Vitamin K (Aquamephyton) injection; erythromycin eye ointment; and resuscitation medications (if any) including date, time, and route of administration.	R2	1.3.6.1.4.1.19376.1.5.3.1.3.21
Event Outcomes This section SHALL include the outcomes of the procedures and interventions such as a resuscitation event.	R2	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9
Coded Event Outcomes	C	1.3.6.1.4.1.19376.1.7.3.1.1.13.7
Coded Results	C	1.3.6.1.4.1.19376.1.5.3.1.3.28
Intake and Output This section SHALL include any intake and output while the newborn is in the delivery suite (excluding estimated blood loss) such as: first urine/void; stool; gastric output	C	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.3

1900	<component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.4' /> <id root=' ' extension=' ' /> <code code='57075-4' displayName='Newborn delivery information from newborn' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' /> <text> Text as described above </text>
1905	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1' /> <!-- Required Coded Detailed Physical Examination Section --> </section>
1910	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.6' /> <!-- Required if known Active Problems Section --> </section>
1915	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.13.2.11' /> <!-- Required if known Procedures and Interventions Section --> </section>
1920	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.21' /> <!-- Required if known Medications Administered Section --> </section>
1925	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9' /> <!-- Required if known Event Outcomes Section --> </section>
1930	</component> <component>
1935	</component> </section> </component>

Figure 6.3.3.2.40-1: Specification for Newborn Delivery Information Section

1940

*Add Section 6.3.3.2.41***6.3.3.2.41 Postpartum Hospitalization Treatment Section****1.3.6.1.4.1.19376.1.5.3.1.1.21.2.7**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.7	
Parent Template		
General Description	The Postpartum Treatment Section shall include a narrative description of the treatment delivered to the mother subsequent to the delivery.	
LOINC Code	Opt	Description
57076-2	R	Postpartum hospitalization TREATMENT
Subsections	Opt	Description
Immunizations This section SHOULD contain the immunization given to the mother prior to the discharge from the birthing facility.	O	1.3.6.1.4.1.19376.1.5.3.1.4.12
Medications Administered This SHOULD include commonly prescribed maternal medications including contraceptive medication.	R2	1.3.6.1.4.1.19376.1.5.3.1.3.21
Procedures and Interventions This section SHALL include the procedures and interventions received by the mother during the immediate post-partum period e.g., transfusion or curettage.	R	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11
Coded Results This section SHOULD contain laboratory results and procedures as pertaining to the mother while discharged such as the hemoglobin or the hematocrit level.	R2	1.3.6.1.4.1.19376.1.5.3.1.3.28
Care plan This section SHOULD include the plan of care for the mother upon her discharge such as the feeding method or the contraceptive plan	O	1.3.6.1.4.1.19376.1.5.3.1.3.31
Discharge Diet This section SHALL include the diet that the mother was recommended upon her discharge.	R	1.3.6.1.4.1.19376.1.5.3.1.3.33

1945	<component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.7' /> <id root=' ' extension=' ' /> <code code='57076-2' displayName='POST PARTUM HOSPITALIZATION TREATMENT' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' /> <text> Text as described above </text>
1950	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.6' /> <!-- Required Active Problems Section --> </section> </component>
1955	<component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.12' /> <!-- Optional Immunizations Section --> </section>
1960	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.13.2.11' /> <!-- Required if known Hospital Discharge Medication Section --> </section>
1965	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.22' /> <!-- Required Procedures and Interventions Section --> </section>
1970	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.28' /> <!-- Required if known Coded Results Section --> </section>
1975	</component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31' /> <!-- Optional Care Plan Section -->
1980	</section> </component> <component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.33' /> <!-- Required Discharge Diet Section -->
1985	</section> </component>
1990	</component> <component> <section>
1995	</section> </component>

Figure 6.3.3.2.41-1: Specification for Postpartum Treatment Section

2000

*Add Section 6.3.3.2.42***6.3.3.2.42 Event Outcomes Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9	
Parent Template		
General Description	The Event Outcome Section shall include a narrative description of the outcomes following a procedure, an intervention or a problem.	
LOINC Code	Opt	Description
42545-4	R	EVENT OUTCOME

2005

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9' />
    <id root=' ' extension=' ' />
    <code code='42545-4' displayName='EVENT OUTCOME'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </component>

```

2010

Figure 6.3.3.2.42-1: Specification for Event Outcomes Section

2015

*Add Section 6.3.3.2.43***6.3.3.2.43 Newborn Status at Maternal Discharge 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.8**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.8	
Parent Template		
General Description	The Newborn Status and Maternal Discharge section shall contain a narrative description of the status and disposition of the newborn at the time of maternal discharge.	
LOINC Code	Opt	Description
57077-0	R	Newborn status at maternal discharge from newborn

2020

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.8' />
    <id root=' ' extension=' ' />
    <code code='57077-0' displayName='Newborn status at maternal discharge from newborn'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </component>

```

2025

Figure 6.3.3.2.43-1: Specification for Newborn Status at Maternal Discharge Section

2030

*Add Section 6.3.3.2.44***6.3.3.2.44 History of Surgical Procedures Section 1.3.6.1.4.1.19376.1.5.3.1.1.16.2.2**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.16.2.2	
Parent Template		
General Description	The History of Surgical Procedures Section shall contain a narrative description of the surgical procedures performed on the patient.	
LOINC Code	Opt	Description
10167-5	R	History of surgical procedures

2035

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.16.2.2' />
    <id root=' ' extension=' ' />
    <code code='10167-5' displayName='History of surgical procedures'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </component>
```

2040

Figure 6.3.3.2.44-1: Specification for History of Surgical Procedures Section

2045

*Add Section 6.3.3.2.45***6.3.3.2.45 Operative Note Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.6**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.6	
Parent Template		
General Description	The Operative Note Section shall contain a narrative description of the current operation or surgical procedure in detail.	
LOINC Code	Opt	Description
10223-6	R	Surgical operation note surgical procedure

2050

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.6' />
    <id root=' ' extension=' ' />
    <code code='10223-6' displayName='Surgical operation note surgical procedure'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </component>
```

2055

Figure 6.3.3.2.45-1: Specification for Operative Note Section

2060

Add Section 6.3.3.2.46

6.3.3.2.46 Child Functional Status Assessment 1.3.6.1.4.1.19376.1.7.3.1.1.13.3

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.13.3	
General Description	This section provides a description of the child's status of normal functioning at the time the document was created. This section includes the psychomotor and the eating and sleeping assessments. This section shall include the Psychomotor Test Observation entry.	
LOINC Code	Opt	Description
47420-5	R	Functional Status Assessment
Subsections	Opt	Description
1.3.6.1.4.1.19376.1.7.3.1.1.13.4	O	Psychomotor Development
1.3.6.1.4.1.19376.1.7.3.1.1.13.5	O	Eating and sleeping assessment

2065

Example

2070

```
<component>
  <section>
    <templateId root="1.3.6.1.4.1.19376.1.7.3.1.1.13.3"/>
    <id root="16696797-f854-443d-8819-231ee09cad71"/>
    <code code="47420-5" displayName="Functional Status Assessment"
      codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
    <title/>
    <text/>
    <component>
      <section>
        <!-- Optional Psychomotor Development section -->
        <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.4'>
        :
      </section>
    </component>

    <component>
      <section>
        <!-- Eating and sleeping assessment section -->
        <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.5'>
        :
      </section>
    </component>
  </section>
</component>
```

2075

2080

2085

2090

Add Section 6.3.3.2.47

6.3.3.2.47 Psychomotor Development Section 1.3.6.1.4.1.19376.1.7.3.1.1.13.4

2095

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.13.4	
General Description	This section describes a test battery in order to evaluate the psychomotricity of the newborn.	
LOINC Code	Opt	Description
xx-MCH-PsychoMDev	R	Psychomotor development
Entries	Opt	Description
Simple Observation	R	1.3.6.1.4.1.19376.1.5.3.1.4.13

2100	<component> <section> <!-- Psychomotor Development section templateId --> <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.4' /> <id root=' ' extension=' '/> <code code='47420-5' displayName=' Functional Status Assessment ' codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" /> <text> : </text> <entry> : <!--Required simple Observation element --> <templateId root=1.3.6.1.4.1.19376.1.5.3.1.4.13> : </entry> </section> </component>
2105	
2110	

2115

Add Section 6.3.3.2.48

6.3.3.2.48 Eating and Sleeping Assessment Section 1.3.6.1.4.1.19376.1.7.3.1.1.13.5

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.13.5	
General Description	This section describes a test battery in order to evaluate the psychomotricity of the newborn.	
LOINC Code	Opt	Description
47420-5	R	Functional Status Assessment
Entries	Opt	Description
Simple Observation	R	1.3.6.1.4.1.19376.1.5.3.1.4.13

```

2120 <component>
    <section>
        <!--Eating and Sleeping assessment section templateId -->
        <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.5' />
        <id root=' ' extension=' '/>
        <code code='47420-5' displayName=' Functional Status Assessment '
              codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" />
        <text>
            :
        </text>
        <entry>
            :
            <!-- Required Simple Observation element -->
            <templateId root=1.3.6.1.4.1.19376.1.5.3.1.4.13/>
            :
        </entry>
    </section>
</component>

```

Add Section 6.3.3.2.49

2140 **6.3.3.2.49 Coded Event Outcomes 1.3.6.1.4.1.19376.1.7.3.1.1.13.7**

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.13.7	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.9	
General Description	The Coded Event Outcome Section shall include a narrative description of the outcomes following a procedure, an intervention or a problem, and outcomes related to the labor and delivery process such as live birth or stillborn. It shall include entries for observation as described in the Simple Observation entry, or optionally as Problem Entry observations.	
LOINC Code	Opt	Description
42545-4	R	EVENT OUTCOME
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.13	R	Simple Observation
1.3.6.1.4.1.19376.1.5.3.1.1.25.1.4.1	R2	Patient Transfer
1.3.6.1.4.1.19376.1.5.3.1.4.5	O	Problem Entry

```

2145 <component>
    <section>
        <!--Coded Event Outcomes assessment section templateId -->
        <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.13.7' />
        <id root=' ' extension=' '/>
        <code code='42545-4' displayName='Event Outcome'
              codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" />
        <text>
            :
        </text>
        <entry>
            :
        <!-- Required Simple Observation element -->
        <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.13"/>
            :
        </entry>
        <entry>
            :
        <!-- Required if known Patient Transfer element -->
        <templateId root="1.3.6.1.4.1.19376.1.5.3.1.1.25.1.4.1"/>
            :
        </entry>
        <entry>
            :
        <!-- Optional Problem Entry element -->
        <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.5"/>
            :
        </entry>
    </section>
</component>

```

2175

Add Section 6.3.3.2.50 (Occupational History - removed 2011-09 at the request of QRPH)

6.3.3.2.50 Intentionally blank

Add Section 6.3.3.2.51 (Patient Status - removed 2011-09 at the request of QRPH)

6.3.3.2.51 Intentionally blank

2180

Add Section 6.3.3.2.52 Cancer Control - removed 2011-09 at the request of QRPH)

6.3.3.2.52 Intentionally blank

Add Section 6.3.3.2.53

6.3.3.2.53 Notifications, Alerts, and Reminders Section

1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1.x

2185

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1.x	
General Description	The Notifications, Reminders and Alerts section highlights areas of care plan non-conformance and directs the need for follow-up communications.	
LOINC Code	Opt	Description
XXX	R	Notifications, Alerts, and Reminders
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.7	C	<u>Medications</u> Medications entries shall appear for all pending medications when present. These entries shall be in intent mood.
1.3.6.1.4.1.19376.1.5.3.1.4.19	C	<u>Procedure</u> Procedure entries shall appear for all pending procedures when present. These entries shall be in intent mood.
1.3.6.1.4.1.19376.1.5.3.1.4.14	C	<u>Encounter</u> Encounter entries should appear for all pending follow-up encounters. These entries shall be in promise or appointment request mood.
1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1	C	<u>Observation Request</u> Observation request entries should appear for all pending follow-up observations. These entries shall appear in intent mood.

Add Section 6.3.3.2.54

6.3.3.2.54 Pain Assessment Panel Section 1.3.6.1.4.1.19376.1.5.3.1.1.20.2.4

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.4	
General Description	This contains a narrative description of the patient's pain, including such items as severity, quality, location, time of onset, radiation, etc.	
LOINC Code	Opt	Description
38212-7	R	Pain Assessment Panel

2190

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.2.4' />
    <id root=' ' extension=' '/>
    <code code='38212-7' displayName='Pain Assessment Panel'
      codeSystem='1.2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

2195

2200

Figure 6.3.2.54-1: Specification for Pain Assessment Panel Section

Add Section 6.3.3.2.55

2205 **6.3.3.2.55 History of Cognitive Function Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.11**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.11	
General Description	This contains a narrative description of a patient's mental status.	
LOINC Code	Opt	Description
11332-4	R	History of Cognitive Function

2210

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.11' />
    <id root=' ' extension=' ' />
    <code code='11332-4' displayName='History of Cognitive Function'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2215

Figure 6.3.2.55-1: Specification for History of Cognitive Function Section

2220

Add Section 6.3.3.2.56 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

6.3.3.2.56 Isolation Status Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.8

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.8	
General Description	The Isolation Status section describes a patient with an active infectious disease requiring additional personal protective equipment for healthcare providers.	
LOINC Code	Opt	Description
55017-8	R	ISOLATION OR QUARANTINE FOR ACTIVE INFECTIOUS DISEASE

2225

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.8' />
    <id root=' ' extension=' ' />
    <code code='55017-8' displayName=' ISOLATION OR QUARANTINE FOR ACTIVE INFECTIOUS DISEASE '
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2230

2235

Figure 6.3.3.2.56-1: Sample Isolation Status Section

Add Section 6.3.3.2.57 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

2240

6.3.3.2.57 Restraints Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.10

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.10	
General Description	The Restraints section describes the type of restraints currently in use on the patient to be transported.	
LOINC Code	Opt	Description
46067-5	R	DEVICES AND RESTRAINTS SET

2245

```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.10'>
      <id root=' ' extension=' ' />
      <code code='46067-5' displayName='DEVICES AND RESTRAINTS SET'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
      <text>
        Text as described above
      </text>
    </section>
  </component>
```

2250

Figure 6.3.3.2.57-1: Sample Restraints Section

2255

Add Section 6.3.3.2.58. Added 2011-09 from QRPH EHCP Profile

6.3.3.2.58 Risk Indicators for Hearing Loss

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.15.3.1	
General Description	This section SHALL include at least one entry describing hearing risk indicators for the subject	
LOINC® Code	Opt	Description
58232-0	R	HEARING LOSS RISK INDICATOR

Entries	Opt	Description
1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1	R	Risk Indicators for Hearing Loss Entry

```

2260 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.15.3.1' />
<id root=' ' extension=' ' />
<code code='58232-0' displayName='HEARING LOSS RISK INDICATOR '
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
<!-- Required Risk Indicators for Hearing Loss Entry element -->
<templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1' />
  :
</entry>
</section>
</component>

```

2275 **Figure 6.3.3.2.58-1: Sample Coded Risk Indicators for Hearing Loss Section**

Add Section 6.3.3.2.59. Added 2011-09 from QRPH PRPH-Ca Profile

6.3.3.2.59 Cancer Diagnosis Section 1.3.6.1.4.1.19376.1.7.3.1.3.14.1

Template ID	1.3.6.1.4.1.19376.1.7.3.1.3.14.1	
Parent ID	PCC Active Problem Section 1.3.6.1.4.1.19376.1.5.3.1.3.6 CCD 3.5 2.16.840.1.113883.10.20.1.11	
General Description	This section contains specific detailed information about cancer diagnosis(es) that are currently being monitored for the patient. A separate entry for each cancer diagnosis SHALL be provided.	
LOINC Code	Opt	Description
72135-7	R	Cancer Diagnosis
Entries	Opt	Description
1.3.6.1.4.1.19376.1.7.3.1.4.14.1	R	Cancer Diagnosis Entry

```

2280 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.7.3.1.3.14.1' />
<id root=' ' extension=' ' />
<code code='72135-7' displayName='Cancer Diagnosis'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
<!-- Required Cancer Diagnosis Entry element -->
<templateId root='1.3.6.1.4.1.19376.1.7.3.1.4.14.1' />
  :
</entry>
</section>
</component>

```

2295 **Figure 6.3.3.2.59-1: Specification for Cancer Diagnosis Section**

Add Section 6.3.3.3

6.3.3.3 Medications

2300

Add Section 6.3.3.3.1

6.3.3.3.1 Medications Section

Add Section 6.3.3.3.2

6.3.3.3.2 Admission Medication History Section

Add Section 6.3.3.3.3

2305

6.3.3.3.3 Medications Administered Section

Add Section 6.3.3.3.4

6.3.3.3.4 Hospital Discharge Medications Section

Add Section 6.3.3.3.5

6.3.3.3.5 Immunizations Section

2310

Add Section 6.3.3.4

6.3.3.4 Physical Exams

Note: Sections 6.3.3.4.1 through 6.3.3.4.29 reside in IHE PCC TF-2:6.3.3.4

Add Section 6.3.3.4.30

2315

6.3.3.4.30 Coded Detailed Physical Examination Section

1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1	
Parent Template	Detailed Physical Examination (1.3.6.1.4.1.19376.1.5.3.1.1.9.15)	
General Description	The Coded Detailed Physical Examination section shall contain a narrative description of the patient's physical findings. It shall include subsections, if known, for the exams that are performed.	
LOINC Code	Opt	Description
29545-1	R	PHYSICAL EXAMINATION
Subsections	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.5.3.2	R2	Coded Vital Signs

		Vital signs may be a subsection of the physical examination or they may stand alone
1.3.6.1.4.1.19376.1.5.3.1.1.9.16	R2	General Appearance
1.3.6.1.4.1.19376.1.5.3.1.1.9.48	R2	Visible Implanted Medical Devices
1.3.6.1.4.1.19376.1.5.3.1.1.9.17	R2	Integumentary System
1.3.6.1.4.1.19376.1.5.3.1.1.9.18	R2	Head
1.3.6.1.4.1.19376.1.5.3.1.1.9.19	R2	Eyes
1.3.6.1.4.1.19376.1.5.3.1.1.9.20	R2	Ears, Nose, Mouth and Throat
1.3.6.1.4.1.19376.1.5.3.1.1.9.21	R2	Ears
1.3.6.1.4.1.19376.1.5.3.1.1.9.22	R2	Nose
1.3.6.1.4.1.19376.1.5.3.1.1.9.23	R2	Mouth, Throat, and Teeth
1.3.6.1.4.1.19376.1.5.3.1.1.9.24	R2	Neck
1.3.6.1.4.1.19376.1.5.3.1.1.9.25	R2	Endocrine System
1.3.6.1.4.1.19376.1.5.3.1.1.9.26	R2	Thorax and Lungs
1.3.6.1.4.1.19376.1.5.3.1.1.9.27	R2	Chest Wall
1.3.6.1.4.1.19376.1.5.3.1.1.9.28	R2	Breasts
1.3.6.1.4.1.19376.1.5.3.1.1.9.29	R2	Heart
1.3.6.1.4.1.19376.1.5.3.1.1.9.30	R2	Respiratory System
1.3.6.1.4.1.19376.1.5.3.1.1.9.31	R2	Abdomen
1.3.6.1.4.1.19376.1.5.3.1.1.9.32	R2	Lymphatic System
1.3.6.1.4.1.19376.1.5.3.1.1.9.33	R2	Vessels
1.3.6.1.4.1.19376.1.5.3.1.1.9.34	R2	Musculoskeletal System
1.3.6.1.4.1.19376.1.5.3.1.1.9.35	R2	Neurologic System
1.3.6.1.4.1.19376.1.5.3.1.1.9.36	R2	Genitalia
1.3.6.1.4.1.19376.1.5.3.1.1.9.37	R2	Rectum
1.3.6.1.4.1.19376.1.5.3.1.1.16.2.1	R2	Extremities
1.3.6.1.4.1.19376.1.5.3.1.1.21.2.10	R2	Pelvis

```

2320 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.15' />
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.15.1' />
        <id root=' ' extension=' ' />
        <code code='29545-1' displayName='PHYSICAL EXAMINATION'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
        <component>
            <section>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.25' />
                <!-- Optional Vital Signs Section content -->
            </section>
        </component>
    </section>
</component>

```

Figure 6.3.3.4.30-1: Coded Detailed Physical Examination Section

Add Section 6.3.3.4.31

2340 **6.3.3.4.31 Pelvis Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.10**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.10	
General Description	The Pelvis section shall include a narrative description of any type of exam of the reproductive organs.	
LOINC Code	Opt	Description
10204-6	R	PELVIS
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.5	O	Problem Entry

```

2345 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.10' />
        <id root=' ' extension=' ' />
        <code code='10204-6' displayName='PELVIS'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
        <entry>
            :
            <!-- Optional Problem Entry element -->
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5' />
            :
        </entry>
    </section>
</component>

```

Figure 6.3.3.4.31-1: Pelvis Section

2360

Add Section 6.3.3.4.32

6.3.3.4.32 Admission Physical Exam Section 1.3.6.1.4.1.19376.1.5.3.1.1.22.1.1.2.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.22.1.1.2.1	
General Description	The Admission physical exam section shall include a narrative description of the physical exams given during the admission to a hospital or similar type of facility.	
LOINC Code	Opt	Description
XX-AdmissionPhysicalExam	R	Admission physical exam

2365

```
<component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.22.1.1.2.1' />
<id root=' ' extension=' '/>
<code code='XX-AdmissionPhysicalExam' displayName='Admission physical exam'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
</section>
</component>
```

2370

2375

Figure 6.3.3.4.32-1: Admission Physical Exam Section

Add Section 6.3.3.4.33

6.3.3.4.33 Discharge Status 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.12

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.12	
Parent Template		
General Description	Discharge status should contain a narrative description of the status/condition of the patient at the time of discharge, such as stable, critical, etc.	
LOINC Code	Opt	Description
52523-8	R2	Discharge Status

2380

```
<component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.12' />
<id root=' ' extension=' '/>
<code code='52523-8' displayName=Discharge status'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
</component>
```

2385

2390

Figure 6.3.3.4.33-1: Discharge Status Section

6.3.3.5 Relevant Studies*Add Section 6.3.3.5.1*

2395

6.3.3.5.1 Results*Add Section 6.3.3.5.2***6.3.3.5.2 Coded Results***Add Section 6.3.3.5.3***6.3.3.5.3 Hospital Studies Summary**

2400

*Add Section 6.3.3.5.4***6.3.3.5.4 Coded Hospital Studies Summary***Add Section 6.3.3.5.5***6.3.3.5.5 Consultations 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.8**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.8	
General Description	The ED Consultations section shall contain a narrative description of the consultations obtained during an encounter of care.	
LOINC Code	Opt	Description
18693-2	R	ED CONSULTANT PRACTITIONER

2405

```
<component>
<section>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.8' />
  <id root=' ' extension=' ' />
  <code code='18693-2' displayName='ED CONSULTANT PRACTITIONER'
    codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
  <text>
    Text as described above
  </text>
</section>
</component>
```

2410

2415

Figure 6.3.3.5.5-1: Specification for ED Consultations Section*Add Section 6.3.3.5.6*

2420 6.3.3.5.6 Antenatal Testing and Surveillance Section

1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5	
Parent Template		
General Description	The Antenatal Testing and Surveillance section shall contain a narrative description of reports and data from tests and surveillance performed during the pregnancy (e.g., Ultrasound, Biophysical Profile, Non-Stress Test, Contraction Stress Test)	
LOINC Code	Opt	Description
57078-8	R	Antenatal testing and surveillance

2425

```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5' />
    <id root=' ' extension=' ' />
    <code code='57078-8' displayName='ANTENATAL TESTING AND SURVEILLANCE'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </component>
```

2430

Figure 6.3.3.5.6-1: Specification for and Surveillance Section

Add Section 6.3.3.5.7

2435 6.3.3.5.7 Coded Antenatal Testing and Surveillance Section

1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5.1	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5	
General Description	The Antenatal Testing and Surveillance section shall contain a narrative and coded description of reports and data from tests and surveillance performed during the pregnancy (e.g., Ultrasound, Biophysical Profile, Non-Stress Test, Contraction Stress Test). It shall contain an Antenatal Testing and Surveillance Battery.	
LOINC Code	Opt	Description
57078-8	R	ANTENATAL TESTING AND SURVEILLANCE
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10	R	Antenatal Testing and Surveillance Battery

```

2440 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.5.1' />
<id root=' ' extension=' ' />
<code code='57078-8' displayName='ANTENATAL TESTING AND SURVEILLANCE'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
  <!-- Required Antenatal Testing and Surveillance Battery -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10' />
  :
</entry>
</component>

```

2455 **Figure 6.3.3.5.7-1: Specification for Coded Antenatal Testing and Surveillance Section**

Add Section 6.3.3.5.8 (Diagnosis - Removed 2011-09 at the request of QRPH)

6.3.3.5.8 Intentionally blank

2460 *Add Section 6.3.3.5.9 (TNM Stage – removed 2011-09 at the request of QRPH)*

6.3.3.5.9 Intentionally blank

Add Section 6.3.3.5.10 (Cancer Supporting Documentation - removed 2011-09 at the request of QRPH)

2465 **6.3.3.5.10 Intentionally blank**

Add Section 6.3.3.5.11. (Added 2011-09 from QRPH EHCP Profile)

6.3.3.5.11 Hearing Screening Coded Results

2470 The Hearing Screening Coded Results section SHALL contain the hearing screening results of pass or refer for the right ear and pass or refer for the left ear, expressed as LOINC® codes as well as the coded methodology to complete the screening. Coded methodology includes (LOINC 54106-0) Automated Auditory Brainstem Response, Auditory Brainstem Response, Otoacoustic Emissions, Transient Otoacoustic Emissions, and Distortion Product Otoacoustic Emissions. If the methodology is unknown, the coded result of unknown method SHALL be used. Where the screening results are not available, the reason the results are not available SHALL be present. This could include unsuccessful, technical fail; not performed, not performed, medical exclusion. The Hearing Screening Coded Results section is required.

Template ID	1.3.6.1.4.1.19376.1.7.3.1.1.15.3.2	
Parent Template	Coded Results (1.3.6.1.4.1.19376.1.5.3.1.3.28)	
General Description	The Hearing Screening Code Results section SHALL include at least one observation entry describing the hearing screening results as described in the Entry Content Module. Where there are no hearing screening results performed, then the reason SHALL be indicated	
LOINC Code	Opt	Description
30954-2	R	Relevant diagnostic tests/laboratory data
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.19	R	Procedure Entry
1.3.6.1.4.1.19376.1.5.3.1.4.4	R2	References Entry
1.3.6.1.4.1.19376.1.5.3.1.4.13	R	Simple Observation

2480 **6.3.3.5.11.1 Parent Template**

The parent of this template is Coded Results.

