

Unified Study Definitions Model Implementation Guide (USDM-IG)

Version 4.0 (Draft)

Prepared by the **DDF Team**

Notes to Readers

- This is the draft version of the Unified Study Definitions Model Implementation Guide (intended to be USDMIG v4.0).
- This version has been created using a simple print from the USDMIG Wiki version and not the full copy edited version. This copy editing step will take place before public review and publication.
- Note that the Data Dictionary section contains a table that is truncated. Please refer to the the <u>Wiki</u> <u>version</u> to see the full table.

Revision History

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1 Introduction

CDISC, in collaboration with TransCelerate Biopharma and Accenture as a part of <u>TransCelerate's Digital Data Flow (DDF) Project</u>, have developed a Study Definition Reference Architecture called the Unified Study Definitions Model (USDM).

The aim of TransCelerate's DDF initiative is to optimize study start-up (SSU) processes and automate system configuration and readiness. The current state typically involves disconnected study design services and assets and transcription or re-entry of the same information into many systems across sponsors, contract research organizations, and systems vendors. This inefficiency results in systems configuration falling onto the critical path for SSU and adds risks for transcription errors and unnecessary delays.

Ideally, a solution would enable interoperability across multiple systems in a clinical study, improve efficiency and data quality, and reduce cycle times. That solution should capture protocol elements and present them in standardized formats to enable automated configuration of downstream systems and efficient consumption of protocol information across the study ecosystem.

The challenge is that SSU system configuration workflow and asset creation is currently not automated, which makes it inefficient and increases the risk of error. Current workflows also include a number of redundant, manual activities. Sponsors are not able to utilize resources efficiently due to the siloed, document-based environment. Additional information can be found on the <u>TransCelerate Digital Data Flow Solutions</u> web page.

The collaborative effort between TransCelerate and CDISC has enabled the development of the USDM reference architecture in conjunction with development of a Study Definitions Repository (a reference implementation of the USDM architecture). For more information on the SDR, visit the <u>TransCelerate DDF GitHub site</u> and the <u>SDR Github site</u>.

1.1 Purpose

The USDM Implementation Guide (USDM-IG) is intended for companies and individuals involved in the set-up of clinical studies—sponsors or stakeholders involved in upstream (protocol and content authoring tools)—and downstream consumers of system (e.g., electronic data capture (EDC), clinical trial management, trial master file) and document (e.g., protocol, clinical study reports, statistical analysis plans) standardized digitized study definitions.

This document provides users with sufficient information to understand the USDM and also its potential implementations with the study design process by showing examples of the types of study definition information that can be represented in the USDM.

1.2 Organization of this Document

This document is divided into the following sections:

- Section 1, Introduction, provides an overall introduction to the purpose and goals of the USDM-IG.
- Section 2, <u>Fundamentals of the USDM</u>, provides a boundary of the scope of this version of the USDM and what use cases this version is intended to support.
- Section 3, <u>Relationship to Other Standards and Formats</u>, describes at a high level how the USDM relates to other standards (both CDISC and non-CDISC) and to the TransCelerate Common Protocol Template.
- Section 4, <u>USDM Features</u>, provides an overview of enhancements that support increased trial complexity.
- Section 5, <u>USDM Data Dictionary</u>, illustrates the types of information that can be represented using the USDM, and includes various study designs ranging in complexity.
- Section 6, <u>USDM API</u>, provides information on the USDM application programming interface.
- Section 7, <u>Mapping to Other Standards and Formats</u>, describes the alignment between the USDM and SDTM Trial Design domains and controlled terminology elements, and provides definitions for protocol registration data elements submitted to ClinicalTrials.gov.
- Appendices provide additional background material and describe other supplemental material relevant to the USDM.

Examples of use of the model in JSON, .PNG, and .XLS format as well as other information can be found here.

1.3 How to Read this Document

- First, become familiar with the DDF project; see the <u>TransCelerate DDF Project web page</u> and <u>CDISC DDF</u> resources. If new to DDF, visit the TranCelerate <u>YouTube channel</u>, which includes several videos describing DDF.
- 2. Read this guide all the way through (without skipping any sections) at least once.
- 3. Finally, revisit any sections of particular interest.

2 Fundamentals of the USDM

The USDM comprises 4 parts, which are official CDISC standards:

- 1. Unified Study Definitions Model (USDM) class diagram represented as a unified modeling language (UML) class diagram
- 2. Application programming interface (API) specification
- 3. CDISC Controlled Terminology
- 4. Unified Study Definitions Model Implementation Guide (USDM-IG)

3 USDM v1.0

USDM v1.0 (released August 2022) provided a base model of structured study design.

Please note that USDM v1.0 did not have a corresponding implementation guide. The USDM-IG was initially developed for USDM v2.0 and further updated for USDM v3.0.

4 USDM v2.0

Building on the USDM v1.0 foundation, USDM v2.0 (released June 2023) was developed to satisfy an agreed set of use cases based around

- updates to the USDM that enable greater population of SSU elements and represent structured study design information for more complex trials,
- updates to the USDM that support EDC automation, and
- updates to the USDM that demonstrate population of the TransCelerate Common Protocol Template (CPT).

4.1 Support for More Complex Trials

The first version of the USDM provided a model for simple study designs. Version 2.0 implemented additional elements that allow for representation of more complex study designs in USDM. Section 4, <u>USDM Features</u>, provides an overview of enhancements that support increased trial complexity. One main area of development has been the implementation of study timing (see <u>Section 4.14</u>) within the model, allowing for complex timing and visit structures to be represented.

4.2 Enabling EDC Automation

In order to support EDC automation, the CDISC <u>Biomedical Concepts model</u> was adapted and included as a submodel in the USDM. The addition of biomedical concepts to the model adds a machine-readable "data" layer to the study design. This data layer can be used in a variety of ways to inform about what data relates to particular assessments within a study design. This biomedical concepts model not only assists in informing an EDC system as to the individual data items required for an assessment (e.g., automating identification of a form in an EDC library with the same/similar set of biomedical concepts) but also provide basic information required to build a new form should there be no EDC library, or no form that matches.