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.28' />
    <id root=' ' extension=' ' />
    <code code='30954-2' displayName='Relevant diagnostic tests/laboratory data'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required Procedure Entry element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19' />
      :
    </entry>
    <entry>
      :
      <!-- Required if known References Entry element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Simple Observation element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
      :
    </entry>
  </section>

```

Figure 6.3.3.5.11-1: Hearing Screening Coded Results Section

Add Section 6.3.3.6

6.3.3.6 Plans of Care

2515

Add Section 6.3.3.6.1

6.3.3.6.1 Care Plan

Add Section 6.3.3.6.2

6.3.3.6.2 Assessment and Plan

Add Section 6.3.3.6.3

2520

6.3.3.6.3 Discharge Disposition

Add Section 6.3.3.6.4

6.3.3.6.4 Discharge Diet

Add Section 6.3.3.6.5

6.3.3.6.5 Advance Directives

2525

Add Section 6.3.3.6.6

6.3.3.6.6 Coded Advance Directives

Add Section 6.3.3.6.7

6.3.3.6.7 Transport Mode

Add Section 6.3.3.6.8

2530

6.3.3.6.8 Procedure Care Plan Status Report Section

1.3.6.1.4.1.19376.1.5.3.1.1.9.45

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.45	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.1.9.40	
General Description	The procedure care plan status report section shall contain a description of the progress towards completing expectations for care including actions completed in fulfillment of proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient prior to the procedure.	
LOINC Code	Opt	Description
18776-5	R	TREATMENT PLAN

Sample Procedure Care Plan Status Report Section

```

2535 <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.40' />
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.45' />
        <id root=' ' extension=' ' />
        <code code='18776-5' displayName='TREATMENT PLAN'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
          Text as described above
        </text>
      </section>
    </component>
  
```

Add Section 6.3.3.6.9

6.3.3.6.9 Health Maintenance Care Plan Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.50

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.50	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.3.31	
General Description	The health maintenance care plan section shall contain a description of the expectations for wellness care including proposals, goals, and order requests for monitoring, tracking, or improving the lifetime condition of the patient with goals of educating the patient on how to reduce the modifiable risks of the patient's genetic, behavioral, and environmental pre-conditions and otherwise optimizing lifetime outcomes.	
LOINC Code	Opt	Description
18776-5	R	TREATMENT PLAN

Sample Health Maintenance Care Plan Section

```

2555 <component>
      <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31' />
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.50' />
        <id root=' ' extension=' ' />
        <code code='18776-5' displayName='TREATMENT PLAN'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
          Text as described above
        </text>
      </section>
    </component>
  
```

Add Section 6.3.3.6.10

2570

6.3.3.6.10 Health Maintenance Care Plan Status Report Section

1.3.6.1.4.1.19376.1.5.3.1.1.9.41

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.41	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.1.9.50	
General Description	The health maintenance status report section shall contain a description of the progress towards completing expectations for care including actions completed in fulfillment of proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient.	
LOINC Code	Opt	Description
18776-5	R	TREATMENT PLAN

2575

Sample Health Maintenance Care Plan Status Report Section

```
<component>
<section>    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.50'>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.41'>
        <id root=' ' extension=' '/>
        <code code='18776-5' displayName='TREATMENT PLAN'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'>
            <text>
                Text as described above
            </text>
        </code>
    </templateId>
</section>
</component>
```

2580

2585

Add Section 6.3.3.6.11

2590

6.3.3.6.11 Provider Orders Section

1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1	
General Description	The provider orders shall contain a list of all pertinent orders from healthcare providers.	
LOINC Code	Opt	Description
46209-3	R	PROVIDER ORDERS
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.7	C	<u>Medications</u> Medications entries shall appear for all ordered medications when present. These entries shall be in intent mood.
1.3.6.1.4.1.19376.1.5.3.1.4.19	C	<u>Procedure</u> Procedure entries shall appear for all ordered procedures when present. These entries shall be in intent mood.
1.3.6.1.4.1.19376.1.5.3.1.4.14	O	<u>Encounter</u> Encounter entries should appear for all ordered encounters. These entries shall be in promise or appointment request mood.

1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1	O	Observation Requests Observation request entries should appear for all ordered observations. These entries shall appear in intent mood.
------------------------------------	---	--

Sample Provider Orders Section

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.2.1'>
    <id root=' ' extension=' '/>
    <code code='46209-3' displayName='PROVIDER ORDERS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required if known Medications element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7'>
      :
    </entry>
    <entry>
      :
      <!-- Required if known Procedure element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19'>
      :
    </entry>
    <entry>
      :
      <!-- Optional Encounter element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.14'>
      :
    </entry>
    <entry>
      :
      <!-- Optional Observation Requests element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1'>
      :
    </entry>
  </section>
</component>

```

2630 Add Section 6.3.3.6.12

6.3.3.6.12 Birth Plan Section 1.3.6.1.4.1.19376.1.5.3.1.1.21.2.1

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.21.2.1	
Parent Template		
General Description	The Birth Plan section shall contain a narrative description of the patient's requests and expectations with respect to care she is expecting during the labor and delivery process.	
LOINC Code	Opt	Description
57079-6	R	Birth plan

2635 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.2.1' />
 <id root=' ' extension=' ' />
 <code code='57079-6' displayName='Birth plan'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 </component>

2640

Figure 6.3.3.6.12-1: Specification for Birth Plan Section

2645 *Add Section 6.3.3.6.13*

6.3.3.6.13 Immunization Recommendations 1.3.6.1.4.1.19376.1.5.3.1.1.18.3.1

Add Section 6.3.3.6.14

6.3.3.6.14 Patient Education Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.38

Template Id	1.3.6.1.4.1.19376.1.5.3.1.1.9.38	
General Description	The patient education section shall contain a description of the patient education the patient received as well as the results of the education.	
LOINC Code	Opt	Description
34895-3	R	EDUCATION NOTE
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.12.2	R	Immunization Recommendation Entry At least one Immunization Plan Entry shall be present in Proposal mood to indicate what the proposed care is for the patient. Other Immunization Plan entries may appear in intent mood to indicate the current plan.

2650 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.38' />
 <id root=' ' extension=' ' />
 <code code='34895-3' displayName='EDUCATION NOTE'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 </section>
 </component>

2655

2660

Figure 6.3.3.6.14-1: Specification for Patient Education and Consents Section

2665 *Add Section 6.3.3.6.15*

6.3.3.6.15 Coded Care Plan Section 1.3.6.1.4.1.19376.1.5.3.1.3.36

Template ID	1.3.6.1.4.1.19376.1.5.3.1.3.31	
Parent Template	2.16.840.1.113883.10.20.1.10	
General Description	The care plan section shall contain a narrative description of the expectations for care including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient.	
LOINC Code	Opt	Description
18776-5	R	TREATMENT PLAN
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1	O	<p><u>Observation Requests</u> The care plan may include observation requests in intent, goal or proposal mood to identify intended observations that are part of the care plan, goals of the plan, or proposed observations (e.g., from clinical decision support).</p>
1.3.6.1.4.1.19376.1.5.3.1.4.7	O	<p><u>Medication</u> The care plan may include medication entries to identify those medications that are or are proposed to be part of the care plan.</p>
1.3.6.1.4.1.19376.1.5.3.1.4.12	O	<p><u>Immunization</u> The care plan may include immunization entries to identify those immunizations that are or are proposed to be part of the care plan.</p>
1.3.6.1.4.1.19376.1.5.3.1.4.19	O	<p><u>Procedure</u> The care plan may include procedure entries to identify those procedures that are or are proposed to be part of the care plan.</p>
1.3.6.1.4.1.19376.1.5.3.1.4.14	O	<p><u>Encounter</u> The care plan may include encounter entries in to identify those encounters that are or are proposed to be part of the care plan.</p>

```

<component>
  <section>
    <templateId root='2.16.840.1.113883.10.20.1.10' />
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.36' />
    <id root=' ' extension=' ' />
    <code code='18776-5' displayName='TREATMENT PLAN'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Optional Observation Requests element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Medication element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Immunization element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.12' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Procedure element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19' />
      :
    </entry>
    <entry>
      :
      <!-- Optional Encounter element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.14' />
      :
    </entry>
  </section>
</component>

```

Figure 6.3.3.6.15-1: Specification for Care Plan Section**6.3.3.6.16 Diet and Nutrition Section 1.3.6.1.4.1.19376.1.5.3.1.1.20.2.2**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.2	
General Description	This shall contain a narrative description of the diet restrictions necessary due to disease.	
LOINC Code	Opt	Description
XX-DietAndNutrition	R	Diet and nutrition

2715 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.2.2' />
 <id root=' ' extension=' ' />
 <code code='XXDiet-Restrictions' displayName='Diet and nutrition'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 </section>
 </component>

2720

Figure 6.3.3.6.16-1: Specification for Diet Restrictions Section**6.3.3.6.17 Intake and Output Section 1.3.6.1.4.1.19376.1.5.3.1.1.20.2.3**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.20.2.3	
General Description	This section shall contain a narrative description of specific fluid inputs or fluid outputs for the patient.	
LOINC Code	Opt	Description
XX-IntakeAndOutput	R	Intake and output

2730 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.2.3' />
 <id root=' ' extension=' ' />
 <code code='XX-FluidManagement' displayName='Intake and output'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 </section>
 </component>

2735

Figure 6.3.3.6.17-1: Specification for Fluid Management Section

Add Section 6.3.3.6.18 (Cancer Course of Treatment – removed 2011-09 at the request of QRPH)

6.3.3.6.18 Intentionally blank

2745

Add Section 6.3.3.6.19 (Cancer Treatment Plan – removed 2011-09 at the request of QRPH)

6.3.3.6.19 Intentionally blank

Add section 6.3.3.6.20

2750

6.3.3.6.20 Procedure Care Plan Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.40

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.40	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.3.31 (1.3.6.1.4.1.19376.1.5.3.1.3.31)	
General Description	The procedure care plan section shall contain a description of the expectations for care including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient prior, during and after a procedure with goals of educating the patient, reducing the modifiable risks of the procedure and anesthesia and otherwise optimizing the outcomes. The care plan will often be updated immediately following the addition of new impressions during the course of pre-procedure evaluation.	
LOINC Code	Opt	Description
18776-5	R	TREATMENT PLAN
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1	O	<u>Observation Requests</u> The care plan may include observation requests in intent, goal or proposal mood to identify intended observations that are part of the care plan, goals of the plan, or proposed observations (e.g., from clinical decision support).
1.3.6.1.4.1.19376.1.5.3.1.4.7	O	<u>Medication</u> The care plan may include medication entries to identify those medications that are or are proposed to be part of the care plan.
1.3.6.1.4.1.19376.1.5.3.1.4.12	O	<u>Immunization</u> The care plan may include immunization entries to identify those immunizations that are or are proposed to be part of the care plan.
1.3.6.1.4.1.19376.1.5.3.1.4.19	O	<u>Procedure</u> The care plan may include procedure entries to identify those procedures that are or are proposed to be part of the care plan.
1.3.6.1.4.1.19376.1.5.3.1.4.14	O	<u>Encounter</u> The care plan may include encounter entries in to identify those encounters that are or are proposed to be part of the care plan.

2755

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31' />
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.40' />
    <id root=' ' extension=' '/>
    <code code='18776-5' displayName='TREATMENT PLAN'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

2760

2765

Figure 6.3.3.6.20-1: Sample Care Plan Section

2770 *Add section 6.3.3.6.21 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)*

6.3.3.6.21 Protocols Used Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.5

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.5	
General Description	The Protocols Used section describes the protocol used by EMS personnel to direct the clinical care of the patient.	
LOINC Code	Opt	Description
52019-7	R	DESCRIPTION OF SERVICES PERFORMED TO SUPPORT LEVEL OF SERVICE

2775 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.5' />
 <id root=' ' extension=' ' />
 <code code='52019-7' displayName='DESCRIPTION OF SERVICES PERFORMED TO SUPPORT LEVEL OF SERVICE'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 </section>
 </component>

2780

2785

Figure 6.3.3.6.21-1: Sample Protocols Used Section

2790 *Add section 6.3.3.6.22 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)*

Modified by CP PCC 0205

6.3.3.6.22 Invasive Airway Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.7

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.7	
General Description	The Invasive Airway section describes if, and what type, of advanced airway used. The Invasive Airway section is derived from DEEDS LL1832-6 NEMESIS_45_protocol used / Airway/Airway-failed/Airway-obstruction/foreign body/Airway-paralytic (RSI), Airway-Rapid Sequence Induction (RSI-Paralytic)/Airway-sedation assisted (nonparalytic)/Cardiac arrest- asystole, etc. PROTOCOLS USED	
LOINC Code	Opt	Description
67537-1	R	PROTOCOLS USED NEMESIS

```

2795 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.7' />
        <id root=' ' extension=' ' />
        <code code='67537-1' displayName='PROTOCOLS USED NEMESIS'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
    </section>
</component>

```

2805 **Figure 6.3.3.6.22-1: Sample Invasive Airway Section**

Add section 6.3.3.6.23 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

2810 **6.3.3.6.23 Ventilator Usage Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.11**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.11	
General Description	The Ventilator Usage section describes	
LOINC Code	Opt	Description
20124-4	R	VENTILATION MODE

```

2815 <component>
    <section>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.11' />
        <id root=' ' extension=' ' />
        <code code='2012404' displayName='VENTILATION MODE'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <text>
            Text as described above
        </text>
    </section>
</component>

```

2820

Figure 6.3.3.6.23-1: Sample Ventilator Usage Section

2825

*Add Section 6.3.3.7***6.3.3.7 Administrative and Other Information***Add Section 6.3.3.7.1***6.3.3.7.1 Payers***Add Section 6.3.3.7.2*

2830 **6.3.3.7.2 Referral Source***Add Section 6.3.3.7.3***6.3.3.7.3 Transport Mode***Add Section 6.3.3.7.4***6.3.3.7.4 ED Disposition**2835 *Add Section 6.3.3.7.5 (Cancer Payers – Removed 2011-09 at the request of QRPH)***6.3.3.7.5 Intentionally blank***Add Section 6.3.3.7.6 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)*2840 **6.3.3.7.6 Sending Facility Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.1**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.1	
General Description	The Sending Facility section contains the name and address of the healthcare facility that is sending the patient for transport.	
LOINC Code	Opt	Description
52023-9	R	ORIGINATION SITE NAME AND ADDRESS

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.1' />
    <id root=' ' extension=' '/>
    <code code='52023-9' displayName='ORIGINATION SITE NAME AND ADDRESS'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

2845

2850

Figure 6.3.3.7.6-1: Sample Sending Facility Section2855 *Add Section 6.3.3.7.7 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)*

6.3.3.7.7 Receiving Facility Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.2

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.2	
General Description	The Receiving Facility section contains the name and address of the healthcare facility that is receiving the transported patient.	
LOINC Code	Opt	Description
52026-2	R	DESTINATION SITE NAME & ADDRESS

2860

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.2' />
    <id root=' ' extension=' ' />
    <code code='52026-2' displayName='DESTINATION SITE NAME & ADDRESS'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2865

2870

Figure 6.3.3.7.7-1: Sample Receiving Facility Section

Add Section 6.3.3.7.8 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

2875

Modified by CP PCC 0205

6.3.3.7.8 Mass Casualty Incident Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.3

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.3	
General Description	The Mass Casualty Incident Section indicates if this event would be considered a mass casualty incident overwhelming existing EMS and ED resources.	
LOINC Code	Opt	Description
67490-3	R2	Mass casualty incident NEMSIS

2880

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.3' />
    <id root=' ' extension=' ' />
    <code code='67490-3' displayName=' Mass casualty incident NEMSIS '
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2885

2890

Figure 6.3.3.7.8-1: Sample Mass Casualty Incident Section

Add Section 6.3.3.7.9 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

6.3.3.7.9 Unit Response Level Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.4

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.4	
General Description	The Unit Response Level section describes the level of service provided for this transport.	
LOINC Code	Opt	Description
51995-9	R	RATIONALE FOR TYPE OF TRANSPORT

2895

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.4' />
    <id root=' ' extension=' ' />
    <code code='51995-9' displayName='RATIONALE FOR TYPE OF TRANSPORT'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2900

2905

Figure 6.3.3.7.9-1: Sample Unit Response Level Section

2910

Add Section 6.3.3.7.10 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

6.3.3.7.10 Extra Attendants Information Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.6

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.6	
General Description	The Protocols Used section describes the protocol used by EMS personnel to direct the clinical care of the patient.	
LOINC Code	Opt	Description
52074-2	R2	EXTRA ATTENDANTS INFORMATION

2915

2920

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.6' />
    <id root=' ' extension=' ' />
    <code code='52074-2' displayName='EXTRA ATTENDANTS INFORMATION'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

Figure 6.3.3.7.10-1: Sample Extra Attendants Information Section

2925

Add Section 6.3.3.7.11 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)

Modified by CP PCC 0205

6.3.3.7.11 Provider Level Section 1.3.6.1.4.1.19376.1.5.3.1.1.25.2.9

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.25.2.9	
General Description	The Provider Level section describes the certification or licensure level of the healthcare provider.	
LOINC Code	Opt	Description
71580-5	R	Crew member level NEMSIS

2930

```
<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.25.2.9' />
    <id root=' ' extension=' '/>
    <code code='71580-5' displayName=' Crew member level NEMSIS '
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>
```

2935

2940

Figure 6.3.3.7.11-1: Sample Provider Level Section

Add Section 6.3.3.8

2945

6.3.3.8 Interventions

This section contains section content modules that describe interventions, procedures, therapeutic treatments, et cetera, performed on the patient.

Add Section 6.3.3.8.3

2950

6.3.3.8.3 Procedures and Interventions Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11	
General Description	The Procedures and Interventions section shall contain a narrative description of the actions performed by a clinician.	
LOINC Code	Opt	Description
29554-3	R	PROCEDURE

Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.19	R	Procedure This entry provides coded values for procedures performed during the encounter.

2955	<component> <section> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.11'> <id root=' ' extension=' ' /> <code code='X-PROC 29554-3' displayName='PROCEDURES PERFORMED' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' /> <text> Text as described above </text> <entry> : <!-- Required Procedure element --> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19'> : </entry> </section> </component>
2960	
2965	
2970	

Figure 6.3.3.8.3-1: Specification for Procedures and Interventions Section*Add Section 6.3.3.8.4 (Added 2011-06 from the PCC Transport Record Summary Profiles supplement)*

2975

6.3.3.8.4 Intravenous Fluids Administered Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.6

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.6	
General Description	The intravenous fluids administered section shall contain a narrative description of fluids administered to a patient during the course of an encounter. It may include entries for IV fluid administration as described in the Entry Content Module.	
LOINC Code	Opt	Description
57072-1	R	Intravenous fluids administered

Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.13.3.2	R	Intravenous Fluids

```

2980 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.6' />
<id root=' ' extension=' ' />
<code code='57072-1' displayName='Intravenous fluids administered'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
  <!-- Required Intravenous Fluids element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.2' />
  :
</entry>
</section>
</component>

```

2995 **Figure 6.3.3.8.4-1: Specification for Intravenous Fluids Administered Section**

3000 **Add Section 6.3.3.9**

6.3.3.9 Impressions

This section contains section content modules that describe assessments, impressions, diagnoses, or other reporting of clinical opinions or judgment.

3005 **Add Section 6.3.3.9.1**

6.3.3.9.1 Pre-procedure Impressions Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.42 (Deprecated)

3005 **Add Section 6.3.3.9.2**

6.3.3.9.2 Pre-procedure Risk Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.9.44

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.9.44	
General Description	The pre-procedure risk section shall contain a description of the risks the patient faces because of the planned procedure and associated anesthesia, especially in the context of modifiable risks identified by patient findings. It shall include entries for patient risks as described in the Entry Content Module.	
LOINC Code	Opt	Description
11450-4	R	PROBLEM LIST
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.5	R	Problem Entry

The parent of this template is 1.3.6.1.4.1.19376.1.5.3.1.3.6

```

3010 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.6' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.9.44' />
<id root=' ' extension=' ' />
<code code='11450-4' displayName='PROBLEM LIST'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
<entry>
  :
  <!-- Required Problem Entry element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5' />
  :
</entry>
</section>
</component>

```

Figure 6.3.3.9.2-1: Specification for Pre-procedure Risk Assessment Section*Add Section 6.3.3.9.3***6.3.3.9.3 Antepartum Visit Summary Flowsheet Section****1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.2**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.2	
General Description	This section is a running history of the most important elements noted for a pregnant woman.	
LOINC Code	Opt	Description
57059-8	R	Pregnancy visit summary
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.13	R	<u>Simple Observation</u> The flowsheet contains one simple observation to represent the Prepregnancy Weight. This observation SHALL be valued with the LOINC code 8348-5, BODY WEIGHT^PRE PREGNANCY-MASS-PT-QN-MEASURED. The value SHALL be of type PQ. The units may be either "lb_av" or "kg".
1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2	R	<u>Antepartum Flowsheet Panel</u> Other entries on the flowsheet are "batteries" which represent a single visit.

3035

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.2.2' />
    <id root=' ' extension=' ' />
    <code code='57059-8' displayName='Pregnancy visit summary'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required Simple Observation element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
      :
    </entry>
    <entry>
      :
      <!-- Required Antepartum Flowsheet Panel element -->
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2' />
      :
    </entry>
  </section>
</component>

```

Figure 6.3.3.9.3-1: Specification for Antepartum Visit Summary Flowsheet Section*Add Section 6.3.3.9.4***6.3.3.9.4 Progress Note Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.7**

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.7	
General Description	The Progress Note section shall contain a narrative description of the sequence of events from initial assessment to discharge for an encounter.	
LOINC Code	Opt	Description
18733-6	R	SUBSEQUENT EVALUATION NOTE (ATTENDING PHYSICIAN)

3065

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.7' />
    <id root=' ' extension=' ' />
    <code code='18733-6' displayName='SUBSEQUENT EVALUATION NOTE (ATTENDING PHYSICIAN)'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

3070

3075

Figure 6.3.3.9.4-1: Specification for Progress Note Section*Add Section 6.3.3.9.5*

6.3.3.9.5 ED Diagnosis Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.9

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.9	
General Description	The ED diagnosis section shall contain a narrative description of the conditions that were diagnosed or addressed during the ED course, as well as those active conditions that modify the complexity of the patient encounter. It should include entries for patient conditions as described in the Entry Content Module.	
LOINC Code	Opt	Description
11301-9	R	ED DIAGNOSIS
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.4.5	R	Problem Entry

3085 <component>
 <section>
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.9' />
 <id root=' ' extension=' ' />
 <code code='11301-9' displayName='ED DIAGNOSIS'
 codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
 <text>
 Text as described above
 </text>
 <entry>
 :
 <!-- Required Problem Entry element -->
 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5' />
 :
 </entry>
 </section>
 </component>

3090

3095

3100

Figure 6.3.3.9.5-1: Specification for ED Diagnosis Section

Add Section 6.3.3.9.6

6.3.3.9.6 Acuity Assessment Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.2

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.2	
General Description	The Acuity Assessment section contains a description of the acuity of the patient upon presentation to the Emergency department.	
LOINC Code	Opt	Description
11283-9	R	ACUITY ASSESSMENT
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1	R	Acuity This entry provides coded values giving the triage acuity.

```

3105 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.2' />
<id root=' ' extension=' ' />
3110 <code code='11283-9' displayName='ACUITY ASSESSMENT'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
3115 <entry>
  :
  <!-- Required Acuity element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1' />
  :
</entry>
3120 </section>
</component>

```

Figure 6.3.3.9.6-1: Specification for Acuity Assessment Section

3125 *Add Section 6.3.3.9.7*

6.3.3.9.7 Assessments Section 1.3.6.1.4.1.19376.1.5.3.1.1.13.2.4

Template ID	1.3.6.1.4.1.19376.1.5.3.1.1.13.2.4	
General Description	The assessments section contains narrative assessments of the patient status.	
LOINC Code	Opt	Description
51848-0	R	ASSESSMENT
Entries	Opt	Description
1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4	O	Nursing Assessments Battery

```

3130 <component>
<section>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.2.4' />
<id root=' ' extension=' ' />
<code code='51848-0' displayName='ASSESSMENT'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text>
  Text as described above
</text>
3135 <entry>
  :
  <!-- Optional Nursing Assessments Battery element -->
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4' />
  :
</entry>
3140 </section>
</component>

```

Figure 6.3.3.9.7-1: Specification for Assessments Section

Add section 6.3.3.10

6.3.3.10 Section Content Modules-Non-categorized

3150 **Please note:** As of 2013, section content modules are no longer being categorized into one of the nine existing categories (6.3.3.1 through 6.3.3.9). Instead, going forward, all section content modules will be placed under the 6.3.3.10 heading.

Add section 6.3.3.10.1. Added 2013-09 from QRPH VRDR supplement.
--

3155 **6.3.3.10.1 VRDR Death Report Section- Section Content Module
(1.3.6.1.4.1.19376.1.7.3.1.3.23.2)**

The sections and clinical statements which have additional implementation guidance further constrained are listed here showing their new IHE template ID.

Table 6.3.3.10.1-1: VRDR Death Report Section

Template Name		VRDR Death Report Section			
Template ID		1.3.6.1.4.1.19376.1.7.3.1.3.23.2			
Parent Template		Death Report Document Body (2.16.840.1.113883.10.20.24.1.2)			
General Description		The VRDR Death Report section shall contain a coded entries describing the decedent's death			
Section Code		64297-5, LOINC, "Death Certificate"			
Opt and Card	Condition	Data Element or Section Name	Template ID	Specification Document	Vocabulary Constraint
Entries					
R[0..1]		Time of Death	2.16.840.1.113883.10.20.24.1.3	HL7 VRDR CDA CH4	
R[1..1]		Location of Death	2.16.840.1.113883.10.20.24.1.4	HL7 VRDR CDA CH4	
O[0..1]	QRPH 3:6.3.3.10.S1 .3	Death Certification	2.16.840.1.113883.10.20.24.1.5	HL7 VRDR CDA CH4	
R[1..1]		Manner of Death	2.16.840.1.113883.10.20.24.1.7	HL7 VRDR CDA CH4	
C[0..1]	QRPH 3: 6.3.3.10.S1.1	Pregnancy Status	2.16.840.1.113883.10.20.24.1.8	HL7 VRDR CDA CH4	
R2[0..1]		Tobacco Use	2.16.840.1.113883.10.20.24.1.9	HL7 VRDR CDA CH4	
R2[0..1]		Injury	2.16.840.1.113883.10.20.24.1.10	HL7 VRDR CDA CH4	
R[1..1]	QRPH 3: 6.3.3.10.S1.4	Death Causal Information	2.16.840.1.113883.10.20.24.1.6	HL7 VRDR CDA CH4	

R[1..1]		Autopsy Performance	2.16.840.1.113883.10.20.24.1. 11	HL7 VRDR CDA CH4	
C[0..1]	QRPH 3: 6.3.3.10.S1.2	Autopsy Results	2.16.840.1.113883.10.20.24.1. 13	HL7 VRDR CDA CH4	
O[0..1]		Coroner Referral	2.16.840.1.113883.10.20.24.1. 14	HL7 VRDR CDA CH4	
R[1..1]		Coroner Case Transfer	2.16.840.1.113883.10.20.24.1. 12	HL7 VRDR CDA CH4	
R[1..1]		Death Location Type	1.3.6.1.4.1.19376.1.7.3.1.4.23. 2	QRPH 3: 6.3.4.E2	
R[1..1]		Death Pronouncement	1.3.6.1.4.1.19376.1.7.3.1.4.23. 1	QRPH 3: 6.3.4.E1	

3160

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.5' />
    <id root=' ' extension=' ' />
    <code code='64297-5/displayName='Death certificate'
          codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required Time of Death -->
      <templateId root='2.16.840.1.113883.10.20.24.1.3' />
      :
    </entry>
    <entry>
      :
      <!-- Required Location of Death -->
      <templateId root='2.16.840.1.113883.10.20.24.1.4' />
      :
    </entry>
    <entry>
      :
      <!--Optional Death Certification -->
      <templateId root='2.16.840.1.113883.10.20.24.1.5' />
      :
    </entry>
    <entry>
      :
      <!--Required Manner of Death -->
      <templateId root='2.16.840.1.113883.10.20.24.1.7' />
      :
    </entry>
    <entry>
      :
      <!--Conditional Pregnancy Status -->
      <templateId root='2.16.840.1.113883.10.20.24.1.8' />
      :
    </entry>
    <entry>
      :
      <!--Required if known Tobacco Use -->
      <templateId root='2.16.840.1.113883.10.20.24.1.9' />
      :
    </entry>
    <entry>
      :
    </entry>
  </section>

```

3165

3170

3175

3180

3185

3190

3195

3200

3205

```

3210      <!--Required if known Injury -->
3211      <templateId root='2.16.840.1.113883.10.20.24.1.10' />
3212      :
3213      </entry>
3214      <entry>
3215          :
3216          <!--Required Death Causal Information -->
3217          <templateId root='2.16.840.1.113883.10.20.24.1.6' />
3218          :
3219          </entry>
3220          <entry>
3221              :
3222              <!--Required Autopsy Performance -->
3223              <templateId root='2.16.840.1.113883.10.20.24.1.11' />
3224              :
3225              <!--Conditional Autopsy Results -->
3226              <templateId root='2.16.840.1.113883.10.20.24.1.13' />
3227              :
3228          </entry>
3229          <entry>
3230              :
3231              <!--Optional Coroner Referral -->
3232              <templateId root='2.16.840.1.113883.10.20.24.1.14' />
3233              :
3234          </entry>
3235          <entry>
3236              :
3237              <!--Required Coroner Case Transfer -->
3238              <templateId root='2.16.840.1.113883.10.20.24.1.12' />
3239              :
3240          </entry>
3241          <entry>
3242              :
3243              <!--Required Death Location Type -->
3244              <templateId root=' 1.3.6.1.4.1.19376.1.7.3.1.4.23.2' />
3245              :
3246          </entry>
3247          <entry>
3248              :
3249              <!--Required Death Pronouncement-->
3250              <templateId root=' 1.3.6.1.4.1.19376.1.7.3.1.4.23.1' />
3251              :
3252          </entry>
3253
3254      </section>
3255  </component>

```

Figure 6.3.3.10.1-1: Sample VRDR Death Report Section

3260

6.3.3.10.1.1 Pregnancy Status Entry Condition

The Pregnancy Status clinical statement SHALL be Required if the person is female and in the age range 5 to 75 years.

6.3.3.10.1.2 Autopsy Results Entry Condition

3265 The Autopsy Results clinical statement SHALL be Required if autopsy performed.

6.3.3.10.1.3 Death Certification Entry Condition

The License Number of Person Certifying Death SHALL be reflected in Performer/assigned person.

6.3.3.10.1.4 Death Causal Information Entry Condition

3270 The Name of person completing COD SHALL be reflected in author/assignedAuthor/name.

Add section 6.3.3.10.2 (added 2013-09 from QRPH VRDR supplement).