Implementation of the biomedical concepts model in the USDM provides a machine-readable data specification that can support other data-source use cases such as digital health technologies, electronic patient-reported outcomes (ePROs), and electronically supplied data (e.g., central lab, central ECG data).

4.3 Populating protocol standards

In Version 2.0, additional elements were added to the model as a proof-of-viability (POV) exercise, demonstrating that structured study design information could be moved from an upstream study design application into USDM format and then used to populate the TransCelerate CPT. Additional information on the USDM elements used for this POV can be found in Section 7.3, <u>Use of USDM for Populating Protocol Content</u>. Note that only a selected set of CPT elements is included for the POV.

5 USDM v3.0

USDM v3.0 development topics included:

- Ability to represent the draft ICH Clinical electronic Structured Harmonised Protocol (CeSHarP) developed by the ICH M11 group in USDM
- Add elements to expand population of SDTM trial design datasets
- Identify elements within USDM that can assist in population of trial planning elements for clinical trial registration in trial registries
- Addition of elements and model amendments required to represent structured study design information for more complex studies, including complex cohort trial designs
- Model enhancements to support use of the USDM and ensure consistency within the model

5.1 Representation of ICH M11 CeSHarP in USDM

Working closely with ICH, USDM v3.0 has been aligned to cover the breadth of sections found in the ICH M11 CeSHarP template. This will allow a USDM study design to be represented in the ICH CeSHarP template. **Note:** At the time of publication of USDM v3.0, ICH CeSHarP was still in the development phase. In future phases of USDM development, CDISC will continue to collaborate with the ICH team in order to ensure that USDM remains aligned with the ICH M11 CeSHarP template.

5.2 SDTM Trial Design Population

During development of USDM v2.0, elements within the USDM were identified that would allow data from a USDM compliant system to be used to populate SDTM Trial Design datasets related to trial planning. This was expanded during USDM v3.0 development to include additional elements that can be used for SDTM Trial Design population. Additional information can be found in Section 7.1, Creation of SDTM Trial Design Domains.

5.3 Clinical Trial Registry Population

Working alongside clinical trial registry subject-matter experts (SMEs), an evaluation was performed to determine how USDM can be utilized to assist in the population of elements required for clinical trial registries. In Version 3.0, this was restricted to ClinicalTrials.gov. Additional information can be found in Section 7.2, <u>Informing ClinicalTrials.gov Registry</u>

5.4 Support for More Complex Trials

An evaluation was performed to determine model changes that could support more complex cohort trials designs. This resulted in new USDM classes being developed (i.e., Population Definitions, Study Cohort, Characteristic) to support these types of studies. Additional information can be found in Section 4.19, Populations, Cohorts, and Eligibility Criteria.

5.5 Model Enhancements

Version 3.0 includes model enhancements to support use of the USDM and ensure consistency within the model, such as updating the UML to make it a more logical model, removing the API implementation elements and links, and making naming more consistent between classes. Additional information can be found in Section 4.2, Principles, Section 4.3, Naming Conventions, Section 4.4, Internal Identifiers Within the Model, and Section 4.5, Controlled Terminology.

6 Relationship to Other Standards and Formats

The USDM covers a wide range of concepts related to study design that also appear in other published standards such as trial registry standards (<u>EudraCT</u>, <u>ClinicalTrials.gov</u>), <u>HL7 FHIR</u> standards, and <u>ICH</u> guidance documents. As part of the development process, these standards were used as input in order to try to ensure harmonization with these standards, where possible.

6.1 Relationship to Other CDISC Standards

The USDM development process relies on published CDISC standards and other products that serve as references for modeling and naming conventions. To the extent possible, an effort has been made to align or be compatible with these sources where the content was determined to be conceptually identical or closely related to those being developed for the USDM.

6.1.1 BRIDG

The Biomedical Research Integrated Domain Group (BRIDG) is a CDISC, <u>HL7</u>, and <u>ISO</u> "standard for biomedical research concepts designed to support computable semantic interoperability."[1] BRIDG can be used for various purposes: as a reference model, a data integration/mapping solution, an exchange format, an ontology, or to create a BRIDG-based database. The use of BRIDG helps support the meaningful exchange of data between software systems and databases.

When BRIDG is used as a reference model to create or add new content to a standard, it can help ensure that relationships between and among biomedical research concepts represented using the standard are consistently modeled.

6.1.2 PRM

The <u>Protocol Representation Model</u> (PRM) provides a standard for planning and designing a research protocol with focus on study characteristics such as study design; eligibility criteria; and requirements from <u>ClinicalTrials.gov</u>, <u>World Health Organization</u> (WHO) registries, and <u>EudraCT</u> registries. The PRM assists in automating CRF creation and EHR configuration to support clinical research and data sharing.

Note: The PRM was released in 2012 and includes some overlap with the USDM. It is anticipated that the USDM will develop to be more content rich and implementable as a model and will therefore supersede the PRM.

6.1.3 SDTM and SDTMIG

The Study Data Tabulation Model (SDTM) provides a standard for organizing and formatting data to streamline processes in collection, management, analysis, and reporting. Implementing SDTM supports data aggregation and warehousing, fosters mining and reuse, facilitates sharing, helps perform due diligence and other important data review activities, and improves the regulatory review and approval process. The SDTM provides a standard model for organizing and formatting data for human and animal studies; the SDTM Implementation Guide (SDTMIG) is intended to guide the organization, structure, and format of standard clinical trial tabulation datasets. The SDTMIG was developed to support data submitted to a regulatory authority, such as the US Food and Drug Administration (FDA), but is not restricted to use in regulated submissions. The SDTM is one of the required standards that sponsors must use, as specified in the FDA's Data Standards Catalog, [2] for New Drug Applications (NDAs), Abbreviated New Drug Applications (ANDAs), and certain Biologics License Applications (BLANDAs). The SDTMIG includes a section related to Trial Design Model datasets. Section 9.1 (Annex IIIa and Annex IIIb) of the ICH Guideline for Industry: Structure and Content of Clinical Study Reports[3] calls for a brief, clear description of the overall plan and design of the study, and supplies examples of charts and diagrams for this purpose. Each annex corresponds to an example trial and provides a diagram describing the study design and a table showing the schedule of assessments. The Trial Design Model provides a standardized way to describe aspects of the planned conduct of a clinical trial shown in the study design diagrams of these examples. Standard Trial Design datasets allow reviewers to

- clearly and quickly grasp the design of a clinical trial,
- compare the designs of different trials,
- search a data warehouse for clinical trials with certain features, and
- compare planned and actual treatments and visits for subjects in a clinical trial.

Modeling a clinical trial in this standardized way requires the explicit statement of certain decision rules that may not be addressed or may be vague or ambiguous in the usual prose protocol document. Prospective modeling of the design of a clinical trial should lead to a clearer, better protocol. Retrospective modeling of the design of a clinical trial should ensure a clear description of how the trial protocol was interpreted by the sponsor.

Automated creation of SDTM Trial Design datasets is possible using data structured in USDM v3.0 format as detailed in Section 7.1, Creation of SDTM Trial Design Domains.

6.1.4 Controlled Terminology

CDISC, in collaboration with the National Cancer Institute's (NCI) Enterprise Vocabulary Services (EVS), supports the controlled terminology (CT) needs of the CDISC standards. Controlled terminology is the set of codelists, definitions, and valid values used with CDISC model elements. Within CDISC there are many volunteer teams that evaluate and manage CDISC CT. For example, the Protocol Entities Terminology Team develops and publishes the semantics for concepts found in clinical research protocols; the CDISC Glossary Team harmonizes the semantics and definitions for concepts commonly found in CDISC standards documents. The DDF terminology subset of CDISC CT is one of the main deliverables supporting the USDM, and development of CDISC CT for the USDM has been harmonized with existing, published CDISC CT (including SDTM, Protocol, and CDISC Glossary) in order to ensure maximum reuse of terms and definitions. Any new CT that has been developed for the USDM has undergone review from the Protocol Entities and CDISC Glossary Teams. USDM-related CT is developed and

published using the same process as all other CDISC CT, in order to ensure a consensus based, fit for use, and harmonized set of terms.

6.1.5 CTR

<u>Clinical Trial Registry (CTR)-XML</u> lets technology vendors implement tools that support a "write once, use many times" solution based on a single XML file that holds the information needed to generate submissions for multiple clinical trials for clinical trial registry submissions, primarily to the World Health Organization (WHO), the European Medicines Agency (EMA), the EudraCT Registry, and United States <u>ClinicalTrials.gov</u>. Working alongside clinical trial registry SMEs, an evaluation was performed to determine how USDM could be utilized to assist in the population of elements required for clinical trial registries. In Version 3.0, this was restricted to <u>ClinicalTrials.gov</u>. Additional information can be found in Section 7.2, <u>Informing ClinicalTrials.gov</u> Registry.

6.1.6 ODM

Operational Data Model (ODM)-XML is a vendor-neutral, platform-independent format for exchanging and archiving clinical and translational research data, along with their associated metadata, administrative data, reference data, and audit information. The ODM-XML facilitates the regulatory-compliant acquisition, archival, and exchange of metadata and data. It has become the language of choice for representing CRF content in many EDC tools. ODM-XML v2.0 (released August 2023) added significant functionality to the ODM standard, including:

- Multilingual support
- Data query support
- Traceability (Trace-XML features) support
- HL7 FHIR interoperability
- Study/Trial Design Model in XML (SDM-XML) integration and enhancement
- CDISC 360 support
- Data capture

Although the USDM is a reference model and the ODM is a transport model, there is overlap between the standards in terms of elements related to study design (e.g., biomedical concepts) and elements related to EDC build (e.g., visits, forms, variables). Therefore, during the development of the USDM, areas of development for ODM-XML v2.0 were investigated and, where possible, aligned with USDM.

6.1.7 SDM

Study/Trial Design Model in XML (SDM-XML) is an extension of the ODM-XML and allows organizations to provide rigorous, machine-readable, interchangeable descriptions of the designs of their clinical studies, including treatment plans, eligibility, and times and events. SDM-XML defines 3 key submodules (i.e., structure, workflow, timing), permitting various levels of detail in any representation of a clinical study's design.

Note: SDM v1.0, released in 2011, was incorporated into ODM-XML v2.0. The SDM was used as an input reference model during the development of the USDM.

6.2 Relationship to Other Standards

6.2.1 ICH M11 Guideline, Clinical Study Protocol Template, and Technical Specifications

The ICH M11 guideline[4] introduced CeSHarP; the technical specification ensures that protocols are prepared in a consistent fashion and provided in a harmonized data-exchange format acceptable to regulatory authorities. The guideline, clinical study protocol template, and technical specifications were released in October 2022 for public review; where possible, these were used as reference input during USDM v3.0 development. Working closely with ICH, USDM v3.0 has been aligned to cover the breadth of sections found in the ICH M11 CeSHarP template. This allows a USDM study design to be represented in the ICH CeSHarP template. **Note:** At the time of publication of USDM v3.0, the ICH CeSHarP was still in the development phase. In future phases of USDM development, CDISC

will continue to collaborate with the ICH team in order to ensure that USDM remains aligned with the ICH M11 CeSHarP template.

6.2.2 HL7 FHIR SOA

The <u>Vulcan Schedule of Activities (SOA) Project</u> defines a pattern for a clinical trial SOA structure using FHIR resources and processes that enables sharing, interpretation, and implementation in healthcare (EHR, PHR) systems. When a subject is enrolled in a study, research personnel will be able to attach them to the ResearchSubject and ResearchStudy, connecting the CarePlan with the schedule of activities (the research visits and corresponding tests/activities).