6.3.3.10.2 Coded Hospital Course Section 1.3.6.1.4.1.19376.1.7.3.1.3.23.1

3275 **Table 6.3.3.10.2-1: Coded Hospital Course Section**

Template Name		Coded Hospital Course Section			
Template ID		1.3.6.1.4.1.19376.1.7.3.1.3.23.1			
Parent Template		Hospital Course Section (1.3.6.1.4.1.19376.1.5.3.1.3.5)			
General Description		The hospital course section shall contain a narrative description and coded entries describing the sequence of events from admission to discharge in a hospital facility.			
Section Code		8648-8, LOINC, HOSPITAL COURSE			
Opt and Card	Condition	Data Element or Section Name	Template ID	Specification Document	Vocabulary Constraint
Entries					
R2[0..1]	HL7	Time of Death	2.16.840.1.113883.10.20.24.1. 3		

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.7.3.1.3.23.1' />
    <id root=' ' extension=' ' />
    <code code='8648-8' displayName='HOSPITAL COURSE'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
    <entry>
      :
      <!-- Required if known Time of Death element -->
      <templateId root='2.16.840.1.113883.10.20.24.1.3' />
      :
    </entry>
  </section>
</component>

```

3295 **Figure 6.3.3.10.2-1: Sample Coded Hospital Course Section**

Add section 6.3.3.10.3 (added 2013-09 from the QRPH HW supplement).

6.3.3.10.3 Resources to Support Goals Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.1

3300 **Table 6.3.3.10.3-1: Resources to Support Goals Section**

Template ID	1.3.6.1.4.1.19376.1.7.3.1.3.24.1	
General Description	The Resources to Support Goals Section shall contain a narrative description of the community, health, and wellness resources available or provided to the patient to support their care plan goals.	
LOINC Code	Opt	Description
46802-5	R	Communication with community resources.knowledge

```

<component>
  <section>
    <templateId root='1.3.6.1.4.1.19376.1.7.3.1.3.24.1' />
    <id root=' ' extension=' ' />
    <code code='46802-5' displayName='Communication with community resources.knowledge'
      codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
    <text>
      Text as described above
    </text>
  </section>
</component>

```

3310 **Figure 6.3.3.10.3-1: Sample Resources to Support Goals Section**

Add section 6.3.3.10.4 (added 2013-09 from the QRPH HW supplement).

6.3.3.10.4 Healthy Weight Care Plan Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.2

Table 6.3.3.10.4-1: Healthy Weight Care Plan Section

Template ID	1.3.6.1.4.1.19376.1.7.3.1.3.24.2	
Parent Template	1.3.6.1.4.1.19376.1.5.3.1.3.31	
General Description	<p>The healthy weight care plan section shall contain a narrative description of the expectations for care for healthy weight management including proposals, goals, and order requests for monitoring, tracking, or improving the condition of the patient. The Healthy Weight care plan includes the following Goal Setting documentation:</p> <ul style="list-style-type: none"> Identification of goals for behavior change (increasing healthy behaviors and/or decreasing unhealthy behaviors) that are appropriate for the patient based on discussion during the visit and patient-reported readiness to change. Messaging related to an ideal (targeted) level for the behavior Goal selection may be selected from structured lists or selected in an open-ended manner Documentation of barriers and supports to attaining selected goals, may be selected from structured lists or selected in an open-ended manner Monitoring of progress against goals set during previous visits 	
LOINC Code	Opt	Description
18776-5	R	PATIENT PLAN OF CARE

3320	<component> <section> <templateId root='2.16.840.1.113883.10.20.1.10' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.3.31' /> <templateId root='1.3.6.1.4.1.19376.1.7.3.1.3.24.2' /> <id root=' ' extension=' ' /> <code code='18776-5' displayName='PATIENT PLAN OF CARE' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' /> <text> Text as described above </text> </section> </component>
3325	
3330	

Figure 6.3.3.10.4-1: Sample Healthy Weight Care Plan Section

<i>Replace the following section 6.3.3.10.5:</i>
--

3335

6.3.3.10.5 Occupational Data for Health Section 1.3.6.1.4.1.19376.1.7.3.1.3.24.1

Table 6.3.3.10.5-1: Occupational Data for Health Section

Template Name		Occupational Data for Health			
Template ID		1.3.6.1.4.1.19376.1.7.3.1.3.24.1			
Parent Template					
General Description		<p>The Occupational Data for Health section shall contain a narrative description of the person's employment status, retirement status, combat zone work, and usual occupation, as well as the person's history of employment. Employment information includes occupation and industry, supervisory level, and the employer's name and location. It should also include compensation and sector employment type and work schedule with hours per day and days per week, and may include job duties, occupational hazards information.</p> <p>When represented in a document containing a Social History section, the Occupational Data for Health section shall be encoded as a sub-section of the Social History section</p>			
Section Code		74166-0, LOINC, "Occupational Data for Health"			
Author		If not the author from the encompassing context, include author. Role and entity must be specified if not inherited.			
Informant		If not the informant from the encompassing context, include informant. Role and entity must be specified if not inherited.			
Subject		If not the subject from the encompassing context, include subject. Role and entity must be specified if not inherited.			
Opt and Card	Condition	Data Element or Section Name	Template ID	Specification Document	Vocabulary Constraint
Entries					
R2 [0..*]		History of Employment Status Observation	1.3.6.1.4.1.19376.1.7.3.1.4.24.18	CDA Content Module: 6.3.4.64	
R2 [0..1]		Usual Occupation Observation	1.3.6.1.4.1.19376.1.7.3.1.4.24.20	CDA Content Module: 6.3.4.69	
R2 [0..*]		Past or Present Occupation Observation	1.3.6.1.4.1.19376.1.7.3.1.4.24.19	CDA Content Module: 6.3.4.66	
R2 [0..*]		Date of Retirement Observation	1.3.6.1.4.1.19376.1.7.3.1.4.24.12	CDA Content Module: 6.3.4.bb	
R2 [0..*]		Combat Zone Period Observation	1.3.6.1.4.1.19376.1.7.3.1.4.24.22	CDA Content Module: 6.3.4.cc	

6.3.3.10.5.1 Occupational Data for Health Section < 74166-0>

3340

[section: templateId 1.3.6.1.4.1.19376.1.7.3.1.3.24.31 (open)]

The Occupational Data for Health section describes all aspects of the subject's employment history. It may contain the history of employment status, the usual occupation (longest held occupation) and related observations, the self-identified date of retirement, any time periods spent working in a combat zone, and the job history.

- 3345 1. **SHALL** contain exactly one [1..1] **templateId** such that it
 a. **SHALL** contain exactly one [1..1]
 @root="1.3.6.1.4.1.19376.1.7.3.1.3.24.31".
 2. **SHALL** contain exactly one [1..1] **code/@code**="74166-0" Occupational Data
 (CodeSystem: LOINC 2.16.840.1.113883.6.1).
3350 3. **SHALL** contain exactly one [1..1] **title**.
 4. **SHALL** contain exactly one [1..1] **text**.
 5. **SHOULD** contain zero or more [0..*] History of Employment Status Observation **entry**
 6. **SHOULD** contain at least one [1..1] Usual Occupation Observation **entry**
 7. **SHOULD** contain at least one [1..*] Past or Present Occupation Observation entry
 8. **SHOULD** contain zero or one [0..*] Date of Retirement Observation entry
 9. **SHOULD** contain zero or more [0..*] Combat Zone Period Observation
- 3355

```

<section>
...
3360    <!-- Sub section for Occupational Data For Health -->
    <component>
        <section>
            <templateId root="2.16.840.1.113883.10.20.22.2.17"/>
            <!-- ODH SECTION TEMPLATE ID-->
            <templateId root="1.3.6.1.4.1.19376.1.7.3.1.3.24.31"/>
            <code code="74166-0" codeSystem="2.16.840.1.113883.6.1"
codeSystemVersion="0" codeSystemName="LOINC" displayName="Occupational
Data for Health"/>
            <text>...</text>
            <entry>
                :
3365                <!-- HISTORY OF EMPLOYMENT STATUS OBSERVATION ENTRY TEMPLATE
ID-->
                <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.24.18"/>
                :
3370                <!-- USUAL OCCUPATION OBSERVATION ENTRY TEMPLATE ID-->
                <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.24.20"/>
                :
3375                <!-- PAST OR PRESENT OCCUPATION OBSERVATION ENTRY TEMPLATE
ID-->
                <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.24.19"/>
                :
3380                <!-- DATE OF RETIREMENT OBSERVATION ENTRY TEMPLATE ID-->
                <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.24.12"/>
                :
3385                <!-- COMBAT ZONE PERIOD OBSERVATION ENTRY TEMPLATE ID-->
                <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.24.22"/>
                :
3390            </entry>
        </section>
    </component>
...
3395 </section>

```

Figure 6.3.3.10.5-1: Occupational Data for Health Section example

6.3.4 CDA Entry Content Modules

Please note: Section 6.3.4.1 through 6.3.4.24 are defined in IHE PCC TF-2: 6.3.4.

3400

Add Section 6.3.4.25

6.3.4.25 Family History Observation 1.3.6.1.4.19376.1.5.3.1.4.13.3

A family history observation is a [Simple Observation](#) that uses a specific vocabulary, and inherits constraints from CCD®⁴. Family history observations are found inside [Family History Organizers](#).

3405

6.3.4.25.1 Standards

CCD [ASTM/HL7 Continuity of Care Document](#)

6.3.4.25.2 Parent Template

The parent of this template is [Simple Observation](#). This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.22

3410

6.3.4.25.3 Specification

```
<observation typeCode='OBS' moodCode='EVN'>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.3' />
<templateId root='2.16.840.1.113883.10.20.1.22' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.3' />
<id root=' ' extension=' '/>
<code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' ' />
<text><reference value="#xxx" /></text>
<statusCode code='completed' />
<effectiveTime value=' ' />
<repeatNumber value=' ' />
<value xsi:type='CD' .../>
<interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />
<methodCode code=' ' codeSystem=' ' codeSystemName=' ' />
<targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />
</observation>
```

3415

3420

3425

Figure 6.3.4.25.3-1: Family History Specification

6.3.4.25.4 <templateId root='2.16.840.1.113883.10.20.1.22' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.3' />

3430

The <templateId> elements identify this observation as a family history observation, and shall be present as shown above.

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

The <code> indicates the type of observation made (e.g., Diagnosis, et cetera). See the code element in the Problem Entry for suggested values.

3435

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

The <value> element indicates the information (e.g., diagnosis) of the family member. See the value element in the Problem Entry for suggested values.

⁴ CCD is the registered trademark of Health Level Seven International.

Add Section 6.3.4.26

3440 **6.3.4.26 Pregnancy History Organizer 1.3.6.1.4.1.19376.1.5.3.1.4.13.5.1**

Defined in IHE PCC TF-2.

Add Section 6.3.4.27

6.3.4.27 EDD Observation 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1

- 3445 The EDD observation reflects the clinician's best judgment about the estimated delivery date of the patient. It can be supported by patient history (e.g., last menses or quickening), physical examination findings (uterine size), or Ultrasound. The observation is a Simple Observation with a supporting entryRelation of another Observation. The supporting observation may in turn have an entryRelation that gives the original observation as a gestational age or date from which the estimated due date is calculated.
- 3450

6.3.4.27.1 Specification

```

<observation classCode='OBS' moodCode='EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1' />
  <statusCode code='completed' />
  <effectiveTime value=' ' />
  <author typeCode='AUT'>
    <time value=' ' />
    <assignedAuthor>
      <id root=' ' extension=' ' />
    </assignedAuthor>
  </author>
  <id root=' ' extension=' ' />
  <code code='11778-8' displayName='DELIVERY DATE-TMSTP-PT-^PATIENT-QN-CLINICAL-ESTIMATED' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
  <text><reference value='id-foo' /></text>
  <value xsi:type='TS' value=' ' />
  <entryRelationship typeCode='SPRT'>
    <observation classCode='OBS' moodCode='EVN'>
      <id root=' ' extension=' ' />
      <statusCode code='completed' />
      <effectiveTime value=' ' />
      <author typeCode='AUT'>
        <time value=' ' />
        <assignedAuthor classCode=' ' />
          <id root=' ' extension=' ' />
        </assignedAuthor>
      </author>
      <code code='[11779-6] (xx-EDD-by-PE) | 11781-2 | (xx-EDD-by-Qck) | (xx-EDD-by-Fund)' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
      <value type='TS' value=' ' />
      <entryRelationship typeCode='DRIV'>
        <observation classCode='OBS' moodCode='EVN'>
          <id root=' ' extension=' ' />
          <statusCode code='completed' />
          <effectiveTime value=' ' />
          <author typeCode='AUT'>
            <time value=' ' />
            <assignedAuthor>
              <id root=' ' extension=' ' />
            </assignedAuthor>
          </author>
          <informant typeCode='INF'>
            <relatedEntity classCode=' ' />
              <id root=' ' extension=' ' />
            </relatedEntity>
          </informant>
          <code code='[8655-2] (xx-ga-by-pe) | 11888-5 | (xx-date-of-qck) | (xx-date-of-fund-umb)' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
          <value type='[PQ|TS]' value=' ' units='week' />
        </observation>
      </entryRelationship>
    </observation>
  </entryRelationship>
</observation>

```

6.3.4.27.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1' />

The <templateId> identifies the observation as a type of Estimated Delivery Date Observation.
 3510 The root attribute SHALL be valued with '1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.1'.

6.3.4.27.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>

EDD observation SHALL comply with the restrictions of the Simple Observation entry. The observation SHALL NOT include repeatNumber, interpretationCode, methodCode, or targetSiteCode as listed below.

3515 **6.3.4.27.4 <code code='11778-8' codeSystem='2.16.840.1.113883.6.1'>**

The <code> element indicates that this is a "clinically estimated" estimated delivery date (for example, this code is used to represent the field on the last line of the EDD section of the ACOG form). This code SHALL be the LOINC code 11778-8. It is good style to include the displayName and codeSystemName to help debugging.

3520 **6.3.4.27.5 <value xsi:type='TS' value=' ' >**

The value of the EDD SHALL be represented as a point in time.

6.3.4.27.6 <author typeCode='AUT'><assignedAuthor><id root=' ' extension=' /></assignedAuthor></author>

3525 There may be multiple clinicians following the patient and authoring the overall document, however the EDD observation has an individual author. For CDA based content, this author SHALL be listed in the CDA header and referenced from the entry by including the id element of the assignedAuthor. For HL7 Version 3 Messages based content, the author SHALL be included in full through this element.

6.3.4.27.7 <author typeCode='AUT'><time value=' '/></author>

3530 The author.time is used to record the time that the author recorded the observation. It SHALL be included.

6.3.4.27.8 <entryRelationship typeCode='SPRT'>

3535 The <entryRelationship> element binds the clinicians estimated EDD to supporting observations by different methods. Supporting observations SHOULD be included. If included, the typeCode SHALL be 'SPRT'. For HL7 Version 3 Messages based content, the element name is <sourceOf> rather than <entryRelationship>, however the semantics, typeCode, and nested elements remain unchanged.

6.3.4.27.9 <observation>**<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>**3540 :
</observation> [1st nesting]

3545 Observations that support the clinical observation SHALL be included if known. These observations are the supporting calculated dates from various methods such as ultrasound dates or dates calculated from LMP (i.e., the left column of fields on the ACOG form). Supporting observations SHALL also conform to the simple observation template. Supporting observations MAY include a different effectiveTime, author, or informant. Supporting observations SHALL

NOT include repeatNumber, interpretationCode, methodCode, or targetSiteCode. (Method is implied by the LOINC code). The templateId SHALL be valued as
 ‘1.3.6.1.4.1.19376.1.5.3.1.4.13’

3550 **6.3.4.27.10 <code code=' ' codeSystem='2.16.840.1.113883.6.1'> [1st nesting]**

Supporting observations SHALL include one of following LOINC values to indicate the method used to calculate the EDD.

Code	Description
11779-6	Delivery date Estimated from last menstrual period
(xx-EDD-by-PE)	DELIVERY DATE-TMSTP-PT-^PATIENT-QN-ESTIMATED FROM CLINICIANS PHYSICAL EXAM
11781-2	Delivery date composite estimate
57063-0	Delivery date Estimated from quickening date
57064-8	Delivery date Estimated from date fundal height reaches umb

6.3.4.27.11 <entryRelationship typeCode='DRIV'>

3555 Observations of supporting EDD should provide observations from which they were derived such as the patient’s last menses, or gestational age value at a point in time.

For HL7 Version 3 Messages based content, the element name is <sourceOf> rather than <entryRelationship>, however the semantics, typeCode, and nested elements remain unchanged.

6.3.4.27.12 <observation>

<templateId root=' '/>

:

</observation> [2st nesting]

3560 Observations that support the calculation of supporting observation SHALL be included if known. These observations are the supporting dates or ages from various methods such as ultrasound gestational age or the date of last Menses (for example, the right column of fields on the ACOG form). Supporting observations SHALL also conform to the simple observation template. Supporting observations MAY include a different effectiveTime, author, or informant. Supporting observations SHALL NOT include repeatNumber, interpretationCode, methodCode, or targetSiteCode. (Method is implied by the LOINC code)

3570 **6.3.4.27.13 <code code=' ' codeSystem='2.16.840.1.113883.6.1'> [2nd nesting]**

This code is used to represent the either the relevant date, or the gestational age observation from which the EDD is derived. The following table lists the relevant LOINC codes for methods used. For observations that record the gestational age the value is recorded as a physical quantity (PQ) with the units of weeks and the activity time should be recorded to indicate the date at which the gestational age was observed. For observations that simply record a date (e.g., LMP) the observation value is recorded as a point in time (TS).

Code	Description	Type
8655-2	DATE LAST MENSTRUAL PERIOD-TMSTP-PT-^PATIENT-QN-REPORTED	TS
11884-4	GESTATIONAL AGE-TIME-PT-^FETUS-QN-ESTIMATED FROM CLINICIANS PHYSICAL EXAM M	PQ
11888-5	Gestational age composite estimate	PQ
57065-5	Quickening date	TS
57066-3	Date fundal height reaches umbilicus	TS

6.3.4.27.14 <repeatNumber value=' '/> <interpretationCode code=' ' codeSystem=' '/> <targetSiteCode code=' ' codeSystem=' '/>

- 3580 The <repeatNumber> <interpretationCode>, and <targetSiteCode> elements should not be present in an EDD observation.

Add Section 6.3.4.28

6.3.4.28 Antepartum Visit Summary Battery 1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2

- 3585 This entry describes a single row in the Antepartum Visit Summary Flowsheet. The single observation date and provider is applied to all other observations.

6.3.4.28.1 Specification

```

<entry>
    <organizer classCode='BATTERY' moodCode='EVN'>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2' />
        <id root=' ' extension=' ' />
        <code code='57061-4' displayName='Antepartum flowsheet panel'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <statusCode code='completed' />
        <author>
            <time value=' ' />
            <assignedAuthor>
                <id root=' ' extension=' ' />
            </assignedAuthor>
        </author>
        <component>
            <observation classCode='OBS' moodCode='EVN'>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                :
                </observation>
            </component>
            <component>
                <observation classCode='OBS' moodCode='EVN'>
                    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                    :
                    </observation>
                </component>
                :
            </organizer>
</entry>

```

6.3.4.28.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2'>

The <templateId> element specifies that this organizer entry conforms to the APS Profile Antepartum Visit Summary Flowsheet battery. The root attribute SHALL contain the value "1.3.6.1.4.1.19376.1.5.3.1.1.11.2.3.2"

6.3.4.28.3 <organizer classCode='BATTERY' moodCode='EVN'>

Each row in the visit Summary flowsheet of the Antepartum Summary SHALL be represented by an organizer with the classCode of 'BATTERY' and the moodCode of 'EVN'

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

3625 Each battery SHALL have a globally unique identifier.

6.3.4.28.5 <code code='(57061-4)' codeSystem='2.16.840.1.113883.6.1'>

3630 The <code> element specifies the LOINC code that represents the content of the battery. The codeSystem attribute SHALL contain the value '2.16.840.1.113883.6.1'. The code attribute SHALL contain the value='(57061-4)'. It is good practice to include displayName and codeSystemName for clarity and debugging. The corresponding values are 'ACOG VISIT SUMMARY BATTERY--PT--' and 'LOINC' respectively.

6.3.4.28.6 <author><time/><assignedAuthor><id/></assignedAuthor></author>

3635 The <author> relation element points at the author that records the visit battery. This assignedAuthor may be different than the author of the document. The time element is used to record when the assigned author recorded the battery.

6.3.4.28.7 <statusCode code='completed'>

The status code for all batteries SHALL be 'completed'

6.3.4.28.8 <component>

3640 The battery is made of several component simple observations. The following table lists the allowable LOINC codes, displayNames, and observation types, and unit of measures for these observations.

LOINC Code	displayName	xsi:type	units	value set
11884-4	Gestational age Clinical.estimate	PQ	week	
57067-1 or 11727-5 (by US)	Fetal Body weight Estimated by palpation or Fetal weight estimated by US	PQ	g, kg, lb_av, or oz_av	
11881-0	Uterus Fundal height Tape measure	PQ	cm	

LOINC Code	displayName	xsi:type	units	value set
11876-0 (by PE) or 11877-8 (by US)	Fetal presentation by palpitation or Fetal presentation US	CD		SNOMED CT Vertex (70028003) Breech (6096002) Transverse (73161006) Oblique (63750008) Compound (124736009) Brow (8014007) Face (21882006)
11948-7 or 57068-9	Fetal Heart rate US or Fetal Heart rate Auscultation	PQ	/min	
57088-7	Fetal Movement - Reported	CO		SNOMED CT fetal movement activity (finding) CID 364755008 baby kicks a lot (finding) CID 276368003 baby not moving (finding) CID 276370007 reduced fetal movement (finding) CID 276369006 fetal movements present (finding) CID 289431008 fetal movements felt (finding) CID 268470003 fetal movements seen (finding) CID 169731002
57069-7	Preterm labor symptoms	BL		
11709-7 or 11785-3	DILATION-LEN-PT- CERVICAL CANAL.external os -QN-PALPATION or DILATION-LEN-PT- CERVICAL CANAL.external os-QN-US	PQ	cm	
11867-9	Effacement Cervix by palpitation	PQ	percent	
11961-0	Cervix [Length] US	PQ	cm	
8480-6	Systolic blood pressure	PQ	mmHg	
8462-4	Diastolic blood pressure	PQ	mmHg	
3141-9	Body weight Measured	PQ	g, kg, lb_av, or oz_av	
1753-3	Albumin [Presence] in Urine	CO		SNOMED CT Negative (finding) CID 167273002 Trace (finding) CID 167274008 1+ (finding) CID 167275009

LOINC Code	displayName	xsi:type	units	value set
				2+ (finding) CID 167276005 3+ (finding) CID 167277001 4+ (finding) CID 167278006
2349-9 or 25428-4(test strip)	Glucose [Presence] in Urine or Glucose [Presence] in Urine by Test strip	CO		SNOMED CT Negative (finding) CID 167261002 Trace (finding) CID 167262009 1+ (finding) CID 167264005 2+ (finding) CID 167265006 3+ (finding) CID 167266007 4+ (finding) CID 167267003
44966-0	Edema	CO		SNOMED CT Trace 44996-0 1+ pitting edema 420829009 2+ pitting edema 421605005 3+ pitting edema 421346005 4+ pitting edema 421129002
38208-5	Pain severity - Reported	CO		0 (no pain) : 10 (worst possible pain) Note: This observation should correspond to the functional status pain score observation
57070-5	Date next clinic visit	PQ	day,week,mo	
48767-8	Annotation comment	ED		

Add Section 6.3.4.29

3645 **6.3.4.29 Advance Directive Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.7**

An advance directive observation is a simple observation that uses a specific vocabulary, and inherits constraints from CCD.

6.3.4.29.1 Standards

CCD [ASTM/HL7 Continuity of Care Document](#)

6.3.4.29.2 Specification

```

3650 <observation typeCode='OBS' moodCode='EVN'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
      <templateId root='2.16.840.1.113883.10.20.1.17' />
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.7' />
      <id root=' ' extension=' '/>
      <code code=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />
      <text><reference value='#xxx' /></text>
      <statusCode code='completed' />
      <effectiveTime value=' '/>
      <value xsi:type='BL' value='true|false' />
      <reference typeCode='REFR'>
          <templateId root='2.16.840.1.113883.10.20.1.36' />
          <externalDocument classCode='DOC' moodCode='EVN'>
              <id root=' ' extension=' '/>
              <text><reference value=' ' /></text>
          </externalDocument>
      </reference>
</observation>
```

3670 An advanced directive <observation> shall be represented as shown above. They shall not contain any <repeatNumber>, <interpretationCode>, <methodCode> or <targetSiteCode> elements.

6.3.4.29.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' /> <templateId root='2.16.840.1.113883.10.20.1.17' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.7' />

3675 The <templateId> elements shown above shall be present, and indicated that this is an Advance Directive entry.

6.3.4.29.4 <code code=' ' codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />

3680 The <code> element records the type of advance directive. It should use one of the following SNOMED codes in the table below.

Code	Description	Data Type
304251008	Resuscitation	BL
52765003	Intubation	
225204009	IV Fluid and Support	
89666000	CPR	
281789004	Antibiotics	
78823007	Life Support	
61420007	Tube Feedings	

116859006	Transfusion of blood product	
71388002	Other Directive	<value> not permitted

6.3.4.29.5 <value xsi:type='BL' value='true|false'>

The advance directive observation may include a <value> element using the Boolean (xsi:type='BL') data type to indicate simply whether the procedure described is permitted. 3685 Absence of the <value> element indicates that an advance directive of the specified type has been recorded, and must be examined to determine what type of treatment should be performed. The value element is not permitted when the <code> element describes an Other directive.

6.3.4.29.6 <reference typeCode='REFR'>

3690 `<templateId root='2.16.840.1.113883.10.20.1.36'>
<externalDocument classCode='DOC' moodCode='EVN'>
<id root=' ' extension=' '/>
<text><reference value=' '/></text>`

The advanced directive observation may contain a single reference to an external document. That 3695 reference shall be recorded as shown above. The <id> element shall contain the appropriate root and extension attributes to identify the document. The <text> element may be present to provide a URL link to the document in the value attribute of the <reference> element. If the <reference> element is present, the Advance Directive in the narrative shall contain a <linkHTML> element to the same URL found in the value attribute.

3700

Add Section 6.3.4.30

6.3.4.30 Blood Type Observation 1.3.6.1.4.1.19376.1.5.3.1.4.13.6

The blood type observation is a Simple Observation of the patient's blood type. It conforms to the CCD Result observation template.

3705 **6.3.4.30.1 Standards**

CCD [ASTM/HL7 Continuity of Care Document](#)

6.3.4.30.2 Specification

```

<observation typeCode='OBS' moodCode='EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.6' />
  <templateId root='2.16.840.1.113883.10.20.1.31' />
  <id root=' ' extension=' '/>
  <code code='882-1' displayName='ABO+RH GROUP'
    codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
  <text><reference value='#xxx' /></text>
  <statusCode code='completed' />
  <effectiveTime value=' ' />
  <repeatNumber value=' ' />
  <value xsi:type='CE' code=' ' displayName=' ' codeSystem=' ' codeSystemName=' ' />
  <interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />
  <methodCode code=' ' codeSystem=' ' codeSystemName=' ' />
  <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />
<observation>
```

- 3725 **6.3.4.30.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />**
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13.6' />
<templateId root='2.16.840.1.113883.10.20.1.31' />

These <templateId> elements identify this as a blood type observation. They shall be present in the <observation> element as shown above.

- 3730 **6.3.4.30.4 <code code='882-1' displayName='ABO+RH GROUP'**
codeSystem='2.16.840.1.113883.6.1'
codeSystemName='LOINC' />

The <code> element shall be present to represent this as a finding of the patient's composite blood type. It shall use the code and codeSystem attributes shown above.

- 3735 **6.3.4.30.5 <repeatNumber value=' ' />**

The <repeatNumber> element should not be present in a blood type observation.

6.3.4.30.6 <value xsi:type='CE' code=' ' displayName=' '
codeSystem=' ' codeSystemName=' ' />

- 3740 The <value> element shall be present and shall use the CE data type. The code attribute should be valued using a vocabulary that supports encoding of blood types. The table below shows some coding systems that may be used to encode blood type.

Coding System	OID
ISBT 128	2.16.840.1.113883.6.18
SNOMED CT	2.16.840.1.113883.6.96

- 3745 **6.3.4.30.7 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />**
<methodCode code=' ' codeSystem=' ' />

~~codeSystemName=' '/>~~ ~~<targetSiteCode code=' '~~
~~codeSystem=' ' codeSystemName=' '/>~~

The <interpretationCode>, <methodCode>, and <targetSiteCode> should not be present in a blood type observation.

3750

Add Section 6.3.4.31

6.3.4.31 Encounters 1.3.6.1.4.1.19376.1.5.3.1.4.14

An Encounter is an interaction between a patient and care provider(s) for the purpose of providing healthcare-related service(s). Healthcare services include health assessment.

3755

Examples: outpatient visit to multiple departments, home health support (including physical therapy), inpatient hospital stay, emergency room visit, field visit (e.g., traffic accident), office visit, occupational therapy, or telephone call.

6.3.4.31.1 Standards

CCD [ASTM/HL7 Continuity of Care Document](#)

6.3.4.31.2 Specification

3760

```

<encounter classCode='ENC' moodCode='PRMS|ARQ|EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.14' />
  <templateId root='2.16.840.1.113883.10.20.1.21' />
  <templateId root='2.16.840.1.113883.10.20.1.25' />
  <id root='' extension=''/>
  <code code='' codeSystem='2.16.840.1.113883.5.4' codeSystemName='ActEncounterCode' />
  <text><reference value="#xxx"/></text>
  <effectiveTime>
    <low value=''/>
    <high value=''/>
  </effectiveTime>
  <priorityCode code=''/>
  <performer typeCode='PRF'>
    <time><low value=''/><high value=''/></time>
    <assignedEntity>...</assignedEntity>
  </performer>
  <author />
  <informant />
  <participant typeCode='LOC'>
    <participantRole classCode='SDLOC'>
      <id/>
      <code/>
      <addr>...</addr>
      <telecom value='' use=''/>
      <playingEntity classCode='PLC' determinerCode='INST'>
        <name></name>
      </playingEntity>
    </participantRole>
  </participant>
</encounter>

```

6.3.4.31.2.1 <encounter classCode='ENC' moodCode='APT|ARQ|EVN'>

This element is an encounter. The classCode shall be 'ENC'. The moodCode may be PRMS to indicate a scheduled appointment, ARQ to describe a request for an appointment that has been made but not yet scheduled by a provider, or EVN, to describe an encounter that has already occurred.

6.3.4.31.2.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.14'>

The templateId indicates that this <encounter> entry conforms to the constraints of this content module. NOTE: When the encounter is in event mood (moodCode='EVN'), this entry conforms to the CCD template 2.16.840.1.113883.10.20.1.21, and when in other moods, this entry conforms to the CCD template 2.16.840.1.113883.10.20.1.25.

6.3.4.31.2.3 <id root=" extension="/">

This required element shall contain an identifier for the encounter. More than one encounter identifier may be present.

6.3.4.31.2.4 <code code="" codeSystem='2.16.840.1.113883.5.4' codeSystemName='ActEncounterCode' />

This required element should contain a code from the HL7 ActEncounterCode vocabulary describing the type of encounter (e.g., inpatient, ambulatory, emergency, et cetera). Developers should take care to check that rational combinations of encounter.code and encounter.moodCode are used, but this Technical Framework does not restrict any combination.

3810 **6.3.4.31.2.5 <text><reference value="#xxx"/></text>**

The <text> element shall contain a reference to the narrative text describing the encounter.

6.3.4.31.2.6 <effectiveTime><low value="/"><high value="/"></effectiveTime>

This element records the time over which the encounter occurred (in EVN mood), or the desired time of the encounter in ARQ or APT mood. In EVN or APT mood, the effectiveTime element should be present. In ARQ mood, the effectiveTime element may be present, and if not, the priorityCode may be present to indicate that a callback is required to schedule the appointment.

6.3.4.31.2.7 <priorityCode code='CS' />

This element may be present in ARQ mood to indicate a callback is requested to schedule the appointment.

3820 **6.3.4.31.2.8 <performer>**

For encounters in EVN mood, at least one performer should be present that identifies the provider of the service given during the encounter. More than one performer may be present. The <time> element should be used to indicate the duration of the participation of the performer when it is substantially different from that of the effectiveTime of the encounter. In ARQ mood, the performer may be present to indicate a preference for a specific provider. In APT mood, the performer may be present to indicate which provider is scheduled to perform the service.

**6.3.4.31.2.9 <participant typeCode='LOC'>
<participantRole classCode='SDLOC'>**

A <participant> element with typeCode='LOC' may be present to provide information about the location where the encounter is to be or was performed. This element shall have a <participantRole> element with classCode='SDLOC' that describes the service delivery location.

6.3.4.31.2.10 <id/>

The <id> element may be present to identify the service delivery location.

6.3.4.31.2.11 <code/>

3835 The <code> element may be present to classify the service delivery location.

6.3.4.31.2.12 <addr>...</addr>

The <addr> element should be present, and gives the address of the location.

6.3.4.31.2.13 <telecom value="" use="" />

The <telecom> element should be present, and gives the telephone number of the location.

3840 **6.3.4.31.2.14 <playingEntity classCode='PLC'>**
 <name>...</name>
 </playingEntity>

The <playingEntity> shall be present, and gives the name of the location in the required <name> element.

3845

<i>Add Section 6.3.4.32</i>

6.3.4.32 Update Entry 1.3.6.1.4.1.19376.1.5.3.1.4.16

The update entry shall contain references to the entries or sections which are being replaced or updated. This reference shall not be present when the update entry is adding a new entries or sections.

3850 Entries and sections can be added, updated, or removed from a PHR. An update entry indicates the entry in the original PHR Extract that should be replaced or updated with new information contained within the entry. Only one organizer of this type is allowed in a section, and if present, it must be the first entry in the section.

3855 **6.3.4.32.1 Specification**

<pre> <entry> <organizer classCode='BATTERY' moodCode='EVN'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.16' /> <reference typeCode='RPLC'> <externalAct classCode='ACT' moodCode='EVN'> <id root='' extension=''/> </externalAct> </reference> </organizer> </entry> </pre>
--

6.3.4.32.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.16'>

3870 This templateId indicates that the organizer is used to update a PHR Extract.

6.3.4.32.3 <reference typeCode='RPLC'>

A reference element shall be present with typeCode RPLC. The reference element lists the acts that are affected by the update. It indicates that any referenced act is being replaced with new information. This element must be present, and may be repeated to replace more than one act at a time.

6.3.4.32.4 <externalAct classCode='ACT' moodCode='EVN'>

This element must appear as shown above. It indicates that the reference is to an external act (a section or entry contained in the parent document).

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

- 3880 This element identifies the information being replaced or updated. The identifier is of the entry or section being replaced. If the identifier is to a section being replaced, only one reference element is permitted.

Add Section 6.3.4.33

- 3885 **6.3.4.33 Procedure Entry 1.3.6.1.4.1.19376.1.5.3.1.4.19**

The procedure entry is used to record procedures that have occurred, or which are planned for in the future.

6.3.4.33.1 Standards

CCD [ASTM/HL7 Continuity of Care Document](#)

6.3.4.33.2 Specification

```

3890 <procedure classCode='PROC' moodCode='EVN|INT'>
    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19' />
    <templateId root='2.16.840.1.113883.10.20.1.29' /><!-- see text of section 0 -->
    <templateId root='2.16.840.1.113883.10.20.1.25' /><!-- see text of section 0 -->
    <id root=' ' extension=' '/>
    <code code='' codeSystem='2.16.840.1.113883.5.4' codeSystemName='ActCode' />
    <text><reference value='#xxx' /></text>
    <statusCode code='completed|active|aborted|cancelled' />
    <effectiveTime>
        <low value=' '/>
        <high value=' '/>
    </effectiveTime>
    <priorityCode code='' />
    <approachSiteCode code='' displayName='' codeSystem='' codeSystemName='' />
    <targetSiteCode code='' displayName='' codeSystem='' codeSystemName='' />
    <author />
    <informant />
    <entryRelationship typeCode='COMP' inversionInd='true'>
        <act classCode='ACT' moodCode=''>
            <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4.1' />
            <id root=' ' extension=' '/>
        </act>
    </entryRelationship>
    <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>
</procedure>

```

3920

6.3.4.33.2.1 <procedure classCode='PROC' moodCode='EVN|INT'>

This element is a procedure. The classCode shall be 'PROC'. The moodCode may be INT to indicate a planned procedure or EVN, to describe a procedure that has already occurred.

6.3.4.33.2.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.19'>

- 3925 The templateId indicates that this <procedure> entry conforms to the constraints of this content module. NOTE: When the procedure is in event mood (moodCode='EVN'), this entry conforms to the CCD template 2.16.840.1.113883.10.20.1.29, and when in intent mood, this entry conforms to the CCD template 2.16.840.1.113883.10.20.1.25.

6.3.4.33.2.3 <id root=" extension="/>

- 3930 This required element shall contain an identifier for the procedure. More than one procedure identifier may be present.

6.3.4.33.2.4 <code code="" displayName="" codeSystem="" codeSystemName="" />

This element shall be present, and should contain a code describing the type of procedure.

6.3.4.33.2.5 <text><reference value="#xxx"/></text>

- 3935 The <text> element shall contain a reference to the narrative text describing the procedure.

6.3.4.33.2.6 <statusCode code='completed|active|aborted|cancelled' />

- 3940 The <statusCode> element shall be present when used to describe a procedure event. It shall have the value 'completed' for procedures that have been completed, and 'active' for procedures that are still in progress. Procedures that were stopped prior to completion shall use the value 'aborted', and procedures that were cancelled before being started shall use the value 'cancelled'.

6.3.4.33.2.7 <effectiveTime><low value="/"><high value="/"></effectiveTime>

This element should be present, and records the time at which the procedure occurred (in EVN mood), or the desired time of the procedure in INT mood.

6.3.4.33.2.8 <priorityCode code="/" />

- 3945 This element shall be present in INT mood when effectiveTime is not provided, it may be present in other moods. It indicates the priority of the procedure.

6.3.4.33.2.9 <approachSiteCode code="" displayName="" codeSystem="" codeSystemName="" />

This element may be present to indicate the procedure approach.

- 3950 **6.3.4.33.2.10 <targetSiteCode code="" displayName="" codeSystem="" codeSystemName="" />**

This element may be present to indicate the target site of the procedure.

6.3.4.33.2.11 <entryRelationship typeCode='COMP' inversionInd='true'>

3955 This element may be present to point the encounter in which the procedure was performed, and shall contain an internal reference to the encounter. See PCC TF-2: 6.3.4.10 Internal References for more details.

6.3.4.33.2.12 <entryRelationship typeCode='RSON'>

3960 A <procedure> act may indicate one or more reasons for the procedure. These reasons identify the concern that was the reason for the procedure via an Internal Reference (see PCC TF-2: 6.3.4.10 Internal References) to the concern. The extension and root of each observation present must match the identifier of a concern entry contained elsewhere within the CDA document.

Add Section 6.3.4.34

6.3.4.34 Transport 1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1

3965 Defined in IHE PCC TF-2:6.3.4.34

Add Section 6.3.4.35

6.3.4.35 Encounter Disposition 1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2

3970 This element records the intended or actual disposition for the patient (e.g., admit, discharge home after treatment, et cetera).

6.3.4.35.1 Specification

```

<act classCode='ACT' moodCode='INT|EVN'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2' />
  <id root='' extension=''/>
  <code code='' displayName='' codeSystem='' codeSystemName='' />
  <text><reference value='#xxx' /></text>
  <statusCode code='normal|completed' />
  <effectiveTime value=''/>
  <performer typeCode='PRF'>
    <assignedEntity>
      <id root='' extension=''/>
      <addr></addr>
      <telecom value='' use=''/>
      <assignedPerson>
        <name></name>
      </assignedPerson>
    </assignedEntity>
  </performer>
  <participant typeCode='RCV'>
    <time value=''/>
    <participantRole classCode='ROL'>
      <id root='' extension=''/>
      <addr></addr>
      <telecom value='' use=''/>
      <playingEntity>
        <name></name>
      </playingEntity>
    </participantRole>
  </participant>
  <entryRelationship typeCode='COMP'>
    <act classCode='ACT'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1' />
      :
    </act>
  </entryRelationship>
</act>

```

6.3.4.35.1.1 <act classCode='ACT' moodCode='INT|EVN'>

The disposition is recorded in an act element, to describe the disposition action taken during the encounter¹. In intent mood (moodCode='INT'), this records the expected disposition of the patient. In event mood (moodCode='EVN'), this records the actual disposition.

¹ The HL7 RIM allows this portion of the encounter to be recorded in the dischargeDispositionCode RIM Attribute of the Encounter class, but the Encounter class is constrained within CDA. To record the disposition act therefore requires the use of the Act class.

6.3.4.35.1.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.2' />

The templateId indicates that this <encounter> entry conforms to the constraints of this content module.

4015 **6.3.4.35.1.3 <id root=" extension=" />**

This required element shall contain an identifier.

6.3.4.35.1.4 <code code="" displayName="" codeSystem="" codeSystemName="" />

- 4020 This required element indicates the disposition of the patient. The code shall come from a coding system that is able to record common patient dispositions (e.g., Discharged, Transferred, Admitted). The "Administrative Procedure" concept (14734007) of SNOMED CT contains several code values that cover a wide variety of dispositions routinely recorded. Other vocabularies that are commonly in use to describe discharge disposition codes are DEEDS (see section 8.02), and in the US, the Uniform National Billing Code.

6.3.4.35.1.5 <text><reference value="#xxx"/></text>

- 4025 The <text> element shall contain a reference to the narrative text describing the disposition of the patient. <statusCode code='normal|completed' /> When the disposition act has occurred (moodCode='EVN'), the statusCode element shall be present, and shall contain the value 'completed'. When the disposition act is intended (moodCode='EVN') the statusCode element shall contain the value 'normal'.

4030 6.3.4.35.1.6 <effectiveTime><low value="/" /><high value="/" /><effectiveTime/>

When the disposition has occurred, this element shall be sent, and indicates the effective time for the disposition process. This element may be sent to record when the disposition act is intended to occur. The <low> element records the time at which the disposition process was started. The <high> value records the time at which the disposition process was completed.

4035 6.3.4.35.1.7 <performer typeCode='PRF' />

The <performer> element provides information about the person that performs the discharge, admission or transfer of the patient. When the disposition is in intent mood, this element describes any expectations with respect to the performer, and is optional. When the disposition is in event mood, this element is required.

4040 6.3.4.35.1.8 <assignedEntity>

The <assignedEntity> element identifies the performer of the disposition.

6.3.4.35.1.9 <id root="" extension="/" />

The <id> element shall be sent when the disposition has occurred, and identifies the performer of the act.

4045 6.3.4.35.1.10 <addr></addr>

The <addr> element may be sent to provide a contact postal address for the performer of the disposition.

6.3.4.35.1.11 <telecom value="" use="/" />

- 4050 The <telecom> element may be sent to provide a contact postal address for the performer of the disposition.

6.3.4.35.1.12 <assignedPerson><name/></assignedPerson>

The <assignedPerson> element shall be sent to identify the person who performed the disposition of the patient.

6.3.4.35.1.13 <participant typeCode='RCV'>

4055 <time value="/">
 <participantRole classCode='ROL'>
 <id root=" extension="/">
 <addr></addr>
 <telecom value=" use="/">
4060 <playingEntity><name/></playingEntity>

This element identifies the person or organization that is receiving the patient. =====

<entryRelationship typeCode='COMP'>

<act classCode='ACT'>

4065 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.10.4.1'> If the disposition of the patient requires transport to another location, this information shall be recorded in a subordinate act that conforms to the Transport template described above.

Add Section 6.3.4.36

6.3.4.36 Reserved for Coverage Activity

4070 Not yet defined in IHE PCC TF-2:6.3.4

Add Section 6.3.4.37

6.3.4.37 Reserved for Payer Entry

Not yet defined in IHE PCC TF-2:6.3.4

4075

Add Section 6.3.4.38

Section 6.3.4.38.4 updated by CP PCC 0209

6.3.4.38 Pain Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.1

4080 The pain score observation is a [Simple Observation](#) that records the patient's assessment of their pain on a scale from 0 to 10.

6.3.4.38.1 Parent Template

The parent of this template is [Simple Observation](#).

6.3.4.38.2 Specification

```

4085 <observation typeCode='OBS' moodCode='EVN'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
      <id root=' ' extension=' '/>
      <code code='38208-5|38221-8|38214-3' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'>
          <translation code='406127006' displayName='Pain intensity'
                      codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />
      </code>
      <text><reference value='#xxx' /></text>
      <statusCode code='completed' />
      <effectiveTime value=' ' />
      <repeatNumber value=' ' />
      <value xsi:type='CO|REAL' />
      <interpretationCode code= codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />
      <methodCode code=' ' codeSystem=' ' codeSystemName=' ' />
      <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />
</observation>

```

6.3.4.38.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13'>

The <templateId> identifies this as a Pain Score Observation, and shall be present as shown above.

4105 **6.3.4.38.4 <code code='38208 5' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC'>**
<translation code='406127006' displayName='Pain intensity'
codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT' />

4110 The <code> element indicates what kind of pain observation was made. It shall contain the code and codeSystem attribute values shown above. The <translation> element may be present, and provides a mapping to SNOMED CT of the observation. If present, shall have the code and codeSystem attribute values shown above. 38208-5 is used for example purposes. Any of the following codes can be used 38208-5|38221-8|38214-3

Code	Data Type	Description
38208-5	CO	A Pain Score made using the Numerical Rating Scale (NRS), where pain is assessed on a scale from 0 to 10. -->>The code system to use for this observation<<--

4115

6.3.4.38.5 <value xsi:type='CO' value=' ' />

4120 The <value> element records the assessed pain score. If using the NRS the pain is assessed using coded ordinal values that range from 0 to 10. The use of the coded ordinal type is required because while pain assessments are ordered values, and can be compared, the differences between two pain assessment values cannot be compared, and so these values are not really numbers.

Null flavors may be used when clinical values are not present or available. The applicable null flavors SHALL be from the following subset of the HL7 v3 value set:

- UNK – Unknown
 - NI – No Information
 - NA – Not applicable
 - OTH – Other
 - NASK – Not asked
 - ASKU – asked but unknown
- 4125 • MSK – Masked
- NAV - Temp unavailable

4130

6.3.4.38.6 <interpretationCode

```
code='301379001|40196000|76948002|67849003'
codeSystem='2.16.840.1.113883.6.96'
codeSystemName='SNOMED CT'>
```

4135

The <interpretationCode> element should be present to provide an interpretation of the pain scale assessment using SNOMED CT. When the <interpretationCode> element is present, the <translation> element described above shall be present. These interpretations are provided to assist decision support systems that are making secondary use of the assessment information, and are not intended to replace the score values.

4140

Pain Score Range	Code	Description
0	301379001	No Present Pain
1-3	40196000	Mild Pain
4-6	50415004	Moderate Pain
7-9	76948002	Severe Pain
10	67849003	Excruciating Pain

6.3.4.38.7 <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>

4145

The <methodCode> should not be present in a Pain Score Observation, as the method is implied by the <code> element.

6.3.4.38.8 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>

The <targetSiteCode> element should be present, and shall indicate the location of the pain being assessed.

Add Section 6.3.4.39

4150 **6.3.4.39 Braden Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.2**

Add Section 6.3.4.40

6.3.4.40 Braden Score Component 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.3

4155 *Add Section 6.3.4.41*

6.3.4.41 Geriatric Depression Score Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.4

Add Section 6.3.4.42

6.3.4.42 Geriatric Depression Score Component 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.5

4160

Add Section 6.3.4.43

6.3.4.43 Survey Panel 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7

A survey panel collects related survey observations.

6.3.4.43.1 Parent Template

4165 This template is compatible with the ASTM/HL7 Continuity of Care Document template:
2.16.840.1.113883.10.20.1.32

6.3.4.43.2 Uses

See Templates using [Survey Panel](#).

6.3.4.43.3 Specification

4170

```
<organizer classCode='CLUSTER' moodCode='EVN'>
  <templateId root='2.16.840.1.113883.10.20.1.32' />
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7' />
  <id root='' extension=''/>
  <code code=' ' displayName=' '
    codeSystem=' ' codeSystemName=' ' />
  <statusCode code='completed' />
  <effectiveTime value=''/>
  <!-- one or more survey observations -->
  <component typeCode='COMP'>
    <observation classCode='OBS' moodCode='EVN'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6' />
      :
    </observation>
  </component>
</organizer>
```

```
</observation>
</component>
</organizer>
```

6.3.4.43.3.1 <organizer classCode='CLUSTER' moodCode='EVN'>

The survey panel is a cluster of related survey observations.

6.3.4.43.3.2 <templateId root='2.16.840.1.113883.10.20.1.32'>

4175 **<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.7'>**

The survey panel shall have the <templateId> elements shown above to indicate that it inherits constraints from the ASTM/HL7 CCD Specification for results organizers, and the constraints of this specification.

6.3.4.43.3.3 <id root=' ' extension=' '>

4180 The organizer shall have an <id> element.

6.3.4.43.3.4 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '>

The <code> element shall be present, and identifies the survey panel.

4185 **6.3.4.43.3.5 <statusCode code='completed'>**

The observations have all been completed.

6.3.4.43.3.6 <effectiveTime value=' '>

The effective time element shall be present to indicate when the survey panel was taken.

6.3.4.43.3.7 <!-- one or more survey observations -->

4190 **<component typeCode='COMP'>**

The organizer shall have one or more <component> elements that are <observation> elements using the [Survey Observation](#) template.

4195 **Add Section 6.3.4.44**

6.3.4.44 Survey Observation 1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6

Survey observations are used to record responses to assessment instruments. They are simple observations conforming to the CCD Result template. The vocabulary and data type constraints on survey observations is specified elsewhere, either in the specializations of the survey observation template, or by the template that makes use of it.

6.3.4.44.1 Parent Template

The parent of this template is [Simple Observation](#). This template is compatible with the ASTM/HL7 Continuity of Care Document template: 2.16.840.1.113883.10.20.1.31

6.3.4.44.2 Uses

- 4205 See [Templates using Survey Observation](#).

6.3.4.44.3 Specification

```
<observation classCode='OBS' moodCode='EVN'>
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
<templateId root='2.16.840.1.113883.10.20.1.31' />
<templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6' />
<id root=' ' extension=' ' />
<code code=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
<text><reference value='#xxx' /></text>
<statusCode code='completed' />
<effectiveTime value=' ' />
<repeatNumber value=' ' />
<value xsi:type='CO|CD|INT|PQ' />
<interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />
<methodCode code=' ' codeSystem=' ' codeSystemName=' ' />
<targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />
</observation>
```

6.3.4.44.3.1 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' /> <templateId root='2.16.840.1.113883.10.20.1.31' /> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.12.3.6' />

- 4210 A survey observation shall have the <templateId> elements shown above to indicate that it inherits constraints from the ASTM/HL7 CCD Specification for results, and the constraints of this specification.

6.3.4.44.3.2 <code code=' ' codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />

- 4215 A survey observation entry shall contain a code identifying the observation made.

6.3.4.44.3.3 <value xsi:type='CO|CD|INT|PQ' ... />

The <value> element shall be present, and shall be of the appropriate data type specified for the observation.

6.3.4.44.3.4 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />

- 4220 An interpretation code may be present to provide an interpretation of the observation.

**6.3.4.44.3.5 <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>
 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>**

The <methodCode> and <targetSiteCode> element shall not be present, as these are not relevant to survey responses.

4225

Add Section 6.3.4.45

6.3.4.45 Acuity 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1

An acuity entry indicates the triage acuity entry and the triage time of the patient.

6.3.4.45.1 Specification

4230

```
<entry>
    <!-- Acuity Event -->
    <observation classCode='OBS' moodCode='EVN'>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1' />
        <id root='' extension=''/>
        <code code='' displayName=''
            <code code='273887006' displayName='Triage index'
                codeSystem='2.16.840.1.113883.6.96'
                codeSystemName='SNOMED CT' /> <!-- Triage index (assessment scale) FullySpecifiedName --
        ->
            <originalText><reference value='#(ID of text coded)' /></originalText>
        </code>
        <text><reference value='#text' /></text>
        <!-- effectiveTime
        <effectiveTime>
            <low value=''/> <!-- start of triage, may be sent -->
            <high value=''/><!-- end of triage should be sent -->
        </effectiveTime>
    </observation>
</entry>
```

4240

4245

4250

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

This element indicates that the entry is an observation regarding the event of triage assessment. This entry records the observation and the time of the observation.

6.3.4.45.1.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1' />

4255

The <templateId> element identifies this <act> as about Acuity Assessment of the patient. The templateId must have root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.1'.

6.3.4.45.1.3 <id root=" extension="/>

The entry must have an identifier.

**6.3.4.45.1.4 <code code="" displayName=""
codeSystem='2.16.840.1.113883.6.96' codeSystemName='SNOMED CT'>**

- 4260 The code describes the triage acuity scale. IHE recommends the use the Emergency Severity Index (ESI). However, the vocabulary used within an affinity domain may be determined by a policy agreement within the domain.

6.3.4.45.1.5 <originalText><reference value="#xxx"/><originalText>

This is a reference to the narrative text within the section that describes the acuity description.

- 4265 **6.3.4.45.1.6 <text><reference value="#text"/></text>**

This is a reference to the narrative text corresponding to the Observation act.

6.3.4.45.1.7 <effectiveTime>

- 4270 The effectiveTime element shall be sent. It records the interval of time over which triage occurs. The use case for this information requires that the ending time of triage be recorded. However, the <low value=""> element may be sent by systems that capture the beginning and end of the triage process.

6.3.4.45.1.8 <high value="/">

- 4275 This element records the time of completion of triage, and is required. If unknown, it must be recorded using a flavor of null. This element may be sent using the TS data type, as shown above. If there is uncertainty about the time of completion of triage, the sender may record the time using the IVL_TS data type, as shown below.

4280

```
<high xsi:type='IVL_TS'>
  <low value=''/>
  <high value=''/>
</high>
```

Add Section 6.3.4.46

- 4285 **6.3.4.46 Intravenous Fluids 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.2**

This content module describes the general structure for intravenous fluids. All intravenous fluid administration acts should be derived from this content module.

6.3.4.46.1 Specification

```

4290 <substanceAdministration classCode='SBADM' moodCode='INT|EVN'>
4291   <templateId root='2.16.840.1.113883.10.20.1.24' />
4292   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7' />
4293   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.1' />
4294   <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.2' />
4295   <id root='' extension=''/>
4296   <code code='' codeSystem='' displayName='' codeSystemName='' />
4297   <text><reference value='#med-1' /></text>
4298   <statusCode code='completed|active' />
4299   <effectiveTime xsi:type='IVL_TS'>
4300     <low value=''/>
4301     <high value=''/>
4302   </effectiveTime>
4303   <effectiveTime operator='A' xsi:type='TS|PIVL_TS|EIVL_TS|PIVL_PPD_TS|SXPR_TS'>
4304     :
4305   </effectiveTime>
4306   <routeCode code='' codeSystem='' displayName='' codeSystemName='' />
4307   <doseQuantity value='' unit='' />
4308   <approachSiteCode code='' codeSystem='' displayName='' codeSystemName='' />
4309   <rateQuantity value='' unit='' />
4310   <consumable>
4311     :
4312     .
4313   </consumable>
4314   <!-- 0..* entries describing the components -->
4315   <entryRelationship typeCode='COMP' >
4316     <sequenceNumber value=''/>
4317   </entryRelationship>
4318   <!-- An optional entry relationship that indicates the reason for use -->
4319   <entryRelationship typeCode='RSON'>
4320     <act classCode='ACT' moodCode='EVN'>
4321       <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.4.1' />
4322       <id root='' extension=''/>
4323     </act>
4324   </entryRelationship>
4325   <!-- An optional entry relationship that provides prescription activity -->
4326   <entryRelationship typeCode='REFR'>
4327     <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.3' />
4328     :
4329     .
4330   </entryRelationship>
4331   <precondition>
4332     <criterion>
4333       <text><reference value=''/></text>
4334     </criterion>
4335   </precondition>
4336 </substanceAdministration>

```

This content module is derived from the Medication content module to specifically and more easily describe the necessary details of intravenous fluid administration. For the purpose of EDER and other profiles employing this content module, the table below identifies and describes the fields and 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.

4345

6.3.4.46.1.1 Medication Fields

Field	Opt.	CDA Tag	Description
Start and Stop Date	R2	<effectiveTime>	The date and time when the fluid regimen began and is expected to finish. The first component of the <effectiveTime> encodes the lower and upper bounds over which the <substanceAdministration> occurs, and the start time is determined from the lower bound. If the fluid has been known to be stopped, the high value must be present, but expressed as a flavor of null (e.g., Unknown).
Dose	R2	<doseQuantity>	The amount of fluid given. This should be in some known and measurable fluid unit, such as milliliters, or may be measured in "administration" units (such "units" of blood or "packs" of platelets).
Site	O	<approachSiteCode>	The site where the fluid is administered (i.e., "Left Antecubital", or "Central Line").
Rate	R2	<rateQuantity>	The rate is a measurement of how fast the fluid is given to the patient over time (e.g., .5 liter / 1 hr).
Product	R	<consumable> <name> </consumable>	The name of the substance or product. This should be sufficient for a provider to identify the type of fluid. It may be a trade name (Plasmalyte®) or a generic name. This information is required in all fluid entries. The name should not include packaging, strength or dosing information.
Code	R2	<consumable> <code/> </consumable>	A code describing the product from a controlled vocabulary, such as RxNorm, First DataBank, et cetera.

6.3.4.46.1.2 <substanceAdministration classCode='SBADM' moodCode='INT|EVN'>

4350

The general model is to record each fluid administered in a <substanceAdministration> intent (moodCode='INT'). Fluids that have been started but not completely administered should be recorded in a <substanceAdministration> intent (moodCode='INT'). Fluids that have been completed should be recorded as an event (moodCode='EVN').

6.3.4.46.1.3 <templateId root='2.16.840.1.113883.10.20.1.24'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.7.1' />

4355

All intravenous fluid entries use the <templateId> elements specified above to indicate that they are IV fluid administration acts. This element is required.

6.3.4.46.1.4 <id root=" extension="/">

4360

The <substanceAdministration> element must be uniquely identified. If there is no explicit identifier for this observation in the source EMR system, a GUID may be used for the root attribute, and the extension may be omitted. Although HL7 allows for multiple identifiers, this Technical Framework profile requires that one and only one be used.

6.3.4.46.1.5 <code code="" displayName="" codeSystem="" codeSystemName="">

- 4365 The <code> element is required, and is used to supply a code that describes the act of fluid administration, not the fluid being administered. This may be a procedure code, such as those found in CPT-4 (and often used for billing), or may describe the method of administration, such as by intravenous injection.

6.3.4.46.1.6 <text><reference value="/" /></text>

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 fluid administration.

- 4370 **6.3.4.46.1.7 <statusCode code='completed|active' />**

The status of all <substanceAdministration> elements must be "completed" or "active". If "completed", then the administration has occurred, or the request or order has been placed. If "active", then at the time recorded, the fluid was still being administered.

6.3.4.46.8 <effectiveTime xsi:type='IVL_TS'>

- 4375 The first <effectiveTime> element encodes the start and stop time of the administration. This is an interval of time (xsi:type='IVL_TS'), and must be specified as shown. This is an additional constraint placed upon CDA Release 2.0 by this Technical Framework profile, and simplifies the exchange of start/stop and frequency information between EMR systems.

6.3.4.46.1.9 <low value="/" /><high value="/" />

- 4380 The <low> and <high> values of the first <effectiveTime> element represent the start and stop times for the fluid administration. The <low> value represents the start time, and the <high> value represents the stop time. If either the <low> or the <high> value is unknown, this shall be recorded by setting the nullFlavor attribute to UNK. The <high> value records the end of the fluid administration according to the information provided in the initial fluid order or RN documentation. For example, if the fluid order is for one liter, and the fluid is to be delivered at 250 mL/hr, then the high value should contain a datetime that is 4 hours later than the <low> value. The rationale is that a provider, seeing a discontinued fluid could normally assume that the fluid has been stopped, even if the intent of the treatment plan is to continue the fluid continuously.
- 4385

- 4390 **6.3.4.46.1.10 <approachSiteCode code="" codeSystem="" originalText><reference value="/" /></originalText></approachSiteCode>**

- 4395 The <approachSiteCode> element contains a URI in the value attribute of the <reference> that points to the text in the narrative identifying the site. It may be coded to a controlled vocabulary that lists such sites (e.g., SNOMED-CT).

6.3.4.46.1.11 <doseQuantity><low value="" unit="" /><high value="" unit="" /></doseQuantity>

4400 The dose is specified if the <doseQuantity> element. If a dose range is given (e.g., 125-250 mL/hr [i.e., to replace fluid losses]), then the <low> and <high> bounds are specified in their respective elements, otherwise both <low> and <high> have the same value. The unit attribute should be derived from the HL7 UnitsOfMeasureCaseSensitive vocabulary .

6.3.4.46.1.12 <low|high value="" /> <translation> <originalText><reference value="" /></originalText> </translation></low|high >

4405 Any <low> and <high> elements used for <doseQuantity> or <rateQuantity> should contain a <translation> element that provides a <reference> to the <originalText> found in the narrative body of the document .

6.3.4.46.1.13 <rateQuantity><low value="" unit="" /><high value="" unit="" /></rateQuantity>

4410 The rate is specified in the <rateQuantity> element. The rate is given in units that have measure over time. In this case, the units should be specified as a string made up of a unit of measure (see doseQuantity above), followed by a slash (/), followed by a time unit (s, min, h or d) (i.e., mL/hr).

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

4415 **6.3.4.46.1.14 <consumable>**

The <consumable> element shall be present, and shall contain a <manufacturedProduct> entry conforming to the Product Entry template (see PCC TF-2: 6.3.4.19).

<i>Add Section 6.3.4.47</i>

4420 **6.3.4.47 Nursing Assessments Battery 1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4**

This entry describes a single row in the Nursing Assessment flowsheet. The single observation date/time and provider is applied to all other observations.

4425

6.3.4.47.1 Specification