7 USDM Features

- Overview
- Principles
- Naming Conventions
- Internal Identifiers Within the Model
- Controlled Terminology
- Study, Protocols, and Amendments
- Study Identifiers and Titles
- Study Design
- Study Roles and Organizations
- Arms and Epochs
- Activities
- Procedures
- Biomedical Concepts
- Study Timing
- <u>Indications</u>
- Study Interventions
- Study Objectives and Endpoints
- Study Estimands
- Populations, Cohorts, and Eligibility Criteria
- Unstructured Content
- Addressing Footnotes
- Syntax Templates
- XHTML Attributes
- Abbreviations

7.1 Overview

The USDM normative form is a UML model. The USDM provides the ability to define a version of a clinical study that includes:

- 1. The main study details, such as:
 - a. Version of the external protocol that the study relates to
 - b. Various identifiers allocated to the study
- 2. One or more study designs within the study, with each study design detailing:
 - a. Arms and epochs within the design and the relationships between them
 - b. Encounters planned for the study and the relationship with the epochs of the study
 - c. A detailed data specification for the data to be captured as part of the study
 - d. Procedures to be performed as part of the study design

- e. Timing of collection of data and the performance of procedures
- f. Subject populations defined within the study design
- g. Objectives and endpoints defined within the study design
- h. Study estimands defined within the study design
- i. Interventions defined as part of the study design
- j. The relevant indication

Although the USDM is designed to hold a single version of a study, the model can be used to implement systems that hold multiple versions of multiple studies.

Note: The use of the terms above and their respective definitions are defined within the USDM class definitions and the related controlled terms.

7.2 Principles

The main principles applied to the development of the USDM include:

- Try not to reinvent the wheel. At the same time, improve. Use and learn from existing models.
- Align with existing CDISC models as much as possible but do not be constrained by them.
- Where sensible, provide standardized codes from CDISC CT. Allow for aliases.
- Allow for references to any CT where sensible.
- Do not recreate the paper world.
- Be aware of model versus presentation.
- The model should represent a complete protocol, not a partially completed one. Implementators should be able to relax constraints if they are building protocols.
- The model should not prevent implementators from extending the model.
- Keep the approach simple at the start; iterate, learn, and add complexity as it is understood.
- Support the planned design, not subsequent execution.
- Support the whole protocol document (phase 3 onwards; not true for phases 1 and 2).

With respect to terminology, principles include:

- Standardize on a codelist/value set; be prescriptive.
- Where there is misalignment, standardize on the best global standard.
- Allow for regional differences (e.g., FDA in the US).

7.3 Naming Conventions

7.4 General

USDM v3.0 defines standard naming conventions. This includes improving the names of classes and, in particular, attributes to make the model more implementation friendly.

This section details the conventions used for naming and the use of attribute data types.

7.5 Class and Attribute Naming

The naming convention as currently used is:

- Nouns are used for class names.
- Every class has an attribute named "id" such that a unique identifier, within the scope of a study, can be allocated to instances of the class.
- A class can have a number of standard attributes. The attribute names should not be used for any other purpose than:
 - o name: the literal identifier (i.e., distinctive designation) for an instance of the class
 - o description: a narrative representation for an instance of the class
 - o label: the short descriptive designation for an instance of the class

 notes: a USDM relationship between the class and the CommentAnnotation class which provides the set of notes related to the class

Note: a class may employ these attributes if they are required and thus not all classes use them.

• A class can have additional attributes.

7.6 Data Types

Attributes have been provided with simple data types. The USDM generally avoids the use of complex data types. Where there is a need for a complex data type, a separate class is created.

7.7 Relationships

Relationships have, in general, been formed from the names of the class at either end of the relationship with singular names used for one-to-one relationships and plural names used for one-to-many relationships.

7.8 Internal Identifiers Within the Model

Each class defined within the UML has an identification attribute that can be used to provide a unique identifier for an instance of the class. The identifier should be unique and self-consistent within the scope of a version of a study. No attempt is made to define the form, type, or structure of these identifiers; the attributes are defined as strings. The only exception is the identifier at the head of the model within the Study class. Implementations are free to allocate the value to this field using, for example, a UUID, to ensure uniqueness within the implementation.

7.9 Controlled Terminology

Controlled terminology is referenced in multiple places across the USDM. So as to provide a mechanism to refer to controlled terms in a consistent manner, the USDM employs the Code class. The Code class uses 4 attributes to define the term being used (a code and decode pair), the terminology from which the term is taken, and the version of that terminology. This allows for any controlled term—whether CDISC, SNOMED, LOINC, or other—to be referred to in a consistent manner.

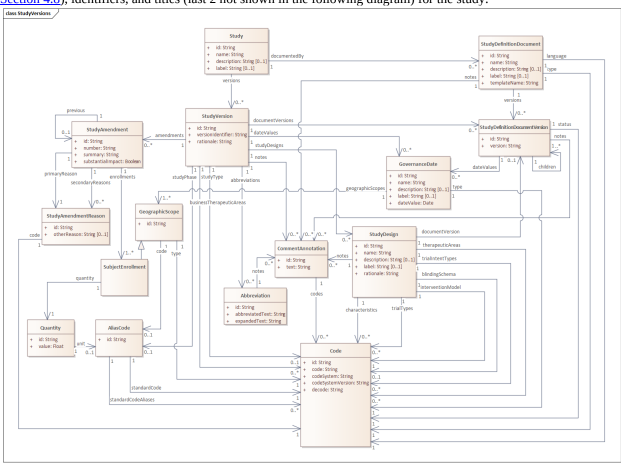
Certain attributes within the USDM Code class have been constrained to using terms from a given codelist from specified terminologies; these are specified in the controlled terminology spreadsheet. Although most of the terms referenced are CDISC CT, some other controlled vocabularies are referenced.

Where a standard code (typically a CDISC code but not always) is demanded by the model but flexibility is desirable / needed, users may include other terms (aliases) using the AliasCode class. Here one standard term is required but zero, 1, or more aliases can be provided. One particular instance is geographic references. The standard code should be from ISO 3166; other code aliases (e.g., GENC) can be provided.