```

4430 <entry>
    <organizer classCode='BATTERY' moodCode='EVN'>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4' />
        <id root=' ' extension=' ' />
        <code code='XX-ASSESS' displayName='Nursing Assessments Battery'
              codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
        <statusCode code='completed' />
        <author>
            <time value=' ' />
            <assignedAuthor>
                <id root=' ' extension=' ' />
            </assignedAuthor>
        </author>
        <component>
            <observation classCode='OBS' moodCode='EVN'>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                :
                </observation>
            </component>
            <component>
                <observation classCode='OBS' moodCode='EVN'>
                    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                    :
                    </observation>
                </component>
                :
            </organizer>
        </entry>

```

6.3.4.47.1.1 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4' />

The <templateId> element specifies that this organizer entry conforms to the Nursing Interventions battery. The root attribute SHALL contain the value "1.3.6.1.4.1.19376.1.5.3.1.1.13.3.4"

6.3.4.47.1.2 <organizer classCode='BATTERY' moodCode='EVN' />

Each row in the Nursing Interventions battery SHALL be represented by an organizer with the classCode of 'BATTERY' and the moodCode of 'EVN'

4465 6.3.4.47.1.3 <id root=' ' extension=' ' />

Each battery SHALL have a globally unique identifier.

6.3.4.47.1.4 <code code='X-ASSESS' codeSystem='2.16.840.1.113883.6.1' />

The <code> element specifies the LOINC code that represents the content of the battery. The codeSystem attribute SHALL contain the value '2.16.840.1.113883.6.1'. The code attribute SHALL contain the value='X-ASSESS'. It is good practice to include displayName and codeSystemName for clarity and debugging. The corresponding values are 'Nursing Assessments battery' and 'LOINC' respectively.

6.3.4.47.1.5 <author><time><assignedAuthor><id></assignedAuthor></author>

- 4475 The <author> relation element points at the author that records the visit battery. This assignedAuthor may be different than the author of the document. The time element is used to record when the assigned author recorded the battery.

6.3.4.47.1.6 <statusCode code='completed'>

The status code for all batteries SHALL be 'completed'

6.3.4.47.1.7 <component>

- 4480 The battery is made of several component Simple Observations (see PCC TF-2: 6.3.4.20). The following table lists the allowable LOINC codes, displayNames, and observation types, and unit of measures for these observations.

LOINC Code	displayName	xsi:type	value set
9269-2	GLASGOW COMA CORE.TOTAL	CO	3..15
9268-4	GLASGOW COMA SCORE.MOTOR	CO	1..6
11454-6	LEVEL OF RESPONSIVENESS	CO	ALERT VERBAL RESPONSE PAINFUL RESPONSE UNRESPONSIVE
38208-5	PAIN SEVERITY	CO	0-10
48767-8	(COMMENT FIELD)	ED	

- 4485 Add Section 6.3.4.48

6.3.4.48 Antenatal Testing and Surveillance Battery**1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10**

This entry describes a single row in the Antenatal Testing and Surveillance Section. The single observation date/time and provider is applied to all other observations.

4490 **6.3.4.48.1 Specification**

```

4495 <entry>
    <organizer classCode='BATTERY' moodCode='EVN'>
        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10' />
        <id root=' ' extension=' ' />
        <code code='XX-ANTENATALTESTINGBATTERY' displayName='ANTENATAL TESTING AND SURVEILLANCE
BATTERY'
            codeSystem='2.16.840.1.113883.6.1' codeSystemName='LOINC' />
4500     <statusCode code='completed' />
        <author>
            <time value=' ' />
            <assignedAuthor>
                <id root=' ' extension=' ' />
            </assignedAuthor>
        </author>
4505     <component>
            <observation classCode='OBS' moodCode='EVN'>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                :
                </observation>
            </component>
            <component>
                <observation classCode='OBS' moodCode='EVN'>
                    <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                    :
                    </observation>
                </component>
                :
            </component>
        </organizer>
4520 </entry>

```

6.3.4.48.1.1 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10' />

The <templateId> element specifies that this organizer entry conforms to the Antenatal Testing and Surveillance Battery. The root attribute SHALL contain the value "1.3.6.1.4.1.19376.1.5.3.1.1.21.3.10"

4525 **6.3.4.48.1.2 <organizer classCode='BATTERY' moodCode='EVN'>**

Each row in the Antenatal Testing and Surveillance Battery SHALL be represented by an organizer with the classCode of 'BATTERY' and the moodCode of 'EVN'

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

Each battery SHALL have a globally unique identifier.

4530 **6.3.4.48.1.4 <code code='XX- XX-ANTENATALTESTINGBATTERY'
codeSystem='2.16.840.1.113883.6.1' />**

The <code> element specifies the LOINC code that represents the content of the battery. The codeSystem attribute SHALL contain the value '2.16.840.1.113883.6.1'. The code attribute SHALL contain the value='XX-ANTENATALTESTINGBATTERY'. It is good practice to include displayName and codeSystemName for clarity and debugging. The corresponding values are 'ANTENATAL TESTING AND SURVEILLANCE BATTERY' and 'LOINC' respectively.

6.3.4.48.1.5 <author><time><assignedAuthor><id></assignedAuthor></author>

4540 The <author> relation element points at the author that records the visit battery. This assignedAuthor may be different than the author of the document. The time element is used to record when the assigned author recorded the battery.

6.3.4.48.1.6 <statusCode code='completed'>

The status code for all batteries SHALL be 'completed'

6.3.4.48.1.7 <component>

4545 The battery is made of several component Simple Observations (see PCC TF-2: 6.3.4.20). The following table lists the allowable LOINC codes, displayNames, and observation types, and unit of measures for these observations.

LOINC Code	displayName	xsi:type
11630-1	Biophysical profile.amniotic fluid volume	ED
11631-9	Biophysical profile.body movement	ED
11632-7	Biophysical profile.breathing movement	ED
11633-5	Biophysical profile.heart rate reactivity	ED
11635-0	Biophysical profile.tone	ED
11634-3	Biophysical profile.sum	ED
35096-7	Ultrasound morphologic	ED
49086-2	Nuchal translucency screening	ED
51659-1	Hbs1 Antigen	ED

Add Section 6.3.4.49

4550 **6.3.4.49 Immunization Recommendation**

Defined in IHE PCC TF-2:6.3.4

*Add Section 6.3.4.50***6.3.4.50 Alert Entry**

4555 Defined in IHE PCC TF-2:6.3.4

Add Section 6.3.4.51

6.3.4.51 Antigen Dose

Defined in IHE PCC TF-2:6.3.4

4560

Add Section 6.3.4.52 (Occupation Observation – removed 2011-09 at the request of QRPH)

6.3.4.52 Intentionally blank

Add Section 6.3.4.53 (Industry Observation removed 2011-09 at the request of QRPH)

4565

6.3.4.53 Intentionally blank

Add Section 6.3.4.54

6.3.4.54 Observation Request 1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1

4570

The observation request entry is used to record goals, plans or intention for an observation to be performed (e.g., assessment, laboratory test, imaging study, et cetera).

6.3.4.54.1 Uses

See Templates using Observation Request.

6.3.4.54.2 Specification

4575

```
<observation classCode='OBS' moodCode='INT|PRP|GOL'>
  <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1' />
  <templateId root='2.16.840.1.113883.10.20.1.25' />
  <id root='' extension='' />
  <code code='' displayName='' codeSystem='' codeSystemName='' />
  <!-- for CDA -->
  <text><reference value="#xxx"/></text>
  <!-- For HL7 Version 3 Messages
  <text>text</text>
  -->
  <statusCode code='active' />
  <effectiveTime value='' />
  <repeatNumber value='' />
  <value xsi:type='' .../>
  <interpretationCode code='' codeSystem='' codeSystemName='' />
  <methodCode code='' codeSystem='' codeSystemName='' />
  <targetSiteCode code='' codeSystem='' codeSystemName='' />
  <author typeCode='AUT'>
    <assignedAuthor typeCode='ASSIGNED'><id ... /></assignedAuthor> <!-- for CDA --
  >
```

4600

```

<!-- For HL7 Version 3 Messages
<assignedEntity typeCode='ASSIGNED'
    <Person classCode='PSN'
        <determinerCode root='''>
            <name>...</name>
        </Person>
    <assignedEntity>
        -->
    </author>
</observation>
```

4605

Figure 6.3.4.54.2-1: Observation Request Example**6.3.4.54.2.1 <observation classCode='OBS' moodCode='INT|PRP|GOL'>**

These acts are observations that form the care plan or which can be used in decision support. In intent mood (moodCode='INT') these are what is intended to be performed as part of the care plan. In proposal mood (moodCode='PRP'), these observations are being proposed, for example, as the output of a clinical decision support system. In goal mood (moodCode='GOL'), these observations described the intended goal of a treatment plan.

6.3.4.54.2.2 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.1.20.3.1'>

The <templateId> element identifies this <observation> as an observation request, allowing for validation of the content. The templateId must appear as shown above.

6.3.4.54.2.3 <templateId root=2.16.840.1.113883.10.20.1.25'>

The IHE Observation Request template conforms to the Plan of care activity defined by the HL7 Continuity of Care Document. This template id must be present to indicate conformance.

4620

6.3.4.54.2.4 <id root=' ' extension=' '>

Each observation shall have an identifier.

6.3.4.54.2.5 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '>

4625

Observations shall have a code describing what is to be measured. The code system used is determined by the vocabulary constraints on the types of measurements that might be recorded in a section. Modules that are derived from this one may restrict the code system and code values used for the observation.

6.3.4.54.2.6 <text><reference value="#xxx"/></text> -OR- <text>text</text>

4630

Each observation request entry may contain a <text> element providing the free text that provides the same information as the observation within the narrative portion of the document with a <text> element. For CDA based uses of Observation Requests, this element SHALL be present, and SHALL contain a <reference> element that points to the related string in the

narrative portion of the document. For HL7 Version 3 based uses, the <text> element MAY be included.

6.3.4.54.2.7 <statusCode code='active' />

- 4635 The <statusCode> element shall be present and shall describe the current state of the observation. Goals, intents and proposals that are available for action shall have an 'active' status, but other status values are permitted.

6.3.4.54.2.8 <effectiveTime value=' '/>

- 4640 The <effectiveTime> element shall be present in observation requests to indicate the date and time when the measurement should be taken.

6.3.4.54.2.9 <value xsi:type=' ' .../>

The value of the observation may be recorded using a data type appropriate to the observation to indicate the desired value (e.g., in GOL or PRP mood).

6.3.4.54.2.10 <methodCode code=' ' codeSystem=' ' codeSystemName=' '/>

- 4645 The methodCode element may be used to record the specific method used to make an observation when this information is not already pre-coordinated with the observation code.

6.3.4.54.2.11 <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>

The targetSiteCode may be used to record the target site where the observation should be made when this information is not already pre-coordinated with the observation code.

- 4650 **6.3.4.54.2.12 <author><assignedAuthor classCode='ASSIGNED'>...<assignedAuthor></author>**

In CDA uses, the observation request is assumed to be authored by the same author as the document through context conduction. However, observation requests would often be used to record orders, and in these cases, the author of the order shall be recorded in the author element.

- 4655 For HL7 Version 3 purposes, the <author> element SHOULD be present unless it can be determined by conduction from organizers or higher level structures. When used for HL7 Version 3 the role element name is <assignedEntity> and the author is represented as <assignedPerson> element.

- 4660 **Add Section 6.3.4.55 (Added 2011-09 from QRPH EHCP Profile)**

6.3.4.55 Risk Indicators for Hearing Loss Entry 1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1

This entry describes the Risk Indicators for Hearing Loss.

6.3.4.55.1 Specification

```

<entry>
    <organizer classCode='BATTERY' moodCode='EVN'>
        <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1' />
        <id root=' ' extension=' ' />
        <code code='58232-0' displayName='Hearing Loss Risk Indicators'
              codeSystem='2.16.840.1.113883.6.1'
              codeSystemName='LOINC' />
        <statusCode code='completed' />
        <author>
            <time value=' ' />
            <assignedAuthor>
                <id root=' ' extension=' ' />
            </assignedAuthor>
        </author>
        <component>
            <observation classCode='OBS' moodCode='EVN'>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                :
            </observation>
        </component>
        <component>
            <observation classCode='OBS' moodCode='EVN'>
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.13' />
                :
            </observation>
        </component>
        :
    </organizer>
</entry>

```

4665

Figure 6.3.4.55.1-1: Sample Risk Indicators for Hearing Loss Entry

6.3.4.55.2 <templateId root='1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1' />

The <templateId> element specifies that this organizer entry conforms to the Nursing Interventions battery. The root attribute SHALL contain the value "1.3.6.1.4.1.19376.1.7.3.1.1.15.5.1"

6.3.4.55.3 <organizer classCode='BATTERY' moodCode='EVN' />

Each row in the Nursing Interventions battery SHALL be represented by an organizer with the classCode of 'BATTERY' and the moodCode of 'EVN'

4675 **6.3.4.55.4 <id root=' ' extension=' '/>**

Each battery SHALL have a globally unique identifier.

6.3.4.55.5 <code code='58232-0' codeSystem='2.16.840.1.113883.6.1'/>

The <code> element specifies the LOINC® code that represents the content of the battery. The codeSystem attribute SHALL contain the value '2.16.840.1.113883.6.1'. The code attribute SHALL contain the value='58232-0'. It is good practice to include displayName and codeSystemName for clarity and debugging. The corresponding values are 'Hearing Loss Risk Indicators' and 'LOINC®' respectively.

6.3.4.55.6 <author/><time/><assignedAuthor><id/></assignedAuthor></author>

4685 The <author> relation element points at the author that records the visit battery. This assignedAuthor MAY be different than the author of the document. The time element is used to record when the assigned author recorded the battery.

6.3.4.55.7 <statusCode code='completed'/>

The status code for all batteries SHALL be 'completed'

6.4.4.55.8 <component>

4690 The battery is made of several component Simple Observations. The observation values SHALL be constrained to those coded values and descriptions described by the JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set (1.3.6.1.4.1.19376.1.7.3.1.1.15.2.24).

Add Section 6.3.4.56. (Added 2011-09 from QRPH PRPH-Ca Profile.)

4695 **6.3.4.56 Cancer Diagnosis Entry 1.3.6.1.4.1.19376.1.7.3.1.4.14.1**

A Cancer Diagnosis entry collects details of the patient's cancer diagnosis, including histology, behavior, primary site, laterality, diagnosis date, TNM Stage, and Best Method of Confirmation.

6.3.4.56.1 Parent Template

The parent of this template is Problem Concern Entry (1.3.6.1.4.1.19376.1.5.3.1.4.5.2).

4700 **6.3.4.56.2 Specification**

```

<section>
    <templateId root="2.16.840.1.113883.10.20.1.11"/>
    <templateId root="1.3.6.1.4.1.19376.1.5.3.1.3.6"/>
    <templateId root="1.3.6.1.4.1.19376.1.7.3.1.3.14.1"/>
        <title>"Cancer Diagnosis"</title>
        <text>"Malignant melanoma of the left leg, Stage 1"</text>
        <entry>
            <act classCode='ACT' moodCode='EVN'>
                <templateId root='2.16.840.1.113883.10.20.1.27' />
                <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2' />
                <code nullFlavor='NA' />
                <statusCode code='active' />
                <effectiveTime>
                    <low value="20110101" />
                    <high nullFlavor="NA" />
                </effectiveTime>
                <entryRelationship typeCode="SUBJ" inversionInd="false" >
                    <observation classCode='OBS' moodCode='EVN' negationInd="false" >
                        <templateId root='2.16.840.1.113883.10.20.1.28' />
                        <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5' />
                        <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.14.1" />
                        <code code="282291009" codeSystem="2.16.840.1.113883.6.96"
codeSystemName="SNOMED CT" displayName="Diagnosis"/>
                        <text><reference value="" ></reference></text>
                        <statusCode code="completed" />
                        <effectiveTime>
                            <low value="20110101" />
                            <high nullFlavor="NI" />
                        </effectiveTime>
                    <!--The <value> is the condition that was found.-->
                    <value xsi:type="CD" code="8742" codeSystem="2.16.840.1.113883.3.520.3.2"
codeSystemName="NAACCR Histologic Type" displayName="Lentigo Maligna" >
                    <!--Behavior Qualifier-->
                        <qualifier>
                            <name code="31206-6" codeSystem="2.16.840.1.113883.6.1"
codeSystemName="LOINC" displayName="Behavior ICD-O-3" />
                            <value code="2" codeSystem="2.16.840.1.113883.3.520.3.14"
codeSystemName="NAACCR Behavior Code" displayName="In Situ" />
                        </qualifier>
                        <qualifier>
                            <name code="21861-0" codeSystem="2.16.840.1.113883.6.1"
codeSystemName="LOINC" displayName="Diagnostic Confirmation" />
                            <value xsi:type="CD" code="2"
codeSystem="2.16.840.1.113883.3.520.3.3" codeSystemName="NAACCR Diagnostic Confirmation"
displayName="Positive cytology, no positive histology" />
                        </qualifier>
                    </value>
                    <!--Primary Site -->
                    <targetSiteCode code="C447" codeSystem="2.16.840.1.113883.6.43.1"
codeSystemName="ICD-O-3 (Topography Section)" displayName="Leg" />
                    <!--Laterality-->
                        <qualifier>
                            <name code="20228-3" codeSystem="2.16.840.1.113883.6.1"
codeSystemName="LOINC" displayName="Anatomic part Laterality" />
                            <value code="1" codeSystem="2.16.840.1.113883.3.520.3.1"
codeSystemName="NAACCR Laterality at Diagnosis" displayName="origin of primary: right" />
                        </qualifier>
                    </targetSiteCode>
                    <entryRelationship typeCode="SUBJ" inversionInd="true" >
                    <!--TNM Stage Information-->

```

```

    <observation classCode="OBS" moodCode="EVN">
        <templateId
root="1.3.6.1.4.1.19376.1.7.3.1.4.14.2"/>
        <code code="21908-9" displayName="TNM Clinical Stage
Group" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
<!-- Narrative TNM Clinical Stage -->
        <text> Stage 0 TisNOM0 </text>
        <statusCode code="completed"/>
        <value xsi:type="CD" code="0"
codeSystem="2.16.840.1.113883.3.520.3.9" codeSystemName="NAACCR TNM Clinical Stage Group"
displayName="In Situ">
            <qualifier>

<!--TNM Clinical Stage Descriptor Observation -->
        <name code="21909-7" displayName="TNM
Clinical Stage Descriptor" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
        <value xsi:type="CD" code="0"
codeSystem="2.16.840.1.113883.3.520.3.10" codeSystemName="NAACCR TNM Clinical Stage Descriptor"
displayName="None"/>
            </qualifier>
<!--AJCC TNM Edition Number.-->
        <name code="21917-0" displayName="TNM
Edition Number" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
        <value xsi:type="CD" code="7"
codeSystem="2.16.840.1.113883.3.520.3.5" codeSystemName="NAACCR TNM Edition Number"
displayName="7th Edition"/>
            </value>
        <participant typeCode="PPRF">
            <participantRole>
                <code code="21910-5"
codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Stager.clinical Cancer"/>
                <playingEntity nullFlavor="NA">
                    <code xsi:type="CE" code="1"
codeSystem="2.16.840.1.113883.3.520.3.4" codeSystemName="TNM Clinical Staged By"
displayName="Managing Physician"/>
                </playingEntity>
            </participantRole>
        </participant>
        <entryRelationship typeCode="COMP">

<!-- 6.3.4.62 TNM Clinical Tumor Observation-->
        <observation classCode="OBS" moodCode="EVN">
            <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.13"/>
            <code code="21905-5" displayName="TNM Clinical T"
codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
            <statusCode code="completed"/>
            <value xsi:type="CD" code="Tis"
codeSystem="2.16.840.1.113883.3.520.3.6" codeSystemName="NAACCR TNM Clinical Tumor"
displayName="In Situ"/>
        </observation>
    </entryRelationship>

<!--6.3.4.63 TNM Clinical Nodes Observation -->
        <entryRelationship typeCode="COMP">
            <observation classCode="OBS" moodCode="EVN">
                <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.13"/>
                <code code="21906-3" displayName="TNM Clinical N"
codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
                <statusCode code="completed"/>
                <value xsi:type="CD" code="N0"
codeSystem="2.16.840.1.113883.3.520.3.7" codeSystemName="NAACCR TNM Clinical Nodes"
displayName="None"/>
            </observation>
        </entryRelationship>

```

```

4835 <!--6.3.4.64 TNM Clinical Metastases Observation-->
        <entryRelationship typeCode="COMP">
            <observation classCode="OBS" moodCode="EVN">
                <templateId root="1.3.6.1.4.1.19376.1.5.3.1.4.13"/>
                    <code code="21907-1" displayName="TNM Clinical M"
4840 codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/>
                    <statusCode code="completed"/>
                    <value xsi:type="CD"
4845 codeSystem="2.16.840.1.113883.3.520.3.8" codeSystemName="NAACCR TNM Clinical Metastases"
code="M0" displayName="None"/>
                </observation>
            </entryRelationship>
        </observation>
    </entryRelationship>
</observation>
</entryRelationship>
</act>
</entry>
</section>

```

Figure 6.3.4.56.2-1: Sample Cancer Diagnosis Entry**6.3.4.56.3 <act classCode='ACT' moodCode='EVN'>**

4855 All concerns reflect the act of recording (<act classCode='ACT'>) the event (moodCode='EVN') of being concerned about a problem, allergy or other issue about the patient condition.

6.3.4.56.4 <templateId root='2.16.840.1.113883.10.20.1.27'> <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5.2'>

4860 These template identifiers indicates this entry conforms to the concern content module. This content module inherits constraints from the HL7 CCD Template for problem acts, and so also includes that template identifier.

6.3.4.56.5 <!-- 1..* entry relationships identifying problems of concern --><entryRelationship type='SUBJ'><observation classCode='OBS' moodCode='EVN'><templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5'>...</observation>

4865 >

This entry shall contain one or more problem entries that conform to the Problem Entry template 1.3.6.1.4.1.19376.1.5.3.1.4.5. The typeCode SHALL be “SUBJ” and inversionInd SHALL be “false”.

6.3.4.56.6 <observation classCode="OBS" moodCode="EVN">

4870 The <observation> classCode and moodCode SHALL be recorded as shown above.

6.3.4.56.7 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.5'> <templateId root='1.3.6.1.4.1.19376.1.7.3.1.4.14.1'>

4875 These <templateId> elements identify this <entry> as a cancer diagnosis entry and its parent, Problem Entry, allowing for validation of the content. The <templateId> elements shall be recorded as shown above.

6.3.4.56.8 <code code="282291009" codeSystem=" 2.16.840.1.113883.6.96" codeSystemName="SNOMED CT" displayName="Diagnosis"/>

4880 The <code> element indicates that this is the Diagnosis information. This code SHALL be the SNOMED CT code “282291009” for “Diagnosis”. It is good style to include the displayName and codeSystemName to help debugging.

6.3.4.56.9 <statusCode code='completed' />

The status code for all Cancer Diagnosis Entries SHALL be ‘completed’.

6.3.4.56.10 <effectiveTime value="xxx"/>

4885 This element records the date of initial diagnosis by a recognized medical practitioner for the cancer being reported.

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

4890 The <value> records the Histologic Type, which is the cell type of the tumor/cancer (e.g., carcinoma, melanoma, sarcoma, lymphoma, leukemia). This element is required. It is always represented using the CD datatype (xsi:type='CD'), even though the value may be a coded or uncoded string. If coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

6.3.4.56.12 <qualifier><name code="31206-6" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Behavior ICD-O-3 Cancer"/><value code="" codeSystem="" codeSystemName="" displayName="" /> </qualifier>

4900 This <qualifier> provides Behavior information, indicating whether the tumor is benign, in situ, malignant or metastatic. The code and codeSystem attributes SHALL be recorded exactly as shown above. If coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

6.3.4.56.13 <qualifier><name code="21861-0" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Dx confirmed by Cancer"/><value xsi:type="CD" code="" codeSystem="" codeSystemName="" displayName="" /> </qualifier>

4905 This <qualifier> provides Best Method of Diagnosis information, indicating the best method used to confirm the presence of the cancer being reported. The code and codeSystem attributes SHALL be recorded exactly as shown above. The <value> records the best method of diagnosis, and if coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

4910 **6.3.4.56.14 <targetSiteCode code="" codeSystem="" codeSystemName="" displayName="">**

The <targetSiteCode> element SHALL be present and shall indicate the anatomic location where the primary tumor originated. Vocabulary used SHALL follow the appropriate realm constraints. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

4915 **6.3.4.56.15 <qualifier><name code="20228-3" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Anatomic part Laterality"/> <value code="" codeSystem="" codeSystemName="" displayName="" /></qualifier>**

4920 This <qualifier> provides the laterality, which indicates the side of a paired organ or side of the body on which the reportable tumor originated. The code and codeSystem attributes SHALL be recorded exactly as shown above. The <value> records the laterality, if coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

6.3.4.56.16 <entryRelationship typeCode="SUBJ" inversionInd="false">

4925 One <entryRelationship> element should be present providing information on the TNM Clinical Stage.

When present, this <entryRelationship> element SHALL contain an observation conforming to the TNM Stage Information (1.3.6.1.4.1.19376.1.7.3.1.4.14.2) template. The typeCode SHALL be “SUBJ” and inversionInd SHALL be “false”.

4930 **6.3.4.56.17 <observation classCode="OBS" moodCode="EVN"> <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.14.2"/> [1st nesting]**

Observations that describe the TNM Stage Information SHALL be included if known.

6.3.4.56.18 <code code="75620-5" displayName="TNM Clinical Stage Information" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> [1st nesting]

4935 The <code> element indicates that this observation is the TNM Clinical Stage Information. This code SHALL be the LOINC code 75620-5. It is good style to include the displayName and codeSystemName to help debugging.

6.3.4.56.19 <statusCode code="completed"/> [1st nesting]

The status code for all TNM Clinical Stage Information observations SHALL be ‘completed’.

4940 **6.3.4.56.20 <value xsi:type="CD" code="" codeSystem="" codeSystemName="" displayName="" /> [1st nesting]**

The <value> records the TNM Clinical Stage Group, which is a detailed site-specific code for the clinical stage group as defined by AJCC and recorded by the physician. This element is required. It is always represented using the CD datatype (xsi:type='CD'), even though the value

- 4945 may be a coded or uncoded string. If coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)
- 6.3.4.56.21 <qualifier><name code="21909-7" displayName=" Descriptor.clinical Cancer" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/><value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> [1st nesting]**
- 4950 This <qualifier> provides TNM Clinical Stage Descriptor information, indicating The AJCC clinical stage prefix/suffix recorded by the physician. AJCC stage descriptors identify special cases that require separate analysis. The code and codeSystem attributes SHALL be recorded exactly as shown above. The <value> records the TNM Clinical Stage Descriptor, and if coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)
- 6.3.4.56.22 <qualifier><name code="21917-0" displayName="Version TNM Classification" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/><value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/></qualifier> [1st nesting]**
- 4960 This <qualifier> provides TNM Edition Number information, indicating the edition number of the AJCC Staging Manual. The code and codeSystem attributes of <name> SHALL be recorded exactly as shown above. If coded, it SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)
- 4965 **6.3.4.56.23 <participant typeCode="PPRF"> <participantRole> <code code="21910-5" codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC" displayName="Stager.clinical Cancer"/><playingEntity nullFlavor="NA"> <code xsi:type="CE" code="" codeSystem="" codeSystemName=" " displayName=" "/> [1st nesting]**
- 4970 This <participant> element should specify the person who recorded the AJCC staging elements and stage group in the patient's medical record. The code and codeSystem attributes for <participantRole> SHALL be recorded exactly as shown above. The <code> attribute of <playingEntity> identifies the person who recorded the staging elements, and SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)
- 4975 **6.3.4.56.24 <!-- 0..3 entryRelationships identifying simple observations for TNM Clinic Tumor, TNM Clinical Nodes, and TNM Clinical Metastases--><entryRelationship typeCode="COMP" inversionInd="false"><observation classCode='OBS'moodCode='EVN'><templateIDroot='1.3.6.1.4.1.19376.1.5.3.1.4.13 />...</observation>[2nd nesting]**
- 4980 Each <entryRelationship> element should contain a simple observation that specifies the TNM Clinic Tumor, TNM Clinical Nodes, and TNM Clinical Metastases, each of which is a component of the TNM Stage Group. Simple observations that describe the TNM Clinic Tumor,

4985 TNM Clinical Nodes, and TNM Clinical Metastases SHALL be included if known and inversionInd SHALL be “false”.

6.3.4.56.25 <code code="" displayName=" " codeSystem="2.16.840.1.113883.6.1" codeSystemName="LOINC"/> [2nd nesting]

Observations SHALL include one of following LOINC values to indicate the component of TNM Stage Group represented in the Observation.

4990

LOINC Code	Display Name	Description
21905-5	TNM Clinical T	A detailed site-specific code for the clinical tumor (T) as defined by AJCC and recorded by the physician.
21906-3	TNM Clinical N	A detailed site-specific code for the clinical nodes (N) as defined by AJCC and recorded by the physician.
21907-1	TNM Clinical M	A detailed site-specific staging code for the clinical metastases (M) as defined by AJCC and recorded by the physician.

6.3.4.56.26 <value xsi:type="CD" code="" codeSystem="" codeSystemName=" " displayName=" "/>

4995 The <value> of the observation SHALL be recorded using the vocabulary appropriate to the coded observation according to the table above and SHALL follow the appropriate realm constraints for vocabulary. (See Volume 4 in the QRPH PRPH-Ca Profile found at <http://www.ihe.net>.)

6.3.4.57 Patient Transfer 1.3.6.1.4.1.19376.1.5.3.1.1.25.1.4.1

The Patient Transfer entry shall record the transfer of the patient to an internal department or external entity such as a different hospital or skilled nursing facility.

5000 **6.3.4.57.1 Parent Template**

6.3.4.57.2 Specification