7.10 Study, Protocols, and Amendments

The Study class is the root of the USDM, collecting together the definition of the study and its corresponding versions as a whole. A study is documented by a study definition document which usually is a protocol but could be of other types as well. The overarching study and the study definition document each have their versioning with corresponding governance dates. These dates are to be focused to a specific geographic scope (e.g. global, regional, country).

Because the traditional paper/PDF protocol document has been split into 2 parts (i.e., the document and an electronic design using the USDM), there is a need to link which electronic definition is valid with which version of the document. The Study Version class links to the StudyDefinitionDocumentVersion class to define to which versions of an external protocol document the study definition relates. The study version provides a few basic study details



(e.g., type, phase, rationale) and links the study with its constituent parts that include 1 or more study designs (see Section 4.8), identifiers, and titles (last 2 not shown in the following diagram) for the study.

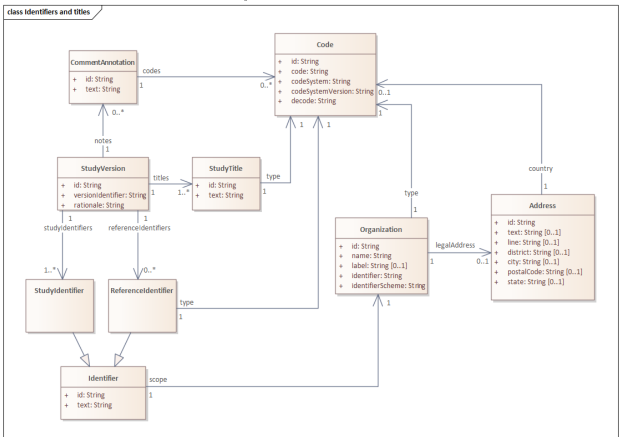
A study version may represent an amendment. Corresponding amendment details—including reasons for the amendment, number or percentage of subjects enrolled at time of amendment, and substantial impact—are captured in the Amendment class. This can be reflected in the corresponding study definition document version via the StudyVersion class. The study definition document version content is captured in the USDM as unstructured content (see Section 4.20).

Abbreviations that are used to describe the study design are defined at the study version level and can be reused (eg referenced) both in the syntax template text (e.g. for eligibility criteria or assessment conditions) as well as in unstructured document content. Some examples are presented in the paragraph <u>Abbreviations</u>. The full list defined for the study can also be used to automatically create the full list of abbreviations in the protocol document. The StudyVersion class also allows for stating the business therapeutic area. **Note:** The business therapeutic area is provided for downstream processes and for sponsor organizations to define the business areas within the enterprise handling the study. It should be noted that business therapeutic area is not the same as the therapeutic area defined in the StudyDesign class.

The Study class allows for 1 or more study designs to be included. This provides a single mechanism for master and umbrella studies. Multiple study designs are permitted so as to accommodate multiple designs that test multiple drugs and/or multiple cancer subpopulations in parallel under a single protocol without a need to develop new protocols for every trial. Typically, there would be a one-to-one relationship between study version and study design with 1 or more protocol versions related to the study covering the different designs. The studyDesign can refer to the study protocol version directly related to the specific design.

7.11 Study Identifiers and Titles

Study identifiers, reference identifiers and titles are stored in separate dedicated classes as presented in the UML below and are referred to from out of the StudyVersion class.



A study identifier specifically identifies the study represented in the data model. The Study Version class allows for links to the 1 or more study identifiers. Although multiple identifiers are permitted, the study definition should have 1, and only 1, sponsor identifier (e.g. linked to an organization with organization type 'Clinical Study Sponsor'). Note the use of ISO 3166-1 country codes within the address field.

A reference identifier may include references to overarching plans like a pediatric investigational plan number and a clinical development plan number.

One or more study titles are required for a study. They can be of different types (e.g., official, scientific, short titles). If available, the acronym should be stored as a title as well, with specifying the type as acronym.

7.12 Study Design

The StudyDesign class is the container for a single design within a study definition. It provides the slots for key parameters such as the trial type, trial intent type, blinding scheme, and intervention model. The class also provides a place to store 1 or more codes defining the therapeutic area to which the study design relates.

No controlled terminology is provided for the population of this therapeutic area field; the following table details controlled vocabularies that are available for users to populate 1 or more values into the attribute. A sponsor's own controlled terms can also be used.

CONTROLLED CONTROL	ond one of the contract of the		
Dictionary/Terminol	URL		
ogy			

EudraCT	https://eudract.ema.europa.eu/docs/technical/EUDRACT_Eutct_Pick_Lists_and_coded_va
	lues_v1_0.xls
ICD-10	https://www.icd10data.com/ICD10CM/Codes
MedDRA	https://www.meddra.org/
MeSH	https://www.ncbi.nlm.nih.gov/mesh/
NCI Thesaurus	https://ncit.nci.nih.gov/ncitbrowser/
SNOMED-CT	https://www.nlm.nih.gov/healthit/snomedct/index.html
US FDA	https://www.fda.gov/drugs/development-resources/spectrum-diseasesconditions