```

<act classCode='ACT' moodCode='EVN'>
  <templateId root='PatientTransferAct' />
  <id/>
  <!-- code is fixed -->
  <code code='107724000' displayName='patient transfer' codeSystem='2.16.840.1.113883.6.96' />
  <effectiveTime value='' />
  <participant typeCode='DST'>
    <templateId root='destinationLocation' />
    <participantRole classCode='SDLOC'>
      <id/>
      <code/>
      <addr/>
      <telecom/>
      <playingEntity classCode='ENT'>
        <name/>
        </playingEntity>
      </participantRole>
    </participant>
  </act>

```

Figure 6.3.4.57.2-1: Sample Cancer Diagnosis Entry

6.3.4.57.3 <act classCode='ACT' moodCode='INT|EVN'>

The transfer is recorded in an act element, to describe a patient transfer. In intent mood (moodCode='INT'), this records the expected transfer of the patient. In event mood

5025 (moodCode='EVN'), this records the actual transfer.

6.3.4.57.4 <templateId root='TBD' />

The templateId indicates that this transfer entry conforms to the constraints of this content module.

6.3.4.57.5 <id root=" extension="/" />

5030 This required element shall contain an identifier.

6.3.4.57.6 <code code="" displayName="" codeSystem="" codeSystemName="" />

The code shall be code='107724000' displayName='patient transfer' codeSystem='2.16.840.1.113883.6.96' />

6.3.4.57.7 <text><reference value="#xxx"/></text>

5035 The <text> element shall contain a reference to the narrative text describing the transfer of the patient.

6.3.4.57.8 statusCode

<statusCode code='normal|completed' /> When the transfer act has occurred (moodCode='EVN'), the statusCode element shall be present, and shall contain the value 'completed'. When the

5040 transfer act is intended (moodCode='EVN') the statusCode element shall contain the value 'normal'.

6.3.4.57.9 <effectiveTime><low value="/"><high value="/"><effectiveTime/>

When the transfer has occurred, this element shall be sent, and indicates the effective time for the transfer. This element may be sent to record when the transfer act is intended to occur. The <low> element records the time at which the transfer process was started. The <high> value records the time at which the transfer was completed.

6.3.4.57.10 participant

The <participant> element encodes the destination with a typeCode of DST

<participant typeCode='DST'>

5050 **6.3.4.57.11 templateId**

The template id identifies the facility or department which is the transfer destination.

<templateId root='destinationLocation' />

6.3.4.57.12 participantRole

The participant role is fixed to <participantRole classCode='SDLOC'>

5055 **6.3.4.57.13 <id root=" extension=""/>**

The <id> element shall be sent when the transfer has occurred, and identifies the performer of the act.

6.3.4.57.14 <code>

The code shall indicate the type of healthcare service location for the transfer destination.

5060 **6.3.4.57.15 <addr></addr>**

The <addr> element may be sent to provide a contact postal address for the performer of the disposition.

6.3.4.57.16 <telecom>

5065 The <telecom> element may be sent to provide a contact postal address for the performer of the disposition.

6.3.4.57.17 playingEntity

The playing entity classCode shall be ENT <playingEntity classCode='ENT'>

6.3.4.57.18 name

5070 The name element of the playing entity shall record the name of the facility or departmental destination.

Add section 6.3.4.58 (added 2013-09 from the QRPH VRDR supplement.)

6.3.4.58 Death Pronouncement Entry Content Module (1.3.6.1.4.1.19376.1.7.3.1.4.23.1)

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.23.1]

5075 The template contains information on the pronouncement of death on the death certificate.

1. **SHALL** contain exactly one [1..1] **@classCode**
2. **SHALL** contain exactly one [1..1] **@moodCode**
3. **SHALL** contain exactly one [1..1] **templateId** (CONF:7136) such that it
 - a. **SHALL** contain exactly one [1..1]
@root="2.16.840.1.113883.10.20.22.4.2" (CONF:9138)
 - b. **SHALL** contain exactly one [1..1] code/@code="58325-2" Provider witnessed decedent's death (CodeSystem: 2.16.840.1.113883.6.1 LOINC).
4. **SHALL** contain zero or one [1..1] **effectiveTime**

5080 Provide the date and time at which the decedent was pronounced dead. The first id represents this specific globally unique result observation.

5. **SHALL** contain exactly one [1..1] **performer**
 - a. This performer **SHALL** contain exactly one [1..1] **@typeCode**="PRF"
 - b. This performer **SHALL** contain exactly one [1..1] **assignedEntity**
 - c. This assignedEntity **SHALL** contain exactly one [1..1] **@classCode**="ASSIGNED"
 - d. This assignedEntity **SHALL** contain exactly one [1..1] **addr**

The postal address used to locate the clinician or pronouncing the death at the time of death certification.

6. This assignedEntity **SHALL** contain exactly one [1..1] assignedPerson
 - a. This assignedPerson **SHALL** contain exactly one [1..1] **@classCode**="PSN"
 - b. This assignedPerson **SHALL** contain exactly one [1..1] **determinerCode**="INSTANCE"
 - c. This assignedPerson **SHALL** contain exactly one [1..1] **ID**

5100 *This field shall contain the License Number of Person Pronouncing Death*

- d. This assignedPerson **SHALL** contain exactly one [1..1] **name**

This field is valued with the person who pronounced the death. The full name of the pronouncer is required.

5105

```

5110 <entry>
      <observation classCode="OBS" moodCode="EVN">
        <templateId root="1.3.6.1.4.1.19376.1.7.3.1.4.23.1"/>
        <id root="" />
        <code code="58325-2" codeSystem="2.16.840.1.113883.6.1"
          codeSystemName="LOINC" displayName=" Provider witnessed decedent's death "/>
        <effectiveTime>
          <low value="201311141201"/>
          <high value="201311141201"/>
        </effectiveTime>
      </observation>
    </entry>
  
```

Figure 6.3.4.58-1: Death Pronouncement Entry Content Module example

5120

*Add section 6.3.4.59 (added 2013-09 from the QRPH VRDR supplement.)***6.3.4.59 Death Location Type Entry Content Module**

[Observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.23.2]

5125 This template makes it possible to record the type of location (e.g., hospital inpatient room) at which the person died.

1. **SHALL** contain exactly one [1..1] **@classCode="OBS"** Observation (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass)
2. **SHALL** contain exactly one [1..1] **@moodCode="EVN"** Event (CodeSystem: 2.16.840.1.113883.5.1001 HL7ActMood)
3. **SHALL** contain exactly one [1..1] **code/@code=" 58332-8"** (CodeSystem: 2.16.840.1.113883.6.1 LOINC)
4. **SHALL** contain exactly one [1..1] **value**, which **SHALL** be selected from ValueSet
5. Death Location Type Codes (1.3.6.1.4.1.19376.1.7.3.1.13.8.4) STATIC, where its data type is CE
6. A code value to indicate the type of location where the patient died.

5135

5140

5145 <observation xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
 xmlns="urn:hl7-org:v3" xsi:schemaLocation="urn:hl7-org:v3 CDA.xsd"
 classCode="OBS" moodCode="EVN">
 <id root="1536492804"/>
 <code code="58332-8" codeSystem="2.16.840.1.113883.6.1"
 codeSystemName="LOINC"/>
 5150 <effectiveTime>
 <low value="2012"/>
 <high value="2012"/>
 </effectiveTime>
 <value xsi:type="CE" code="Value"/>
 </observation>

5155

Figure 6.3.4.59-1: Death Location Type Entry Content Module example

Delete the following section 6.3.4.61 Employment Status Organizer:

5160 *Delete the following section 6.3.4.62 Usual Occupation and Industry Organizer:*

Delete the following section 6.3.4.63 History of Occupation Organizer:

Replace the following section 6.3.4.64 Employment Status Observation

5165 **6.3.4.64 History of Employment Status Observation**

**Table 6.3.4.64-1: History of Employment Status Observation Entry
 1.3.6.1.4.1.19376.1.7.3.1.4.24.18**

Template Name	History of Employment Status Observation Entry
Template ID	1.3.6.1.4.1.19376.1.7.3.1.4.24.18
Parent Template	
General Description	A History of Employment Status Observation entry is a clinical statement about a person's state of being employed at the point in time the statement is recorded. Awareness of the subject's History of Employment Status can assist in understanding the subject's resources, access to benefits, and demands at home and work. Generally, employment status refers to whether or not a person currently has a job. In a healthcare setting employment status may be used to determine appropriate probing questions for occupational hazards and occupational history. For example, someone who is unemployed or has chosen not to work may be prompted to provide information about previous jobs. History of Employment status is not the same as compensation and sector employment type described in the Work Classification Observation Entry.

Class/Mood	Code	Data Type	Value		
ClassCode= "OBS" MoodCode= "EVN"	Code = 74165-2 Display Name = History of Employment Status CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC	Observation	Value xsi:type = "CD" from concept domain CD_EmploymentStatus defined in Table 6.6-1		
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

[observation: templateId 1.3.6.1.4.1.19376.1.5.3.1.4.20.4 (open)]

5170 An Employment Status Entry is a clinical statement about the subject's employment status at the point in time the statement is recorded.

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

3. **SHALL** contain exactly one [1..1] templateId such that it
 - a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.5.3.1.4.20.4".

5180 4. **SHALL** contain at least one [1..*] id.
 5. **SHALL** contain exactly one [1..1] code.

- a. **SHALL** be 74165-2 (History of Employment Status) from LOINC (codeSystem 2.16.840.1.113883.6.1).

5185 6. **SHALL** contain exactly one [1..1] statusCode="completed" (CodeSystem: ActStatus 2.16.840.1.113883.5.14).

7. **SHALL** contain exactly one [1..1] effectiveTime.
 - a. This effectiveTime **SHOULD** contain zero or one [0..1] low.
 - i. Note: The effectiveTime/low asserts when the employment status began.

5190 b. This effectiveTime **SHOULD** contain zero or one [0..1] high

- i. Note: The effectiveTime/high asserts when the employment status ended. If employment status is current, effectiveTime/high should be omitted. Note: The ending time <high> element SHALL not be greater than the time the observation is made.
- ii. Note: If the effectiveTime/high is unknown, use @nullFlavor="UNK" (2.16.840.1.113883.5.1008 (HL7NullFlavor) = UNK)

8. **SHALL** contain exactly one [1..1] value with @xsi:type="CD"

- a. This value SHOULD contain zero or one [0..1] @code
 b. This value **SHALL** be selected from Concept Domain CD_EmploymentStatus

5200

Replace the following section 6.3.4.65: Usual Occupation and Industry Observation

6.3.4.65 Usual Occupation Observation Entry

Table 6.3.4.65-1: Usual Occupation Observation Entry 1.3.6.1.4.1.19376.1.7.3.1.4.24.20

Template Name	Usual Occupation Observation Entry				
Template ID	1.3.6.1.4.1.19376.1.7.3.1.4.24.20				
Parent Template					
General Description	A Usual Occupation Observation Entry contains information about the occupation which the subject has held for the longest duration through his or her working history, at the point in time the statement is recorded. A history of this observation is not retained. Longest-held occupations can be associated with conditions that develop slowly over time or even after the person is no longer performing that type of work, e.g., some respiratory conditions and cancers. It optionally includes a total duration observation because a person can be in and out of a given occupation over time. In addition, knowing when the person began working in this occupation can provide information about potential exposures and allows the clinician to assess whether sufficient time has elapsed for a chronic condition to appear, i.e., the latency period. This guides appropriate use of screening tests to detect early disease.				
Class/Mood	Code		Data Type	Value	
ClassCode= "OBS" MoodCode= "EVN"	Code = 21843-8 Display Name = Usual Occupation CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC		Observation	Value xsi:type = "CD" from concept domain CD_OccupationCode defined in Table 6.6-1	
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

5205

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.20 (open)]

A Usual Occupation Observation Entry contains information about the occupation which the subject has held for the longest duration through his or her working history, at the point in time the statement is recorded

5210

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

- 5215 3. **SHALL** contain exactly one [1..1] **templateId** such that it
- SHALL** contain exactly one [1..1]
`@root="1.3.6.1.4.1.19376.1.7.3.1.4.24.20".`
- 5220 4. **SHALL** contain at least one [1..*] **id**.
- 5220 5. **SHALL** contain exactly one [1..1] **code**.
- SHALL** be 21843-8 (Usual Occupation) from LOINC (codeSystem 2.16.840.1.113883.6.1).
- 5225 6. **SHALL** contain exactly one [1..1] **statusCode**="completed" (CodeSystem: ActStatus 2.16.840.1.113883.5.14).
- 5225 7. **SHALL** contain exactly one [1..1] **effectiveTime**.
- This effectiveTime **SHALL** contain exactly one [1..1] low.
 Note: The effectiveTime/low asserts when the usual occupation began.
- 5230 Note: If the effectiveTime/low is unknown, use @nullFlavor="UNK" (2.16.840.1.113883.5.1008 (HL7NullFlavor) = UNK)
- 5230 b. This effectiveTime **MAY** contain zero or one [0..1] high.
 Note: The effectiveTime/high asserts when the usual occupation ended. If usual occupation is current, effectiveTime/high should be omitted.
- 5235 Note: If the effectiveTime/low is unknown, use @nullFlavor="UNK" (2.16.840.1.113883.5.1008 (HL7NullFlavor) = UNK).
- 5235 8. **SHALL** contain exactly one [1..1] **value** with @xsi:type="CD"
- This value **SHALL** be selected from Concept Domain CD_OccupationCode.
- 5240 9. **MAY** contain zero or one [0..1] **subject**
- The subject, if present, **SHALL** contain exactly one [1..1] **relatedSubject**
 - This relatedSubject **SHALL** contain exactly one [1..1] `@classCode="PRS"` (CodeSystem: HL7EntityClass urn:oid:2.16.840.1.113883.5.41)
 - This relatedSubject **SHALL** contain exactly one [1..1] **code**
 - This code **SHALL** contain exactly one [1..1] `@code` (ValueSet: Family Member Value Set urn:oid:2.16.840.1.113883.1.11.19579 DYNAMIC)
- 5245 10. **SHALL** contain at least one [1..*] **author**
- Such authors **SHALL** contain exactly one [1..1] time
 Note: The author/time asserts when the usual occupation was authored or last updated in the patient's chart.
- 5250 11. **SHOULD** contain zero or one [0..1] **entryRelationship** such that it
- SHALL** contain exactly one [1..1] `@typeCode="REFR"` (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002 STATIC).
 - SHALL** contain exactly one [1..1] Usual Occupation Duration Observation Entry (1.3.6.1.4.1.19376.1.7.3.1.4.24.25)
- 5250 12. **SHOULD** contain zero or one [0..1] **entryRelationship**

- 5255 a. The entryRelationship, if present, **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType urn:oid:2.16.840.1.113883.5.1002)
- 5260 b. The entryRelationship, if present, **SHALL** contain exactly one [1..1] [usual Industry Observation](#) (identifier: urn:oid: 1.3.6.1.4.1.19376.1.7.3.1.4.24.26)

Replace the following section 6.3.4.66 Occupation Observation

6.3.4.66 Past or Present Occupation Observation Entry

5265 **Table 6.3.4.66-1: Past or Present Occupation Observation Entry**
1.3.6.1.4.1.19376.1.7.3.1.4.24.19

Template Name		Past or Present Occupation Observation Entry					
Template ID		1.3.6.1.4.1.19376.1.7.3.1.4.24.19					
Parent Template							
General Description		A Past or Present Occupation Observation entry is a clinical statement about a job or jobs which the subject currently holds or has held in the past. It includes related observations about the occupation (type of work), the type of business (industry) in which that occupation is performed, supervisory level (including military pay grade), and the employer's name and location. It should also include observations about the job's work classification (e.g., self-employed, volunteer) and work schedule, and may also contain observations for job duties and occupational hazards. For a given job, updates to Industry, Occupation, Employer, or Supervisory Level would constitute a new 'job'. The type of work a person performs (occupation) and their industry (type of business in which they work) are critical data elements for patient care, population health, and public health, with the current information being the most important. In the health care encounter, current occupation and industry are important because they provide information regarding the exposures a person may have to substances/environments/hazards that may cause illness/injury or may impact the treatment plan. The combination of occupation and industry serves as a key indicator of the patient's work environment. The entry is designed to ensure that these data remain associated with one-another in perpetuity, even if multiple jobs are included. Note that occupation and industry also describe self-reported service in the armed forces.					
Class/Mood	Code	Data Type	Value	ClassCode= "OBS" MoodCode= "EVN"	Code = 11341-5 Display Name = History of Occupation CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC	Observation	Value xsi:type = "CD" from concept domain CD_OccupationCode defined in Table 6.6-1
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint		

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.19 (open)]

- 5270 A Past or Present Occupation Observation Entry is a clinical statement about a job which the subject currently holds or has held in the past. Multiple Past or Present Occupation Observation Entries may be needed to reflect a person's current jobs, since many people hold more than one job at a time. Over time, a history of jobs is to be built since past jobs can be related to latent health effects.
- 5275 1. **SHALL** contain exactly one [1..1] @classCode="OBS" (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass) .
2. **SHALL** contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001).
3. **SHALL** contain exactly one [1..1] **templateId** such that it
- 5280 a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.7.3.1.4.24.19".
4. **SHALL** contain at least one [1..*] **id**.
5. **SHALL** contain exactly one [1..1] **code**.
- a. **SHALL** be 11341-5 (History of Occupation) from LOINC (codeSystem 2.16.840.1.113883.6.1) .
- 5285 6. **SHALL** contain exactly one [1..1] **statusCode**= (CodeSystem: ActStatus 2.16.840.1.113883.5.14) .
- a. Note: Indicate current job as 'active'. Indicate historical jobs as 'completed'
- 5290 7. **SHALL** contain exactly one [1..1] **effectiveTime**.
- a. This effectiveTime **SHALL** contain exactly zero or one [1..1] **low**.
- i. Note: The effectiveTime/low asserts when the past or present occupation began.
- ii. Note: If the effectiveTime/low is unknown, use @nullFlavor="UNK" (2.16.840.1.113883.5.1008 (HL7NullFlavor) = UNK)
- b. This effectiveTime **MAY** contain exactly zero or one [0..1] **high**.
- i. The ending time <high> element **SHALL** not be greater than the time the observation is made.
- ii. Note: The effectiveTime/high asserts when the past or present occupation ended. If occupation is current, effectiveTime/high should be omitted.
- iii. Note: If the effectiveTime/high is unknown, use @nullFlavor="UNK"
- 5295 8. **SHALL** contain exactly one [1..1] **value** with @xsi:type="CD"
- a. This value **SHALL** be selected from Concept Domain CD_OccupationCode.
- 5300 5305 9. **MAY** contain zero or one [0..1] **subject**
- a. The subject, if present, **SHALL** contain exactly one [1..1] **relatedSubject**

- 5310 i. This relatedSubject **SHALL** contain exactly one [1..1] @classCode="PRS" (CodeSystem: HL7EntityClass urn:oid:2.16.840.1.113883.5.41)
ii. This relatedSubject **SHALL** contain exactly one [1..1] **code**
1. This code **SHALL** contain exactly one [1..1] @code (ValueSet: Family Member Value Set urn:oid:2.16.840.1.113883.1.11.19579 **DYNAMIC**)
5315 Note: This represents the Family Relationship of the person holding this occupation.
10. **SHALL** contain exactly one [1..1] **participant** such that it
5320 a. **SHALL** contain exactly one [1..1] @typeCode="IND"
b. **SHALL** contain exactly one [1..1] **participantRole**
i. Which **MAY** contain exactly one [1..1] @classCode="ROL" (CodeSystem: RoleCode 2.16.840.1.113883.5.111 **STATIC**).
ii. Which **SHOULD** contain exactly zero or one [0..1] **id**
5325 1. Such that the id **SHALL** reference the id of an AssociatedEntity in the header which **SHALL** contain exactly one [1..1] templateId such that it
a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.5.3.1.2.2" (IHE Employer and School Contacts template)
b. **SHALL** contain exactly one [1..1] @extension="2016-11-30".
5330 2. The AssociatedEntity **SHOULD** contain zero or one [0..1] name.
3. The AssociatedEntity **SHOULD** contain zero or one [0..1] addr
5335 iii. This participantRole **SHOULD** contain zero or one [0..1] **addr**
Note: Contains the address of the employer
iv. This participantRole **MAY** contain zero or one [0..1] **playingEntity**
5340 1. The playingEntity, if present, **SHOULD** contain zero or one [0..1] name
Note: Contains the name of the employer
11. **SHALL** contain exactly one [1..1] **entryRelationship** such that it
5345 a. **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType uri:oid:2.16.840.1.113883.5.1002 **STATIC**).
b. **SHALL** contain exactly one [1..1] Past or Present Industry Observation Entry (1.3.6.1.4.1.19376.1.7.3.1.4.24.3)
12. **SHOULD** contain zero or one [0..1] **entryRelationship** such that it
5350 a. **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType uri:oid:2.16.840.1.113883.5.1002 **STATIC**).

- 5355 b. **SHALL** contain exactly one [1..1] Work Classification Observation Entry (1.3.6.1.4.1.19376.1.7.3.1.4.24.4)
13. **SHOULD** contain zero or one [0..1] **entryRelationship** such that it
- a. **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType 2.16.840.1.113883.5.1002 **STATIC**).
 - b. **SHALL** contain exactly one [1..1] Work Schedule Observation (1.3.6.1.4.1.19376.1.7.3.1.4.24.5).
- 5360 14. **SHOULD** contain zero or one [0..1] **entryRelationship**.
- a. The entryRelationship, if present, **SHALL** contain exactly one [1..1] @typeCode="REFR"
 - b. The entryRelationship, if present, **SHALL** contain exactly one [1..1] Supervisory Level (identifier: urn:oid: 1.3.6.1.4.1.19376.1.7.3.1.4.24.16)
- 5365 15. **MAY** contain zero or one [0..*] **entryRelationship**
- a. The entryRelationship, if present, **SHALL** contain exactly one [1..1] @typeCode="REFR"
 - b. The entryRelationship, if present, **SHALL** contain exactly one [1..1] Job Duty Observation (identifier: urn:oid: 1.3.6.1.4.1.19376.1.7.3.1.4.24.14)
- 5370 16. **MAY** contain zero or one [0..*] **entryRelationship**.
- a. The entryRelationship, if present, **SHALL** contain exactly one [1..1] @typeCode="REFR"
 - The entryRelationship, if present, **SHALL** contain exactly one [1..1] Occupational Hazard Observation (identifier: urn:oid: 1.3.6.1.4.1.19376.1.7.3.1.4.24.27)

Replace the following section 6.3.4.67 Work Shift Observation – renamed to Work Schedule

5380

6.3.4.67 Work Schedule Observation Entry

Table 6.3.4.67-1: Work Schedule Observation Entry 1.3.6.1.4.1.19376.1.7.3.1.4.24.5

Template Name	Work Schedule Observation Entry
Template ID	1.3.6.1.4.1.19376.1.7.3.1.4.24.5
Parent Template	
General Description	A clinical statement about the schedule, “shift”, or typical time within a work-day in which a person is scheduled to perform their duties. It includes observations of the hours and days worked per week. Full-time and part-time designations are not defined consistently and would not reflect compressed schedules, long work hours, or overtime

		work. Use cases include care for a patient with diabetes who is on a rotating shift and needs different counseling on diet and medication management than someone working a regular day shift; a patient on a rotating shift who has fatigue interfering with activities at work and home; a patient with obesity working long hours.			
Class/Mood	Code	Data Type	Value		
ClassCode= "OBS" MoodCode= "EVN"	Code = 74159-5 Display Name = Work Schedule CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC	Observation	Value xsi:type = “CD” from concept domain CD_WorkSchedule defined in Table 6.6-1		
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.5 (open)]

5385 A clinical statement about the schedule, “shift”, or typical time within a work-day in which a person is scheduled to perform their duties.

1. **SHALL** contain exactly one [1..1] @classCode="OBS" (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass).
2. **SHALL** contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001).
3. **SHALL** contain exactly one [1..1] templateId such that it
 - a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.7.3.1.4.24.5".
4. **SHALL** contain at least one [1..*] id.
5. **SHALL** contain exactly one [1..1] code.
 - a. **SHALL** be 74159-5 (Workshift) from LOINC.
6. **SHALL** contain exactly one [1..1] statusCode="completed" Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14).
7. **SHALL** contain exactly one [1..1] value with @xsi:type="CD".
 - a. This value **SHALL** contain exactly one [1..1] @code, which **SHALL** be selected from Concept Domain CD_EmploymentStatus
8. **MAY** contain zero or one [0..1] entryRelationship such that it
 - a. **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType uri:oid:2.16.840.1.113883.5.1002 STATIC).
9. **MAY** contain zero or one [0..1] entryRelationship such that it
 - a. **SHALL** contain exactly one [1..1] @typeCode="REFR" (CodeSystem: HL7ActRelationshipType uri:oid:2.16.840.1.113883.5.1002 STATIC).

- 5410 b. **SHALL** contain exactly one [1..1] Weekly Work Days Observation Entry
 (1.3.6.1.4.1.19376.1.7.3.1.4.24.7)

Replace the following section 6.3.4.68 Weekly Work Hours Observation

5415

6.3.4.68 Weekly Work Hours Observation Entry

Table 6.3.4.68-1: Weekly Work Hours Observation Entry 1.3.6.1.4.1.19376.1.7.3.1.4.24.6

Template Name		Weekly Work Hours Observation Entry			
Template ID		1.3.6.1.4.1.19376.1.7.3.1.4.24.6			
Parent Template					
General Description		A clinical statement about the typical number of hours per week that a person spends performing their duties for work. This information is most useful coupled with weekly work days and helps to reveal compressed schedules, long work hours, and overtime.			
Class/Mood		Code	Data Type	Value	
ClassCode= "OBS" MoodCode= "EVN"		Code = 74161-1 Display Name = Weekly Work Hours CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC	Observation	value with @xsi:type="INT"	
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.6 (open)]

5420

A clinical statement about the typical number of hours per week that a person spends performing their duties for work.

1. **SHALL** contain exactly one [1..1] @classCode="OBS" (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass).
2. **SHALL** contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001).
3. **SHALL** contain exactly one [1..1] templateId such that it
 - a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.7.3.1.4.24.6".
4. **SHALL** contain at least one [1..*] id.
5. **SHALL** contain exactly one [1..1] code.

5430

- 5435
- a. **SHALL** be 74161-1 (Weekly Work Hours) from LOINC.
 - 6. **SHALL** contain exactly one [1..1] **statusCode**="completed" Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14).
 - 7. **SHALL** contain exactly one [1..1] **value** with @xsi:type="INT".
 - a. This value **SHALL** contain exactly one [1..1] **@value**, which represents the number of hours in a week that a person usually works.

Replace the following section 6.3.4.69 Usual Occupation Duration

6.3.4.69 Usual Occupation Duration Entry

5440 **Table 6.3.4.69-1: Usual Occupation Duration Entry (1.3.6.1.4.1.19376.1.7.3.1.4.24.10)**

Template Name		Usual Occupation Duration Entry			
Template ID		1.3.6.1.4.1.19376.1.7.3.1.4.24.10			
Parent Template					
General Description		A Usual Occupation Duration Entry is a clinical statement about the total quantity of time a person spent in the occupation they held the longest over the course of their life. Start date alone can be insufficient, because a person may have been in and out of the occupation over time. The length of time a person performed a type of work can assist in assessing the extent of potential exposure to a health hazard.			
Class/Mood		Code	Data Type	Value	
ClassCode= "OBS" MoodCode= "EVN"		Code = 74163-7 Display Name = Usual Occupation Duration CodeSystem = 2.16.840.1.113883.6.1 CodeSystemName=LOINC	Observation	Value xsi:type=PQ representing the number of years or months. Units shall be expressed in UCUM.	
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.10 (open)]

A Usual Occupation Duration Entry is a clinical statement about the quantity of time a person spent in the occupation they held the longest over the course of their life.

5445

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

- 5450 3. **SHALL** contain exactly one [1..1] **templateId** such that it
 - a. **SHALL** contain exactly one [1..1]
`@root="1.3.6.1.4.1.19376.1.7.3.1.4.24.10".`
 4. **SHALL** contain at least one [1..*] **id**.
 5. **SHALL** contain exactly one [1..1] **code**.
- 5455 a. **SHALL** be 74163-7 (Usual Occupation Duration) from LOINC.
- 5455 6. **SHALL** contain exactly one [1..1] **statusCode="completed"** Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14).
- 5455 7. **SHALL** contain exactly one [1..1] **value** with @xsi:type="PQ".
 - a. This value **SHALL** contain exactly one [1..1] **@unit**, which **SHALL** include duration-related units from value set UCUM 2.16.840.1.113883.1.11.12839.

Delete the following section 6.3.4.70 Usual Industry Duration

6.3.4.71 Pregnancy Status Review Organizer (1.3.6.1.4.1.19376.1.5.3.1.4.22)

The pregnancy status review organizer collects observations of the responses the patient gave to a set of routine questions regarding potential pregnancy in females of child-bearing-age.

6.3.4.71.1 Specification