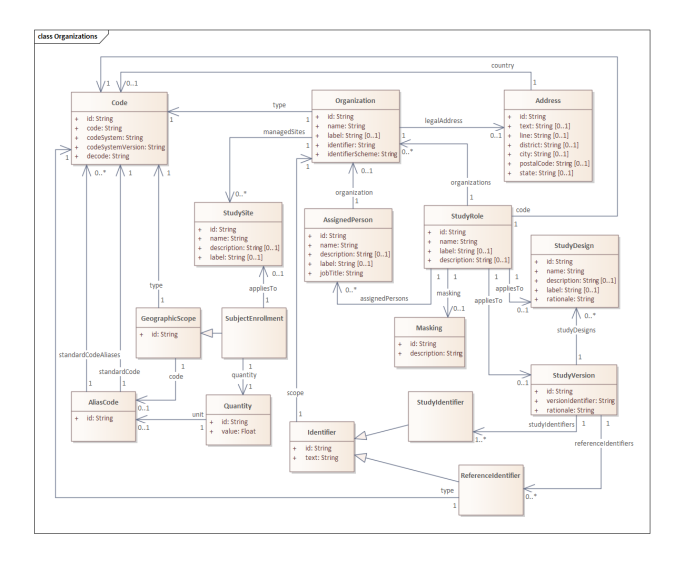
7.13 Study Roles and Organizations

A clinical study may include a number of different roles on different levels. This includes sponsors, investigators, committees, regulatory agencies and more. These roles are stored in the StudyRole class. A role may apply to the study as a whole or to one or more study designs specified within that study. Specific person names linked to a study role are specified in the AssignedPerson class. If no specific persons are assigned then the StudyRole may directly link to an organization being responsible for the role as a whole.

Organizations are organizational entities that are involved in a clinical study. The organization type identifies what kind of organization is specified (e.g., clinical study sponsor, research organization, regulatory agency, etc.). A research organization or clinical study sponsor can optionally manage 1 or more study sites. These study sites may be referred to in case a subject enrollment status for an amendment is specific for a site.

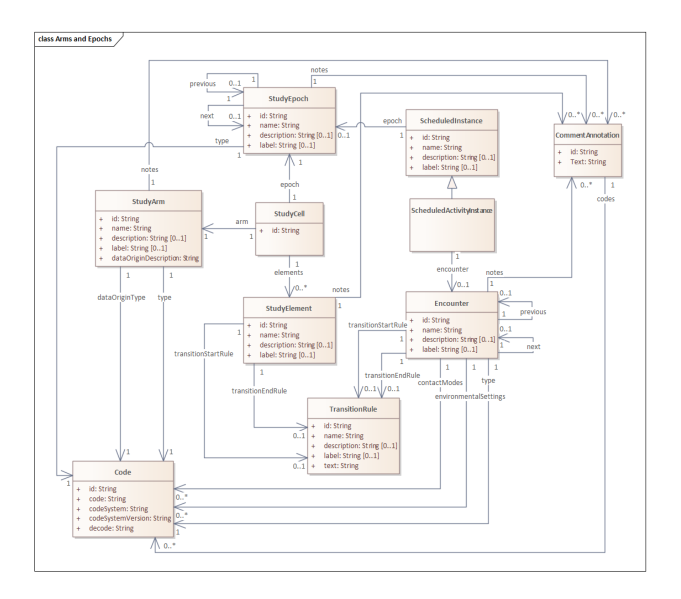
If a role is masked in a study then this should be identified by an entry and corresponding description in the masking class.

An identifier should be referring to one of the defined organizations as it's scope (see Section 4.7).



7.14 Arms and Epochs

The high-level study design consisting of the arms and epochs is defined using the StudyArm, StudyEpoch, StudyCell, and StudyElement classes. The manner in which the classes are used follows the CDISC SDTM. Epochs are related to the study encounters (a more generic term for visits) via ScheduledInstances that form a ScheduleTimeline (for more information see Section 4.14, Study Timing). StudyElements can relate to the corresponding studyInterventions that are planned for the specific StudyArm and in the specific StudyEpoch. StudyElements and Encounters have entry and exit rules that are defined using the TransitionRule class. It should be noted that although the StudyElements and Encounter classes share the use of the TransitionRule class, it is not expected that the instances within any study design will overlap; they are, most likely, distinct sets. Given that the use of the classes is based on the SDTM, the information within these classes can be used to populate the SDTM Trial Design domains (see Section 7.1).

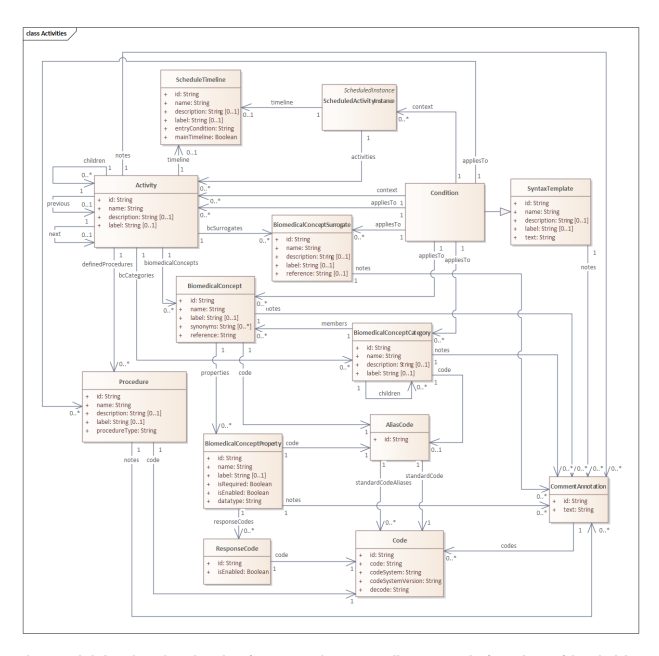


7.15 Activities

Activities are the means by which the procedures to be performed and the data to be captured are specified at a detailed level. The Activity class is used to group together data capture and procedures. The composition of these groupings is left to those designing studies and may align with the activities presented in the schedule of activities. The presentation ordering in the schedule of activities can be handled with the previous and next attributes. Any presentation groupings can be handled with the children attribute. Activities can be reused across multiple points within a study timeline via the ScheduledActivityInstance class (see Section 4.14, Study Timing).

The Activity class can be linked to 1 or more procedures (see Section 4.12), 1 or more biomedical concepts (see Section 4.13), 1 or more groups of biomedical concepts, 1 or more surrogate biomedical concepts and/or a sub timeline. A sub timeline referred to from an activity would typically be a sequency of actions covered by the activity description (e.g. blood glucose profiles, sitting/standing vital signs sequences etc.).