```
<organizer classCode='CLUSTER' moodCode='EVN'>
  <templateId root=''/>
  <id root='' extension=''/>
  <code code='' displayName=''
        codeSystem=''
        codeSystemName=''/>
  <statusCode code='completed'>
  <effectiveTime value=''/>
  <!-- For HL7 Version 3 Messages
  <author classCode='AUT'>
    <assignedEntity1 typeCode='ASSIGNED'>
      :
      <assignedEntity1>
    </author>
    -->
  <!-- One or more components -->
  <component typeCode='COMP'>
    <!-- Or a pregnancy status observation -->
    <observation classCode='OBS' moodCode='EVN'>
      <templateId root=''/>
      :
      </observation>
    </component>
  </organizer>
```

6.3.4.71.2 <organizer classCode='CLUSTER' moodCode='EVN'>

The pregnancy status review organizer is a cluster of pregnancy status review observations.

6.3.4.71.3 <templateId root="/">

- 5495 The pregnancy status review organizer shall have the <templateId> element shown above to indicate that it conforms to this specification.

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

The organizer shall have an <id> element.

6.3.4.71.5 <code code='118185001' displayName='Pregnancy Observations'

- 5500 **codeSystem='2.16.840.1.113883.6.96'**
codeSystemName='SNOMED-CT'/>

The organizer shall contain a code describing the observations present. The recommended code is shown above.

6.3.4.71.6 <statusCode code='completed'/>

- 5505 The observations have all been completed.

6.3.4.71.7 <effectiveTime value=' '/>

The effective time element shall be present to indicate the interval of the pregnancy status review.

6.3.4.71.8 <author typeCode='AUT'><assignedEntity1

- 5510 **typeCode='ASSIGNED'>...</assignedEntity1></author>**

For use with HL7 Version 3, pregnancy status review organizers MAY contain an <author> element to represent the person or device.

6.3.4.71.9 <component typeCode='COMP'>

- 5515 The organizer shall have one or more <component> elements that are instances of pregnancy status review observations.

6.3.4.72 Pregnancy Status Review Observation (1.3.6.1.4.1.19376.1.5.3.1.4.22.1)

A pregnancy Status Review observation is a Simple Observation that uses a specific vocabulary to record observations about a patient's current pregnancy status.

6.3.4.72.1 Parent Template

- 5520 The parent of this template is [Simple Observation](#).

6.3.4.72.1.1 Uses

See [Templates using Pregnancy Status Review Observation](#)

6.3.4.72.2 Specification

Pregnancy Status Review Observation Example