Activities or the corresponding assessments and procedures may be conditional. These conditions, specified in the Condition class, apply to at least 1 activity, biomedical concept, group of biomedical concepts, biomedical concept surrogate or procedure. The context of the condition can be to the activity in general (at every timepoint it is scheduled) or to a specific timepoint in the timeline via ScheduledActivityInstance.



The example below shows how the values for activities that are typically present in the first column of the schedule of activities are stored in the USDM activity class and how "grouping" headings can be accommodated. The previous and next attribute is used to identify the order of presentation while the children attribute is used to identify the group members, for example 'Efficacy' or 'Safety'. This grouping activity (e.g. having children) is typically only used for presentation purposes and is not expected to be referred to from an scheduled activity instance or to point to biomedical concepts or procedures. It is recommended that only two levels of grouping (i.e, parent and child) are used.

	Screening	Day 1	
Subject related Assessments		20,2	
Informed consent	Х		
In/Exclusion criteria	Х	Х	
Demography	X		
Medical history	X		
Randomisation		Х	
Efficacy			
Lab efficacy assessments		Χ	Х
PRO questionnaire		Χ	X
Safety			
Vital signs	Х	Χ	X
ECG	X	Χ	
Hematology	X	Χ	
Biochemistry	X	Х	
Adverse events	X	Х	Х
Intervention			
Drug dispension		Χ	Х
Drug accountability		Х	

label	id	previous	next	children
Subject related Assessments	id_01		id_02	id_02, id_03, id_04, id_05, id_06
Informed consent	id_02	id_01	id_03	
In/Exclusion criteria	id_03	id_02	id_04	
Demography	id_04	id_03	id_05	
Medical history	id_05	id_04	id_06	
Randomisation	id_06	id_05	id_07	
Efficacy	id_07	id_06	id_08	id_08, id_09
Lab efficacy assessments	id_08	id_07	id_09	
PRO questionnaire	id_09	id_08	id_10	
Safety	id_10	id_09	id_11	id_11, id_12, id_13, id_14, id_15
Vital signs	id_11	id_10	id_12	
ECG	id_12	id_11	id_13	
Hematology	id_13	id_12	id_14	
Biochemistry	id_14	id_13	id_15	
Adverse events	id_15	id_14	id_16	
Intervention	id_16	id_15	id_17	id_17, id_18
Drug dispension	id_17	id_16	id_18	
Drug accountability	id_18	id_17		

7.16 Procedures

The procedures linked to the Activity class allow for the procedures required by the activity to be detailed. A procedure consists of a free-text name and description; procedures can be classified using a free-text type attribute and coded using the code attribute. In cases where the procedure includes a study intervention (e.g., drug administration), the corresponding study intervention can be referenced.

7.17 Biomedical Concepts

The CDISC <u>Biomedical Concepts model</u> defines a clinical concept in a standardized and reusable manner; it is a specification focused on the data, not how the data are captured or processed. As such, biomedical concepts (BCs) are atomic entities and should not be split apart; to do so causes a loss of meaning. A BC is identifiable (has an identifier) and is complete (contains everything needed to use it).

A BC defines an observation but it requires context: the context of a clinical study. This is why, in the USDM, BCs are linked to activities and thus the remainder of a study design.

Within the USDM, the BC model has been represented in a manner consistent with the rest of the USDM. For example, controlled terminology references use the Code object to be compatible with all of the CT references across the USDM. Additional attributes have been added to allow for configuration as part of a study to enable or disable certain qualifiers or to constrain terminology responses to match the needs of a study (e.g., constraining units to metric values).

When a BC is included within a study design the BC can be constrained if the BC definition allows for such. When those constraints are applied or by whom is not dictated by the model; that is an implementation and process concern. For example, a study definition may leave everything in the BCs unconstrained and only when the study design is deployed in capture systems will any constraints be applied. Constraints take the form of disabling optional properties; for example, the method used for an observation does not need to be captured, or the terms for a property can be constrained (e.g., body position is always going to be supine for a particular observation and so standing can be disabled as an option). The constraints are applied via a enabled boolean flag. Some properties, such as a result, are always required. Required properties are indicated by a second boolean flag.

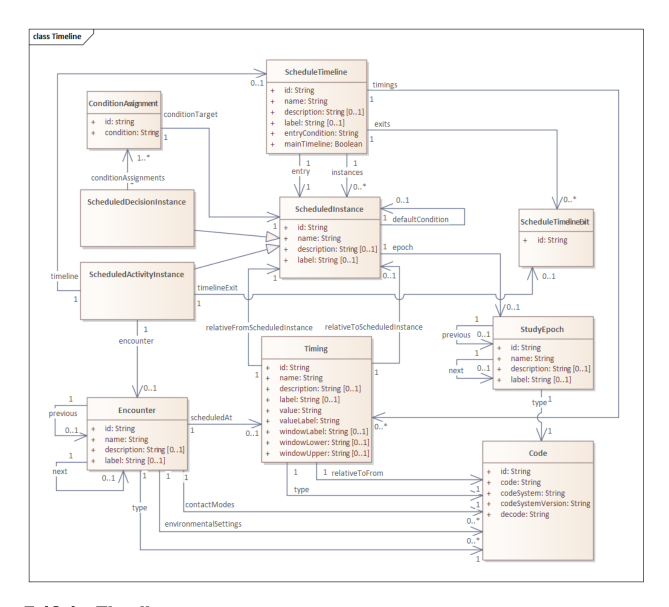
The USDM allows for the inclusion of a single BC (e.g., heart rate), a collection of BCs (e.g., vital signs preconfigured to include height, weight, heart rate, and other tests), or surrogate BCs. Surrogate BCs are a placeholder mechanism for when a BC definition is not available. This allows the name of a test to be specified but no further detail need be provided. Surrogates can contain a name and description pair for the concept required. A reference field is also provided to allow for links to reference materials (e.g., a URL for an external resource). A single BC uses the BiomedicalConcept class as its root instance connected to one or more BiomedicalConceptProperty instances to define the various properties of the BC (e.g., result value, units, qualifiers).