```

5525 <observation classCode='OBS' moodCode='EVN'>
      <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.22.1' />
      <templateId root=''/>
      <id root=' ' extension=' ' />
      <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' ' />
      <text><reference value='#xxx' /></text>
      <statusCode code='completed' />
      <effectiveTime value=' ' />
      <repeatNumber value=' ' />
      <value xsi:type=' ' ... />
      <interpretationCode code=' ' codeSystem=' ' codeSystemName=' ' />
      <methodCode code=' ' codeSystem=' ' codeSystemName=' ' />
      <targetSiteCode code=' ' codeSystem=' ' codeSystemName=' ' />
</observation>

```

5540 **6.3.4.72.3 <templateId root='1.3.6.1.4.1.19376.1.5.3.1.4.22.1'>**
<templateId root="/">

These `<templateId>` elements identify this `<observation>` as a pregnancy status review observation, allowing for validation of the content. The `<templateId>` elements shall be recorded as shown above.

5545 **6.3.4.72.4 <code code=' ' displayName=' ' codeSystem=' ' codeSystemName=' '/>**

A pregnancy status observation shall have a code describing what facet of patient's pregnancy status is being recorded.

5550 **6.3.4.72.5 <repeatNumber value=' ' />**

The `<repeatNumber>` element should not be present in a pregnancy status review observation.

6.3.4.72.6 <value xsi:type=' ' ... />

The value of the observation shall be recording using a data type appropriate to the coded observation according to the table above.

5555 **6.3.4.72.7 <interpretationCode code=' ' codeSystem=' ' codeSystemName=' '/>**
<methodCode code=' ' codeSystem=' ' codeSystemName=' '/>
<targetSiteCode code=' ' codeSystem=' ' codeSystemName=' '/>

The `<interpretationCode>`, `<methodCode>`, and `<targetSiteCode>` should not be present in a pregnancy status review observation.

5560 **6.3.4.73 Performer**

The performer template is used to identify the healthcare provider who was the primary performer of an act. The provider name, address, contact information and identifier are provided to ensure that the performer of the act can be contacted in case there are any questions about the act.

5565 <performer typeCode="PRF">
 <templateId root="1.3.6.1.4.1.19376.1.5.3.1.1.24.3.5"/>
 <assignedEntity classCode="ASSIGNED">
 <id root="" extension="" />
 <addr></addr>
 <telecom></telecom>
 <assignedPerson>
 <name></name>
 </assignedPerson>
 <representedOrganization>
 <name></name>
 <addr></addr>
 <telecom></telecom>
 </representedOrganization>
 </assignedEntity>
</performer>

5570

5575

5580

6.3.4.73.1 <performer typeCode="PRF">

The `performer` element identifies a healthcare provider that performed any activity. A performer is distinct from an author, as the performer is the one who does the work, whereas the author is the person who documented or created it.

- 5585 1. This template SHALL be used only in `performer` elements inside any CDA (V3) act.
 2. The `@typeCode` attribute of the `performer` element SHALL use the value **PRF**.

6.3.4.73.2 <templateId root="1.3.6.1.4.1.19376.1.5.3.1.1.24.3.5"/>

The `performer` element asserts conformance to the Performer template.

- 5590 1. The `performer` SHALL contain a `templateId/@root` attribute containing the value **1.3.6.1.4.1.19376.1.5.3.1.1.24.3.5** to assert conformance to this template.

6.3.4.73.3 <assignedEntity classCode="ASSIGNED">

An `assignedEntity` element appears to identify the performer.

- 5595 1. The `performer` SHALL contain only one **[1..1]** `assignedEntity` element.
 2. The `assignedEntity/@classCode` value in the `performer` element SHALL be **ASSIGNED**.

6.3.4.73.4 <id root="" extension="" />

The identifier of the healthcare provider performing the act should be present.

1. The `performer` element SHALL contain at least one **[1..*]** `id` element.

- 5600 2. The `id` element MAY use the `@nullFlavor` attribute when the information is unknown. (clarify that there SHOULD be an `id/@root`).

6.3.4.73.5 <addr></addr>

The mailing address of the healthcare provider performing the act should be present to enable the provider to be contacted.

- 5605 1. The `performer` element SHALL contain at least one **[1..*]** `addr` element.
 2. The `addr` element MAY use `@nullFlavor` if the information is unknown.

6.3.4.73.6 <telecom></telecom>

The provider telephone number should be provided to enable the performer of the reconciliation to be contacted.

- 5610 1. The `performer` element SHALL contain at least one **[1..1]** `telecom` element.
 2. The `telecom` element MAY use `@nullFlavor` to indicate that information is unknown.

6.3.4.73.7 <assignedPerson>

- 5615 1. The `performer` element SHALL contain only one **[1..1]** `assignedPerson` elements further identifying the person.

6.3.4.73.8 <name></name>

The name of the provider performing the act should be provided.

1. The `performer` SHALL contain at least one **[1..*]** `assignedPerson/name` element.
2. The `name` element MAY use `@nullFlavor` to indicate that the information is unknown.

5620 **6.3.4.73.9 <representedOrganization>**

The name and identifier of the organization represented by the performer should be provided.

1. The `performer` SHALL contain only one **[1..1]** `representedOrganization` element.

6.3.4.73.10 <id root='...' extension='...'/>

- 5625 The identifier of the organization represented must appear.

1. The `representedOrganization` element SHALL contain at least one **[1..*]** `representedOrganization/id` element.
2. The `id` element MAY use `@nullFlavor` to indicate that the identifier is unknown.

6.3.4.73.11 <name></name>

5630 The name of the organization represented must appear.

1. The `representedOrganization` element SHALL contain at least one [1..*] `representedOrganization/name` element.
2. The `name` element SHALL NOT use `@nullFlavor` to indicate that information is unknown.

5635 **6.3.4.73.12 <addr></addr>**

The mailing address of the represented organization should be present to allow the organization to be contacted when the performer is not available.

1. The `performer` element shall contain at least one [1..*] `representedOrganization/addr` element.

5640 2. The `addr` element MAY use `@nullFlavor` attribute to indicate that information is unknown.

6.3.4.73.13 <telecom></telecom>

The telephone number of the represented organization should be present to allow the organization to be contacted when the performer is not available.

1. The `performer` element SHALL contain at least one [1..*] `telecom` element.
2. The `telecom` element MAY use `@nullFlavor` to indicate that the information is unknown.

Add the following section 6.3.4.74 Weekly Work Days Observation

5650

6.3.4.74 Weekly Work Days Observation Entry

Table 6.3.4.71-1 Weekly Work Days Observation Entry 1.3.6.1.4.1.19376.1.7.3.1.4.24.7

Template Name	Weekly Work Days Observation Entry		
Template ID	1.3.6.1.4.1.19376.1.7.3.1.4.24.7		
Parent Template			
General Description	The typical days per week that a person spends working.		
Class/Mood	Code	Data Type	Value
ClassCode=“OBS” MoodCode=	Code = 74160-3 Display Name = Weekly Work Days CodeSystem = 2.16.840.1.113883.6.1	Observation	value with <code>@xsi:type="INT"</code>

“EVN”	CodeSystemName=LOINC				
Opt and Card	entryRelationship	Description	Template ID	Specification Document	Vocabulary Constraint

[observation: templateId 1.3.6.1.4.1.19376.1.7.3.1.4.24.7 (open)]

5655 A clinical statement about the typical number of days per week that a person spends performing their duties for work.

- 1. **SHALL** contain exactly one [1..1] @classCode="OBS" (CodeSystem: 2.16.840.1.113883.5.6 HL7ActClass).
- 2. **SHALL** contain exactly one [1..1] @moodCode="EVN" Event (CodeSystem: ActMood 2.16.840.1.113883.5.1001).
- 3. **SHALL** contain exactly one [1..1] templateId such that it
 - a. **SHALL** contain exactly one [1..1] @root="1.3.6.1.4.1.19376.1.7.3.1.4.24.7".
- 4. **SHALL** contain at least one [1..*] id.
- 5. **SHALL** contain exactly one [1..1] code.
 - a. **SHALL** be 74160-3 (Weekly Work Days) from LOINC.
- 6. **SHALL** contain exactly one [1..1] statusCode="completed" Completed (CodeSystem: ActStatus 2.16.840.1.113883.5.14).
- 7. **SHALL** contain exactly one [1..1] value with @xsi:type="INT".
 - a. This value **SHALL** contain exactly one [1..1] @value, which represents the number of days in a week that a person usually works.

Add Section 6.4

6.4 HL7 Version 2.0 Content Modules

5675 This section contains content modules based upon the HL7 Version 2 Standard, and related standards and/or implementation guides.

Add Section 6.5

6.5 PCC Value Sets

5680 This section contains value sets used by Content Modules. The value sets listed here may be used by other domains (e.g., QRPH) in addition to the PCC domain.

Note: Although some tables in this section include a column for “Units”, units may not be applicable to all table entries and the cell will remain blank.

Add Section 6.5.A

5685

6.5.A Antepartum History of Past Illness Value Set**1.3.6.1.4.1.19376.1.5.3.1.1.16.5.1**

Name	Opt	Type	Units	SNOMED CT
Diabetes	R2	CD		73211009
Hypertension	R2	CD		38341003
Heart Disease	R2	CD		56265001
Autoimmune Disorder	R2	CD		85828009
Kidney Disease	R2	CD		90708001
UTI	R2	CD		68566005
Neurologic	R2	CD		118940003
Epilepsy	R2	CD		84757009
Psychiatric	R2	CD		74732009
Depression	R2	CD		41006004
Postpartum Depression	R2	CD		58703003
Hepatitis	R2	CD		128241005
Liver Disease	R2	CD		235856003
Varicosities	R2	CD		276504003
Phlebitis	R2	CD		61599003
Thyroid Dysfunction	R2	CD		14304000
Trauma	R2	CD		417746004
Violence	R2	CD		225818009
History of Blood Transfusion	R2	CD		116859006
D(Rh) Sensitized	R2	CD		3885002
Pulmonary	R2	CD		19829001
Seasonal Allergies	R2	CD		367498001
Drug Allergy	R2	CD		416098002
Latex Allergy	R2	CD		300916003
Food Allergy	R2	CD		414285001
Breast	R2	CD		79604008
Hospitalizations	R2	CD		32485007
Anesthetic Complications	R2	CD		33211000
History of Abnormal Pap	R2	CD		274688009
Uterine Anomaly/DES	R2	CD		37849005
DES Exposure	R2	CD		413340008 of fetus
Infertility	R2	CD		8619003
Artificial Reproductive Therapy (ART) Treatment	R2	CD		63487001
History of Gestational Diabetes	R2	CD		

Name	Opt	Type	Units	SNOMED CT
History of Incompetent Cervix	R2	CD		17382005 Code is for incompetent cervix rather than history of. Given this condition this should be okay.
History of Infant with Intrauterine Growth Restriction	R2	CD		Need Code for history of.
History of Infant with Macrosomia	R2	CD		Need Code for history of.
History of Pregnancy Induced Hypertension	R2	CD		Need code for history of.
History of Placenta Previa/Abruptio	R2	CD		Need Code for history of.
History of Preterm labor	R2	CD		441493008
History of Premature Rupture of Membranes	R2	CD		Need Code for history of.
Previous Cesarean Section	R2	CD		161805006
History of Stillbirth	R2	CD		161743003
History of Neonatal Death	R2	CD		Need code for history of.
History of Postpartum Hemorrhage	R2	CD		161809000

Add Section 6.5.C

6.5.C Antepartum Family History and Genetic Screening Value Set**1.3.6.1.4.1.19376.1.5.3.1.1.16.5.4**

Name	Opt	Type	Units	SNOMED CT	LOINC
Autism	R2	CD		408856003	
Blood Disorders	R2	CD		414022008	
Canavan Disease	R2	CD		80544005	
Chromosomal Disorder Includes any inherited genetic or chromosomal disorders	R2	CD		409709004	
Congenital Heart Defect	R2	CD		13213009	
Cystic Fibrosis	R2	CD		190905008	
Dysmorphism (Birth Defect) Patient or baby's father has a child with birth defects	R2	CD		276720006	
Down Syndrome	R2	CD		41040004	
Familial Dysautonomia	R2	CD		29159009	
Hemophilia	R2	CD		90935002	
Huntington's Chorea	R2	CD		58756001	
Maternal Metabolic Disorder	R2	CD		75934005	
Mental Retardation	R2	CD		91138005	

Name	Opt	Type	Units	SNOMED CT	LOINC
Muscular Dystrophy	R2	CD		73297009	
Neural Tube Defect	R2	CD		253098009	
Recurrent pregnancy loss/stillbirth	R2	CD		102878001	
Sickle Cell Disease	R2	CD		417357006	
Sickle Cell Trait	R2	CD		16402000	
Tay-Sachs	R2	CD		111385000	
Thalassemia	R2	CD		40108008	

5690

Add Section 6.5.D

6.5.D Antepartum Review of Systems Menstrual History Value Set**1.3.6.1.4.1.19376.1.5.3.1.1.16.5.5**

Name	Opt	Type	Units	SNOMED CT	LOINC
Date of Last Menstrual Period	R	TS		21840007	
Menses Monthly	R	BL		364307006	
Prior Menses Date	R	TS		21840007	
Duration of Menstrual Flow	R	PQ	days	364306002	
Frequency of Menstrual Cycles	R	PQ	days	289887006	
On Birth Control Pills at conception	R	BL		10036567	
Menarche	R	PQ		398700009	
hCG+	R	TS		250423000	

Add Section 6.5.E

5695

6.5.E Antepartum History of Infection Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.6

Name	Opt	Type	Units	SNOMED CT	LOINC
Live with someone with TB or exposed to TB	R2	CD		170464005	
History of Genital Herpes	R2	CD		402888002	
Exposed to Genital Herpes	R2	CD		240480009	
Rash since LMP	R2	CD		49882001	
Viral illness since LMP	R2	CD		34014006	
Rash or viral illness since LMP	R2	CD		49882001	
Hepatitis B	R2	CD		235871004	

Name	Opt	Type	Units	SNOMED CT	LOINC
Hepatitis C	R2	CD		235872006	
History of STD	R2	CD		8098009	
History of Gonorrhea	R2	CD		15628003	
History of Chlamydia	R2	CD		312099009	
History of HPV	R2	CD		302812006	
History of HIV	R2	CD		165816005	
History of Syphilis	R2	CD		76272004	

Add Section 6.5.F

6.5.F Antepartum Laboratory Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.7

Lab	LOINC Code	Comments
Antibody Screen (AB)	890-4 Ab Screen	
Blood Type (ABO Group)	883-9 ABO Group	
Rh	10331-7 Rh	
Hepatitis B virus (HBV) surface Antigen (Ag)	5196-1 HBV surface Ag (EIA) 5195-3 HBV surface Ag 5197-9 HBV surface Ag (RIA) 7905-3 HBV surface Ag (Neut)	
Hemoglobin (Hgb)/Hematocrit (Hct)	718-7 Hgb 4544-3 Hct (Automated count) 30350-3 Hgb	
Hemoglobin (Hgb) Electrophoresis	13514-5 Hemoglobin pattern [interpretation] in Blood by Electrophoresis Narrative	Appropriate code appears to be 13514-5
Aneuploidy Screening (Ultrasound)	XX-ASU Aneuploidy Screening (Ultrasound)	XX-ASU: A LOINC profile code will be requested
Pap Test/Human papilloma virus (HPV)	21440-3 HPV I/H Risk DNA Cervix (Probe) 21441-1 HPV Low Risk DNA Cervix (Probe) 10524-7 Cytology Cervix 18500-9 Thin Prep Cervix 19765-7 Cytology Cervix/Vaginal (Nominal) 19766-5 Cytology Cervix/Vaginal (Narrative)	

Lab	LOINC Code	Comments
Rubella Virus (RUBV) Antibody (Ab)	5334-8 RUBV Ab IgG (EIA) 20458-6 RUBV Ab IgG 40667-8 RUBV Ab IgG (EIA) 8014-3 RUBV Ab IgG	
Urine Culture Screen	630-4 Bacteria Urine Culture	
Purified protein derivative (PPD)	1647-7 Purified protein derivative skin test	
Chlamydia	6347-9 Chlamydia Ag 14510-2 Chlamydia trachomatis Ag (Vaginal) 14474-1 Chlamydia trachomatis Ag (Urine) 6349-5 Chlamydia trachomatis (Unspecified specimen)	
Gonorrhea	691-6 Neisseria Gonorrhoeae (genital specimen) 9568-7 Neisseria Gonorrhoeaea Ab	
Chlamydia Trachomatis/ Neisseria Gonorrhoeae	45067-6 Chlamydia Trachomatis Neisseria Gonorrhoeae (Cervix) 45074-2 Chlamydia Trachomatis Neisseria Gonorrhoeae (Urine)	
Ultrasound	35096-7 OB Ultrasound Panel	
Alpha-Feto Protein (Maternal) (Profile)	30525-0 Age 29463-7 Body Weight 18185-9 Gestational Age 20450-3 Alpha-1-Fetoprotein 48803-1 Neural Tube Defect Risk	
Chorionic Villus Sampling (CVS)	33774-1 Karotype	
Amniotic Fluid (Karotype)	33773-3 Karyotype (Amino Fluid)	
Amniotic Fluid (AFP)	41273-4 Alpha-1-Fetoprotein, Amniotic Fluid Semi-Quantitative 1832-5 Alpha-1-Fetoprotein [Multiple of the median] in Amniotic Fluid 29595-6 Alpha-1-Fetoprotein [Mass/volume] in Amniotic Fluid	
Diabetes Screen	1557-8 Fasting Blood Glucose-Venous 14770-2 Fasting Blood Glucose-Capillary	
Glucose Tolerance Test (GTT)	1507-3 Glucose 1HR post 75 g glucose 14995-5 Glucose 2HR post 75 g glucose	

Lab	LOINC Code	Comments
	20437-0 Glucose 3HR post 75 g glucose	
Rapid Plasma Reagin (RPR)	31147-2 Reagin Ab 20508-8 Reagin Ab by RPR	
Venereal Disease Research Laboratory (VDRL)	5292-8 Reagin Ab by VDRL	
Group B Strep	48683-7 Beta Strep Group B (PCR) 11267-2 Strep Group B	
Beta Human Chorionic Gonadotropin (HCG)	21198-7 Beta HCG	
Varicella zoster virus Ab.IgG	15410-4 Varicella zoster virus Ab.IgG (EIA) 17763-4 Varicella zoster virus Ab.IgG (IF)	
Maternal Serum Triple Screen	30525-0 Age, Patient Quantitative 20450-3 Alpha-1-Fetoprotein Multiple of the Median, Serum Quantitative Calculated 20465-1 Choriogonadotropin/Choriogonatotropin, Control Serum Quantitative 20466-9 Estriol/Estriol, Control Serum Quantitative	
Urinalysis (Urine Screen)	20406-5 Glucose 20505-4 Bilirubin 5797-6 Ketones 5811-5 Specific Gravity 5803-2 pH 5804-0 Protein 20405-7 Urobilinogen 20407-3 Nitrite 5794-3 Hemoglobin 5799-2 Leukocyte esterase 5767-9 Appearance 5778-6 Color 9842-6 Casts 5787-7 Epithelial cells 13945-1 Erythrocytes 5769-5 Bacteria	

Lab	LOINC Code	Comments
First Trimester Maternal Serum Screening with Nuchal Translucency	49588-7 First trimester maternal screen with nuchal translucency [interpretation] Narrative	
Thyroid Stimulating Hormone (TSH)	11580-8 Thyrotropin (3rd generation) 3016-3 TSH 5385-0 Thyrotropin Receptor Ab 27975-2 TSH (serum)	
Triiodothyronine (T3)	3051-0 T3 Free 3052-8 T3 Reverse 3054-4 T3 True 3050-2 T3 Resin Uptake	

5700

*Add Section 6.5.G***6.5.G Antepartum Education Value Set 1.3.6.1.4.1.19376.1.5.3.1.1.16.5.8**

Name	Opt	Type	units	SNOMED CT	LOINC
First Trimester					
Risk factors identified by prenatal history	R2	CD		440047008	
Anticipated course of prenatal care	R2	CD		17629007	
Special Diet	R2	CD		171054004	
Nutrition and weight gain counseling	R2	CD		171054004	
Toxoplasmosis precautions (cats/raw meat)	R2	CD		439733009	
Sexual activity	R2	CD		162169002	
Exercise	R2	CD		171056002	
Influenza vaccine	R2	CD		xx-edu-influenza need code closest is vaccine education 171044003	
Smoking/tobacco counseling	R2	CD		171055003	
Environmental/work hazards	R2	CD		385872009	
Travel	R2	CD		439816006	
Alcohol	R2	CD		171057006	
Illicit/recreational drugs	R2	CD		425014005	
Use of any medications	R2	CD		171058001	

Name	Opt	Type	units	SNOMED CT	LOINC
Indications for ultrasound	R2	CD		440227005	
Domestic violence	R2	CD		413457006	
Seatbelt use	R2	CD		440638004	
Childbirth classes/hospital facilities	R2	CD		66961001	
Second Trimester					
Childbirth classes/hospital facilities	R2	CD		66961001	
Signs and symptoms of preterm labor	R2	CD		440669000	
Abnormal Lab Values	R2	CD		410299006	
Influenza vaccine	R2	CD		xx-edu-fluvaccine need code. Closest is vaccine education 171044003	
Selecting a newborn care provider	R2	CD		439908001	
Postpartum family planning	R2	CD		54070000	
Tubal sterilization	R2	CD		243064009	
Third Trimester					
Anesthesia/analgesia plans	R2	CD		243062008	
Intended Facility for Delivery plan				310585007	
Fetal movement monitoring	R2	CD		440309009	
Labor signs	R2	CD		440671000	
VBAC counseling	R2	CD		440073003	
Signs & Symptoms of Pregnancy-induced hypertension	R2	CD		xx-edu-sspreclampsia need to request code	
Postterm counseling	R2	CD		xx-edu-postterm need to request code	
Circumcision	R2	CD		184002001	
Bottle feeding	R2	CD		169644004	
Breast feeding	R2	CD		169643005	
Postpartum depression	R2	CD		439366005	
Newborn education (Newborn screening, jaundice, SIDS, car seat)	R2	CD		75461000	
Family medical leave or disability forms	R2	CD		40791000	
Tubal sterilization consent signed	R2	CD		408835000	

Add section 6.5.H. (Added 2011-09 from QRPH EHCP Profile)

5705 The value subsets provided in this section are used both to constrain the CDA content, and to assert measure logic. These MAY be supported by the Value Set Repository for value set management as defined by the IHE ITI TF Sharing of Value Sets (SVS) Profile.

6.5.H JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set

6.5.H.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.24
Name	name of the value set	JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the Risk Indicators for Hearing Loss associated with hearing loss using LOINC® concepts
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5710

6.5.H.2 JCIH-EHDI Risk Indicators for Hearing Loss (LOINC®) Value Set Value Set

LOINC® 58232-0 Hearing Loss Risk Indicator

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.24
	Vocabulary:	2.16.840.1.113883.6.1
Sequence	LOINC® Code	Description
1	LA137-2	None
2	LA12667-4	Caregiver concern about hearing
3	LA12668-2	Family Hx of hearing loss
4	LA12669-0	NICU stay > 5 days
5	LA12670-8	ECMO
6	LA12671-6	Assisted ventilation
7	LA12672-4	Ototoxic medication use
8	LA12673-2	Exchange transfusion for Hyperbilirubinemia
9	LA12674-0	In utero infection(s)
10	LA12675-7	Craniofacial anomalies
11	LA12681-5	Physical findings of syndromes that include hearing loss
12	LA12676-5	Syndromes associated with hearing loss
13	LA12677-3	Neurodegenerative disorders
14	LA12678-1	Postnatal infections
15	LA12679-9	Head trauma
16	LA6172-6	Chemotherapy

5715 *Add section 6.5.I. (Added 2011-09 from QRPH EHCP Profile)*

6.5.I JCIH-EHDI Risk Indicators for Hearing Loss Codes

6.5.I.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.11
Name	name of the value set	JCIH-EHDI Risk Indicators for Hearing Loss Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the risk indicators for hearing loss associated with hearing loss using SNOMED-CT Finding/Situation concepts
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5720

6.5.I.2 JCIH-EHDI Risk Indicators for Hearing Loss Value Set

SNOMED-CT Risk Indicators for Hearing Loss Value Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.11
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
1	439750006	Family history of hearing loss (situation)
2	441899004	History of therapy with ototoxic medication (situation)
3	276687002	Conjugated hyperbilirubinemia in infancy (disorder)
4	281610001	Neonatal hyperbilirubinemia (disorder)
5	281612009	Neonatal conjugated hyperbilirubinemia (disorder)
6	281611002	Neonatal unconjugated hyperbilirubinemia (disorder)
7	206363004	Intra-amniotic fetal infection (disorder) (Deprecated, replaced by 11618000)
8	206331005	Infections specific to perinatal period (disorder)
9	206005002	Fetus or neonate affected by maternal infection (disorder)
10	80690008	Degenerative disease of the central nervous system (disorder)
11	178280004	Postnatal infection (disorder)
12	312972009	Neonatal extracranial head trauma (disorder)
13	161653008	History of - chemotherapy (situation)
14	11618000	Intra-amniotic infection of fetus (disorder) (Replaces 206363004)

6.5.I.3 Pending Codes for SNOMED-CT Findings/Situation to support Risk Indicators for Hearing Loss

5725

Note that additional specificity for this value set is under way and will result in an update to this value set. Further coded values are sought to represent the following:

None
Caregiver concern about hearing
Craniofacial anomalies
Physical findings of syndromes that include hearing loss
Syndromes associated with hearing loss

5730

Add section 6.5.J. (Added 2011-09 from QRPH EHCP Profile)

6.5.J JCIH-EHDI Risk Indicators for Hearing Loss - Procedures Codes

6.5.J.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.12
Name	name of the value set	JCIH-EHDI Risk Indicators for Hearing Loss - Procedures Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the risk indicators for hearing loss Procedures associated with hearing loss using SNOMED-CT
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.J.2 JCIH-EHDI Risk Indicators for Hearing Loss - Procedures Value

5735

Risk Indicators for Hearing Loss - Procedures Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.12
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
1	266700009	Assisted breathing (procedure)
2	233573008	Extracorporeal membrane oxygenation (procedure)

Add section 6.5.K. (Added 2011-09 from QRPH EHCP Profile)

6.5.K Newborn Hearing Procedure Codes

6.5.K.1 Metadata

5740

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.17
Name	name of the value set	JCIH-EHDI Newborn Hearing Procedure Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the type of newborn hearing procedure identified using SNOMED-CT Procedure codes (includes both screening and other tests and examinations)
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.K.2 JCIH-EHDI Newborn Hearing Procedure Value Set

Newborn Hearing Procedure Value Set:

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.17
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
3	417491009	Neonatal hearing test (procedure)

5745

Add section 6.5.L. (Added 2011-09 from QRPH EHCP Profile)

6.5.L JCIH-EHDI Newborn Hearing Screening Method Codes

6.5.L.1 Metadata

5750

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.4
Name	name of the value set	JCIH-EHDI Newborn Hearing Screening Method Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the type of newborn hearing screening procedure identified using LOINC® answer codes
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.L.2 JCIH-EHDI Newborn Hearing Screening Method Value Set

Newborn Hearing Screening Method Value Set:

LOINC® 54106-0

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.4	
	Vocabulary:	2.16.840.1.113883.6.1	
Sequence	LOINC® Code	Answer Code	Description
1	LA10387-1	AABR	Automated auditory brainstem response
2	LA10388-9	ABR	Auditory brain stem response
3	LA10389-7	OAE	Otoacoustic emissions
4	LA10390-5	DPOAE	Distortion product otoacoustic emissions
5	LA10391-3	TOAE	Transient otoacoustic emissions
6	LA12406-7		Methodology unknown

5755

Add section 6.5.M. (Added 2011-09 from QRPH EHCP Profile)

6.5.M JCIH-EHDI Hearing Screen Right Codes– Right

6.5.M.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.9
Name	name of the value set	JCIH-EHDI Hearing Screen Right Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the right ear EHDI screening using LOINC® in result type
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active

Metadata Element	Description	Mandatory
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5760

6.5.M.2 JCIH-EHDI Hearing Screen Right Value Set

NB hearing scn -R:Result Type

Hearing Screen Right Value Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.9
	Vocabulary:	2.16.840.1.113883.6.1
Sequence	LOINC® Code	Description
1	53109-4	Newborn Hearing Screen Right

5765

Add section 6.5.N. (Added 2011-09 from QRPH EHCP Profile)

6.5.N JCIH-EHDI Hearing Screen Left Codes

6.5.N.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.8
Name	name of the value set	JCIH-EHDI Hearing Screen Left Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the left ear EHDI hearing screening result type using LOINC®
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org

Metadata Element	Description	Mandatory
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5770 **6.5.N.2 JCIH-EHDI Hearing Screen Left Value Set**

Hearing Screen Left Value Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.8
	Vocabulary:	2.16.840.1.113883.6.1
Sequence	LOINC® Code	Description
1	53108-6	Newborn Hearing Screen Left

Add section 6.5.O. (Added 2011-09 from QRPH EHCP Profile)

5775 **6.5.O JCIH-EHDI Reason for no Hearing Loss Diagnosis or Screening Codes(SNOMED)**

6.5.O.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.15
Name	name of the value set	JCIH-EHDI Reason for no Hearing Loss Diagnosis or Screening Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect Reason for no hearing loss diagnosis coded with SNOMED-CT.

Metadata Element	Description	Mandatory
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.O.2 JCIH-EHDI Reason for no Hearing Loss Diagnosis or Screening Value Set

5780

Reason for no Hearing Loss Diagnosis

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.15	
	Vocabulary:	2.16.840.1.113883.6.96	
Sequence	SNOMED Code	Description	EHDI Concept
1	397709008	Patient died (finding)	No screening or diagnosis: Infant died
2	360885002	Change of residence status (finding)	No diagnosis: Moved or gone elsewhere
3	184112005	Patient address unknown (finding)	No diagnosis: Unable to Contact Family
4	184118009	Patient telephone number unknown (finding)	No diagnosis: Unable to Contact Family
5	183638004	Follow-up refused	No screening diagnosis: Parents Declined Services - Follow-up refused
6	183946001	Procedure refused-uncooperative	No diagnosis: Parents Declined Services -Procedure refused - uncooperative
7	413319007	Persistent non-attender	No diagnosis: Unresponsive - Persistent non-attender
8	399307001	Loss to follow-up	No diagnosis: Unknown - Loss to follow-up

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.15	
	Vocabulary:	2.16.840.1.113883.6.96	
Sequence	SNOMED Code	Description	EHDI Concept
9	419984006	Inconclusive (qualifier value)	No diagnosis: Audiologic Diagnosis in Process
10	185332005	Appointment cancelled by patient (finding)	No diagnosis: Audiologic Diagnosis in Process - Rescheduled appointment
11	185333000	Appointment cancelled by doctor (finding)	No diagnosis: Audiologic Diagnosis in Process - Rescheduled appointment
12	281399006	Did not attend	No diagnosis: Audiologic Diagnosis in Process - Did not attend

Add section 6.5.P. (Added 2011-09 from QRPH EHCP Profile)

6.5.P JCIH-EHDI Newborn Hearing Loss Referrals Codes

5785

6.5.P.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.16
Name	name of the value set	JCIH-EHDI Newborn Hearing Loss Referrals Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect EHDI referrals coded with SNOMED-CT and as a response to care plan recommendations (entered on a list of referrals in a medical summary)
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010

Metadata Element	Description	Mandatory
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.P.2 JCIH-EHDI Newborn Hearing Loss Referrals Value Set

EHDI Newborn Hearing Loss Referrals Value Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.16	
	Vocabulary:	2.16.840.1.113883.6.96	
Sequence	SNOMED Code	Description	EHDI Concept
1	306210008	Referral to pediatric diagnostic audiology service (procedure)	Referral to audiologist
2	415271004	Referral to education service (procedure)	Referral to Early Intervention (Part C/non Part C)

5790

<i>Add section 6.5.Q. (Added 2011-09 from QRPH EHCP Profile)</i>
--

6.5.Q JCIH-EHDI Newborn Hearing Loss Reason for no Follow-up – Patient Reason Codes

6.5.Q.1 Metadata

5795

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.7
Name	name of the value set	JCIH-EHDI Newborn Hearing Loss Reason for no Follow-up – Patient Reason Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the reason that no follow-up is conducted in the case of hearing loss using SNOMED-CT reflected in negation of intent to order the referral.

Metadata Element	Description	Mandatory
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.Q2 JCIH-EHDI Newborn Hearing Loss Reason for no Follow-up – Patient Reason Value Set

EHDI Newborn Hearing Loss Reason for no Follow-up Value Set

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.7	
	Vocabulary:	2.16.840.1.113883.6.96	
Sequence	SNOMED Code	Description	EHDI Concept
1	397709008	Patient died (finding)	Incomplete outpatient screen: Infant died
2	360885002	Change of residence status (finding)	Incomplete outpatient screen: Moved or gone elsewhere
3	184112005	Patient address unknown (finding)	Incomplete outpatient screen: Unable to contact family
4	184118009	Patient telephone number unknown (finding)	Incomplete outpatient screen: Unable to contact family
5	183638004	Follow-up refused	Incomplete outpatient screen: Follow-up refused
6	183946001	Procedure refused-uncooperative	Incomplete outpatient screen: Procedure refused-uncooperative
7	413319007	Persistent non-attender	Incomplete outpatient screen: Persistent non-attender
8	399307001	Loss to follow-up	Incomplete outpatient screen: Loss to follow-up

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.7	
	Vocabulary:	2.16.840.1.113883.6.96	
Sequence	SNOMED Code	Description	EHDI Concept
9	185332005	Appointment cancelled by patient (finding)	Incomplete outpatient screen: Rescheduled appointment
10	185333000	Appointment cancelled by doctor (finding)	Incomplete outpatient screen: Rescheduled appointment
11	281399006	Did not attend	Incomplete outpatient screen: Did not attend

5800

Add section 6.5.R. (Added 2011-09 from QRPH EHCP Profile)

6.5.R Joint Commission Medical Reason Codes

6.5.R.1 Metadata

5805

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.33895.1.3.0.75
Name	name of the value set	Joint Commission Medical Reason Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	The Joint Commission value set is used to reflect medical reason why a test was not performed
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.R.2 Joint Commission Medical Reason Value Set

EHDI specifies the re-use of the existing Medical Reason Value Set used by the Joint Commission measures.

5810

	Value Set :	1.3.6.1.4.1.33895.1.3.0.75
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
1	397745006	Medical contraindication (finding)
2	397773008	Surgical contraindication (finding)

5815

Add section 6.5.S. (Added 2011-09 from QRPH EHCP Profile)

6.5.S JCIH-EHDI Inpatient Screening Results not Performed Codes

6.5.S.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.10
Name	name of the value set	JCIH-EHDI Inpatient Screening Results not Performed Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the right ear EHDI results reported using LOINC® answer lists
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5820 **6.5.S.2 JCIH-EHDI Inpatient Screening Results not Performed Value Set**

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1. 1.15.2.10		
	Vocabulary:	2.16.840.1.113883.6.1		
Sequence	LOINC® Code	Description	Global ID	Global ID Code System
1	LA12408-3	Attempted, but unsuccessful - technical fail	103709008	SN
2	LA7304-4	Not performed	262008008	SN
3	LA12409-1	Not performed, medical exclusion - not indicated	410534003	SN

Add section 6.5.T. (Added 2011-09 from QRPH EHCP Profile)

6.5.T JCIH-EHDI Evidence of Hearing Screening Performed Codes5825 **6.5.T.1 Metadata**

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.18
Name	name of the value set	JCIH-EHDI Evidence of Hearing Screening Performed Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect Evidence of Hearing Screening Performed through the result values of pass-Left, pass-Right, or Refer. This excludes unsuccessful results.
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from LOINC®
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://loinc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010

Metadata Element	Description	Mandatory
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.T.2 JCIH-EHDI Evidence of Hearing Screening Performed Value Set

Evidence of Hearing Screening Performed Value Set

		Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.18		
		Vocabulary:	2.16.840.1.113883.6.1		
Sequence	LOINC® Code	Answer Code	Description	Global ID	Global ID Code System
1	LA10392-1	164059009	Pass		
2	LA10393-9	183924009	Refer		

5830

Add section 6.5.U. (Added 2011-09 from QRPH EHCP Profile)

6.5.U JCIH-EHDI Procedure Declined Value Set Codes

6.5.U.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.20
Name	name of the value set	JCIH-EHDI Procedure Declined Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect that the hearing screening procedure was not performed due to the patient/parent declining the procedure

Metadata Element	Description	Mandatory
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

5835

6.5.U.2 JCIH-EHDI Procedure Declined Value Set Value Set

JCIH-EHDI Procedure Declined Value Set:

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.20
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
1	183949008	Assessment examination refused (situation)
2	183945002	Procedure refused - religion (situation)
3	183948000	Refused procedure - parent's wish (situation)
4	397709008	Patient died (finding)

Add section 6.5.V. (Added 2011-09 from QRPH EHCP Profile)

5840 **6.5.V JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set Codes**

6.5.V.1 Metadata

Metadata Element	Description	Mandatory
Identifier	unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.23

Metadata Element	Description	Mandatory
Name	name of the value set	JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set
Source	source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect abnormal results from the hearing screening procedure
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2010
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2010
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group MAY also have an OID assigned.	IHE EHDI

6.5.V.2 JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set

JCIH-EHDI Newborn Hearing Screening Abnormal Results Value Set:

	Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.15.2.23
	Vocabulary:	2.16.840.1.113883.6.96
Sequence	SNOMED-CT Code	Description
1	313203003	Hearing test abnormal (finding)
2	308409008	Child hearing screening failure (finding)
3	185577006	Child hearing screening first failure (finding)
4	185579009	Child hearing screening second failure (finding)
5	185580007	Child hearing screening failure referred to specialist (finding)

<i>Add section 6.5.W. (Added 2011-09 from QRPH PRPH-Ca Profile)</i>

6.5.W Primary Site Value Set

LOINC = 22035-0	
Code System: ICD-O-3 2.16.840.1.113883.6.43.1	
Code	Meaning
	A code from ICD-O-3 (Topography Section)

*Add section 6.5.X (Added 2011-09 from QRPH PRPH-Ca Profile)*5850 **6.5.X Histologic Type Value Set**

LOINC = 31205-8	
Code System: ICD-O-3 2.16.840.1.113883.6.43.1	
Code	Meaning
	An ICD-O-3 code (Morphology Section)

*Add section 6.5.Y (Added 2011-09 from QRPH PRPH-Ca Profile)***6.5.Y Derived AJCC Descriptor (T,N,M) Value Set**

LOINC = 21908-9	
Code System: 2.16.840.1.113883.15.6	
Code	Meaning
c	clinical
p	pathologic
a	Autopsy classification
yc or yp	Posttherapy classification “y” prefix to utilize with “c” or “p” for denoting extent of cancer after neoadjuvant or primary systemic and/or radiation therapy
r	Retreatment Classification

5855 *Add section 6.5.Z (Added 2011-09 from QRPH PRPH-Ca Profile)***6.5.Z TNM Edition Value Set**

LOINC = 21917-0	
Code System: 2.16.840.1.113883.15.6	
Code	Meaning
5	AJCC Staging Manual, 5 th Edition
6	AJCC Staging Manual, 6 th Edition
7	AJCC Staging Manual, 7 th Edition

<i>Add section 6.5.AA (Added 2011-09 from QRPH PRPH-Ca Profile)</i>

5860

6.5.AA TNM Stage Group Value Set

Note: The AJCC Staging Manual TNM system is propriety and its definitions cannot be included in other documents without permission.

LOINC = 21908-9	
Code System: TNM 5. Edition: 2.16.840.1.113883.15.8 - tnm5 TNM 6. Edition: 2.16.840.1.113883.15.7 - tnm6 TNM 7. Edition: 2.16.840.1.113883.15.6 - tnm7	
Code	Description: Site specific descriptions prevent listing of text equivalents.
0	Site specific descriptions prevent listing of text equivalents.
0a	“
0is	“
I	“
IA	“
IA1	“
IA2	“
IB	“
IB1	“
IB2	“
IC	“
II	“
IIA	“
IIA1	“
IIA2	“
IIB	“
IIC	“
III	“
IIIA	“
IIIB	“
IIIC	“
IS	“
IV	“
IVA	“
IVB	“
IVC	“

<i>Add section 6.5.BB (Added 2011-09 from QRPH PRPH-Ca Profile)</i>

5865

6.5.BB TNM Stage Descriptor Value Set

Note: The AJCC Staging Manual TNM system is propriety and its definitions cannot be included in other documents without permission.

LOINC = 21909-7	
Code System: TNM 5. Edition: 2.16.840.1.113883.15.8 - tnm5 TNM 6. Edition: 2.16.840.1.113883.15.7 - tnm6 TNM 7. Edition: 2.16.840.1.113883.15.6 - tnm7	
Code	Meaning
0	None
1	E (Extranodal, lymphomas only)
2	S (Spleen, lymphomas only)
3	M (Multiple primary tumors in a single site)
4	Y (Classification during or after initial multimodality therapy)—pathologic staging only
5	E & S (Extranodal and spleen, lymphomas only)
6	M & Y (Multiple primary tumors and initial multimodality therapy)

5870

Add section 6.5.CC (Added 2011-09 from QRPH PRPH-Ca Profile)
--

6.5.CC TNM Tumor Value Set

Note: The AJCC Staging Manual TNM system is propriety and its definitions cannot be included in other documents without permission.

LOINC = 21905-5	
Code System: TNM 5. Edition: 2.16.840.1.113883.15.8 - tnm5 TNM 6. Edition: 2.16.840.1.113883.15.7 - tnm6 TNM 7. Edition: 2.16.840.1.113883.15.6 - tnm7	
Code	Description: Site specific descriptions prevent listing of text equivalents.
Ta	Site specific descriptions prevent listing of text equivalents.
Tis	“
T0	“
T1	“
T1mic	“
T1a	“
T1a1	“
T1a2	“
T1b	“
T1b1	“
T1b2	“
T1c	“

T1d	“
T2	“
T2a	“
T2a1	“
T2a2	“
T2b	“
T2c	“
T2d	“
T3	“
T3a	“
T3b	“
T3c	“
T3d	“
T4	“
T4a	“
T4b	“
T4c	“
T4d	“
T4e	“
Tx	“

5875

Add section 6.5.DD (Added 2011-09 from QRPH PRPH-Ca Profile)

6.5.DD TNM Node Value Set

Note: The AJCC Staging Manual TNM system is propriety and its definitions cannot be included in other documents without permission.

LOINC = 21906-3	
Code System: TNM 5. Edition: 2.16.840.1.113883.15.8 - tnm5	
TNM 6. Edition: 2.16.840.1.113883.15.7 - tnm6	
TNM 7. Edition: 2.16.840.1.113883.15.6 - tnm7	
Code	Description: Site specific descriptions prevent listing of text equivalents.
N0	Site specific descriptions prevent listing of text equivalents.
N1	“
N1mi	“
N1a	“
N1b	“
N1b1	“
N1b2	“
N1b3	“
N1b4	“
N1c	“

N2	“
N2a	“
N2b	“
N2c	“
N3	“
N3a	“
N3b	“
N3c	“
N	“

5880

Add section 6.5.EE (Added 2011-09 from QRPH PRPH-Ca Profile)

6.5.EE TNM Metastasis Value Set

Note: The AJCC Staging Manual TNM system is propriety and its definitions cannot be included in other documents without permission.

LOINC = 21907-1	
Code System:	TNM 5. Edition: 2.16.840.1.113883.15.8 - tnm5 TNM 6. Edition: 2.16.840.1.113883.15.7 - tnm6 TNM 7. Edition: 2.16.840.1.113883.15.6 - tnm7
Code	Description: Site specific descriptions prevent listing of text equivalents.
M0	Site specific descriptions prevent listing of text equivalents.
M1	“
M1a	“
M1b	“
M1c	“
M1d	“
M1e	“
Mx	“

5885

Add section 6.5.FF (Added 2013-09 for the QRPH VRDR supplement)

6.5.FF QRPH VRDR Autopsy Procedure Performed Codes

6.5.FF.1 Metadata

Autopsy Procedure Performed Value Set Metadata Shall contain the following content:

5890

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.1

Metadata Element	Description	Mandatory
Name	This is the name of the value set	VRDR Autopsy Procedure Performed Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To reflect that there was an Autopsy Procedure Performed
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	4/3/2013
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

6.5.FF.2 VRDR Autopsy Procedure Performed Value Set

VRDR Autopsy Procedure Performed Value Set will use the SNOMED-CT code system to identify its contents. Codes that are used within the scope of this profile are listed below:

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.1
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
9427006	Autopsy review
16521010	Autopsy review
16522015	Autopsy review, NOS
68184000	Autopsy review, consultation and report
60864000	Autopsy review for conference (procedure)
86693001	Autopsy review for teaching (procedure)
5785009	Forensic autopsy (procedure)
61501008	Forensic autopsy, extensive (procedure)

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.1
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
29240004	Autopsy examination (procedure)
48926013	Autopsy examination
48930011	Autopsy
48927016	Autopsy examination, NOS
48928014	Autopsy, NOS
41770000	Autopsy, gross and microscopic examination (procedure)
56417000	Autopsy, gross and microscopic examination with brain (procedure)
41554000	Autopsy, gross and microscopic examination with brain and spinal cord (procedure)
74348008	Autopsy, gross and microscopic examination, limited (procedure)
57438004	Autopsy, gross and microscopic examination, regional (procedure)
16361008	Autopsy, gross and microscopic examination, stillborn or newborn (procedure)
4447001	Autopsy, gross and microscopic examination, stillborn or newborn without CNS (procedure)
82823006	Autopsy, gross examination with brain (procedure)
47197006	Autopsy, gross examination with brain and spinal cord (procedure)
72598009	Autopsy, gross examination, limited (procedure)
47847005	Autopsy, gross examination, limited, regional (procedure)
50333006	Autopsy, gross examination, macerated stillborn (procedure)
35459000	Autopsy, gross examination, stillborn or newborn (procedure)
26762004	Autopsy, gross examination, teaching, complete (procedure)
22677004	Autopsy, gross examination, teaching, limited (procedure)
5785009	Forensic autopsy (procedure)
430339001	Pediatric autopsy (procedure)
90864005	Special autopsy procedure, explain by report (procedure)

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.1
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
43939005	Autopsy service by diener (procedure)
71604005	Forensic autopsy, coroner's call (procedure)
108259003	Autopsy pathology procedure AND/OR service (procedure)
59543001	Autopsy, clerical procedure (procedure)
29915004	Autopsy, clerical with coding procedure (procedure)
3133002	Patient discharge, deceased, autopsy (procedure)

5895

Add section 6.5.GG (Added 2013-09 form the QRPH VRDR supplement.)

6.5.GG QRPH VRDR Autopsy Not Performed Codes

6.5.GG.1 Metadata

Autopsy Not Performed Value Set Metadata Shall contain the following content:

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.2
Name	This is the name of the value set	VRDR Autopsy Not Performed Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To reflect that there was an Autopsy was not performed
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed/snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	4/3/2013

Metadata Element	Description	Mandatory
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

5900

6.5.GG.2 VRDR Autopsy Not Performed Value Set

VRDR Autopsy Not Performed Value Set will use the SNOMED-CT code system to identify its contents. Codes that are used within the scope of this profile are listed below:

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.2
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
44551000009109	Autopsy not performed (finding)
76231000009111	No post performed
76241000009117	Post mortem examination not performed
76221000009114	Autopsy not performed
408775001	Consent for postmortem declined (finding)
2470636019	Consent for postmortem declined
2477187017	Consent for autopsy declined
79779006	Patient discharge, deceased, no autopsy (procedure)

5905

Add section 6.5.HH (Added 2013-09 from the QRPH VRDR supplement.)

6.5.HH VRDR Discharge Death Codes

6.5.HH.1 Metadata

Discharge Death Value Set Metadata Shall contain the following content:

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3 .1.1.23.8.3
Name	This is the name of the value set	VRDR Discharge Death Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect Discharge disposition of death

Metadata Element	Description	Mandatory
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from UB04
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	www.nubc.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2013
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

5910 **6.5.HH.2 VRDR Discharge DeathValue Set**

Discharge Death Value Set will use the UB-04/NUBC code system to identify its contents. Codes that are used within the scope of this profile are listed below:

Section Template :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.3
Vocabulary:	UB04 OID
UB-04/NUBC Code	Description
20	Expired

5915 Add section 6.5.II (Added 2013-09 form the QRPH VRDR supplement.)

6.5.II VRDR Death Location Type Codes

6.5.II.1 Metadata

Death Location Type Value Set Metadata Shall contain the following content:

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.4

Metadata Element	Description	Mandatory
Name	This is the name of the value set	Death Location Type Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To Reflect the location where the decedent died
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from the HL7 VRDR CDA Death Location Types
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	www.HL7.org
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	8/1/2013
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

5920 6.5.II.2 VRDR Death Location Type Value Set

Death Location Type Value Set will use the HL7 Death Location Type code system to identify its contents. Codes that are used within the scope of this profile are listed below:

Value Set:	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.4
Vocabulary:	2.16.840.1.113883
HL7 Death Location Type Code	HL7DeathLocationType
H-IN	Hospital Inpatient
H-ER/ OP	Hospital Emergency Department or Outpatient
H-DOA	Hospital Dead on Arrival
NH	Nursing Home

RES	Residence
OTH	Other

5925

Add section 6.5.JJ (Added 2013-09 form the QRPH VRDR supplement.)

6.5.JJ VRDR Death Certification Procedure Codes

6.5.JJ.1 Metadata

Death Certification Procedure Value Set Metadata Shall contain the following content:

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3 .1.1.23.8.6
Name	This is the name of the value set	VRDR Death Certification Procedure Performed Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To reflect that there was a Death Certification Procedure Performed
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	4/3/2013
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

5930 **6.5.JJ.2 VRDR Death Certification Procedure Performed Value Set**

Death Certification Procedure Performed Value Set will use the HL7 Transportation Relationship Type code system to identify its contents. Codes that are used within the scope of this profile are listed below:

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.6
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
308646001	Death certification (procedure)

5935

Add section 6.5.KK (Added 2013-09 form the QRPH VRDR supplement.)

6.5.KK VRDR Death Pronouncement Procedure Codes

6.5.KK.1 Metadata

5940 Death Pronouncement Procedure Value Set Metadata Shall contain the following content:

Metadata Element	Description	Mandatory
Identifier	This is the unique identifier of the value set	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.7
Name	This is the name of the value set	VRDR Death Pronouncement Procedure Performed Value Set
Source	This is the source of the value set, identifying the originator or publisher of the information	IHE Quality Research and Public Health Domain
Purpose	Brief description about the general purpose of the value set	To reflect that there was a Death Pronouncement Procedure Performed
Definition	A text definition describing how concepts in the value set were selected	Extensional definition: The value set was constructed by enumerating the codes from SNOMED-CT
Source URI	Most sources also have a URL or document URI that provides further details regarding the value set.	http://www.nlm.nih.gov/research/umls/Snomed_main.html
Version	A string identifying the specific version of the value set.	Version 1.0

Metadata Element	Description	Mandatory
Status	Active (Current) or Inactive	Active
Effective Date	The date when the value set is expected to be effective	8/1/2013
Expiration Date	The date when the value set is no longer expected to be used	N/A
Creation Date	The date of creation of the value set	4/3/2013
Revision Date	The date of revision of the value set	N/A
Groups	The identifiers of the groups that include this value set. A group may also have an OID assigned.	IHE VRDR

6.5.KK.2 VRDR Death Pronouncement Procedure Performed Value Set

Death Pronouncement Procedure Performed Value Set will use the HL7 Transportation Relationship Type code system to identify its contents. Codes that are used within the scope of 5945 this profile are listed below:

Value Set :	1.3.6.1.4.1.19376.1.7.3.1.1.23.8.7
Vocabulary:	2.16.840.1.113883.6.96
SNOMED-CT Code	SNOMED-CT Description
428413005	Death verification (procedure)

Add the following section 6.6-1 Concept Domains

6.6 Concept Domains

5950 This section describes concept domains used by this technical framework. A concept domain describes the purpose of a coding system in an implementation independent way. National extensions can declare bindings from a concept domain to a specific value set.

Table 6.6-1: Concept Domains

UV Concept Domain	Concept Domain Description
CD_EmploymentStatus	The employment status concept domain defines an individual's economic relationship to an occupation/industry. As described by LOINC®: Generally, employment status refers to whether or not a person is currently employed for wages or doing some other unpaid activity, such as volunteering, homemaking, or participating in educational instruction as a student. In a healthcare setting, employment status may be used to determine appropriate probing questions for occupational exposures and occupational history. For example, a person who is currently not employed for wages may be prompted to provide information about previously held occupations ⁵ .
CD_WorkSchedule	The Work Schedule Concept Domain describes an individual's typical arrangement of working hours for an occupation. As described by LOINC®: For example, work schedule may capture that an individual typically works a regular day shift, evening shift, or night shift. It can also specify if an individual has an irregular schedule such as a rotating shift, split shift, etc. In healthcare settings, knowledge of a patient's typical work schedule may assist in diagnosis of healthcare issues related to irregular work hours or sleep patterns. It may also assist in determining appropriate treatment and prevention plans that will coordinate with the patient's work schedule ⁵ .
CD_OccupationCode	The Occupation Code Concept Domain contains a set of codes that describe a set of activities or tasks that individuals are paid to perform or, if unpaid, define a person's contribution to a household/family business/community ⁶ .
CD_IndustryCode	The Industry Code Concept Domain contains a set of codes that describe an economic/business sector comprised of businesses/ enterprises concerned with the output of a specified category of products or services (e.g., the construction industry or the agriculture industry) ⁶ .

5955

¹ This material contains content from LOINC® (<http://loinc.org>). The LOINC table, LOINC codes, and LOINC panels and forms file are copyright © 1995-2013, Regenstrief Institute, Inc. and the Logical Observation Identifiers Names and Codes (LOINC) Committee and available at no cost under the license at <http://loinc.org/terms-of-use>.

² Source: CDC Census 2010