Some of the property nodes will require controlled terminology references; these are placed within ResponseCode instances which then onward refer to a Code instance holding the actual term reference.

One or more BCs can be grouped using a BiomedicalConceptCategory. It is assumed that, to be useful, more than a single BC should be added to a grouping such as the vital signs described above. These groupings are expected to be sponsor defined but, in the future, some can be expected to be industry defined.

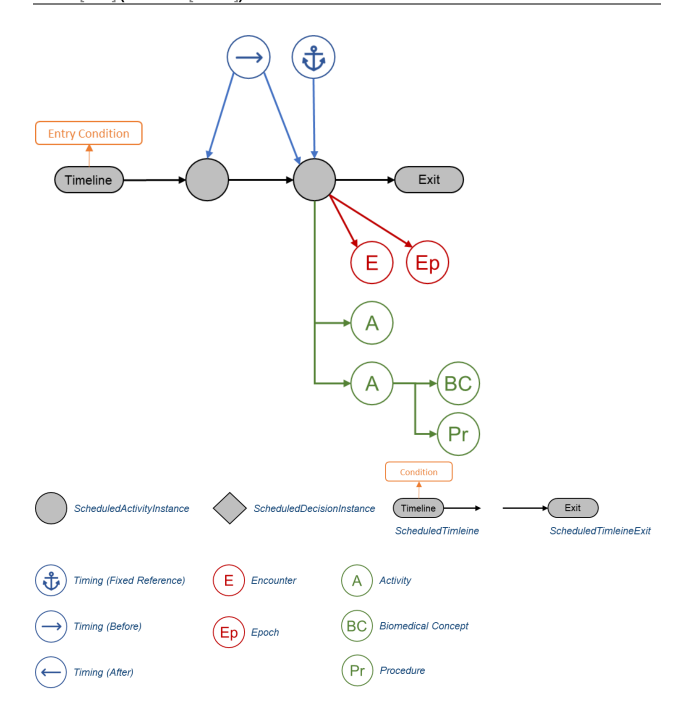
7.18 Study Timing

One of the key aspects of a study design is the timing of encounters (visits) and the activities to be performed within those encounters. The USDM includes a mechanism for building timelines that can be reused within a study and, given external library management, across studies. The corresponding classes and attributes are shown in the following UML diagram. This model allows for multiple planned timings within an encounter as well as for decision points in the study process. The corresponding information is stored in a timeline as scheduled activity instances and scheduled decision instances, respectively. Both inherit all attributes and relationships from the ScheduledInstance class (indicated by the closed arrows in the UML) and can be linked to the corresponding study epoch. The Timing class includes all timing information with details on time between instances and corresponding windowing. One or more scheduled activity instance can be related to a corresponding encounter, which is usually presented as a visit in the schedule of activities.



7.18.1 Timelines

The study timing mechanism depicted in the following figure is based on the notion of a timeline. A *timeline* is composed of an entry point with an associated entry condition (see ScheduleTimeline class), a sequence of steps (the ScheduledActivityInstance class and scheduledDecisionInstance class), timing relating the steps (the Timing class), and 1 or more exits (the ScheduleTimelineExit class) that mark the end of timeline processing. A timeline is named and can be referenced or reused within other timelines. The steps within a timeline link the encounters with the activities required for each step and thus define the timing for the encounters. The ScheduledActivityInstance class is the link between the high-level study design defined by the StudyArms and StudyEpochs classes, the Encounter classes, and the detailed study design defined by the Activity class.



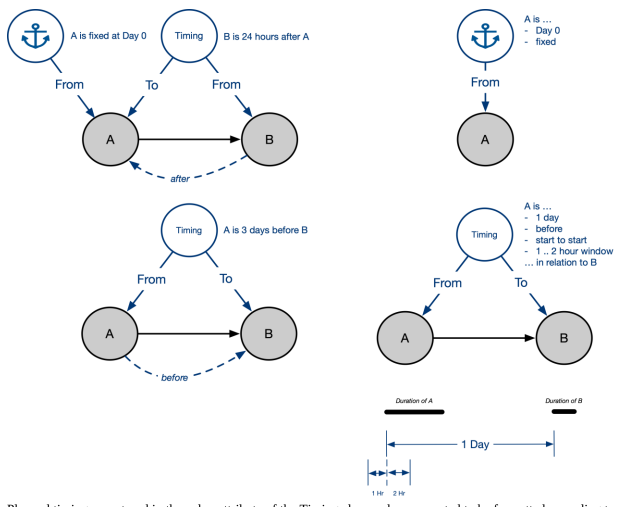
7.18.2 Timing

The timing between steps comprises a relative time of before or after, and an anchor time that is fixed. The following figure illustrates the timing capabilities. The Timing class allows for explicit timing to be built into a timeline using a combination of anchors (fixed timing) and relative timing. The timing definitions should be read as "the <Timing.relativeFromScheduledInstance> node is <Timing.value> <Timing.type of before or after> the <Timing.relativeToScheduledInstance> node". The timing definition allows for further precision in the timing by specifying the relativeToFrom type.

For anchors, the relativeFrom node refers to the scheduled instance that provides the fixed reference. The corresponding relativeTo node should either refer to the same scheduled instance or should be missing.

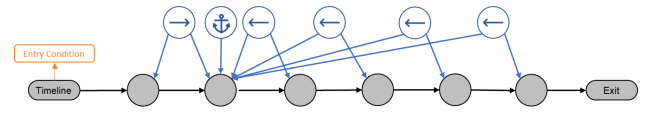
A timing may be referenced from an Encounter using the scheduleAt attribute allowing for a specific encounter timing and corresponding windowing to be defined and presented in a scheduled of activities. An Encounter timing might potentially overarch multiple scheduledInstances representing different blocks of activities within an encounter.

Note that in the timing diagrams the relativeFromScheduledInstance and relativeToScheduledInstance relationships have been shortened ("From" and "To," respectively) so as to make the diagrams readable.



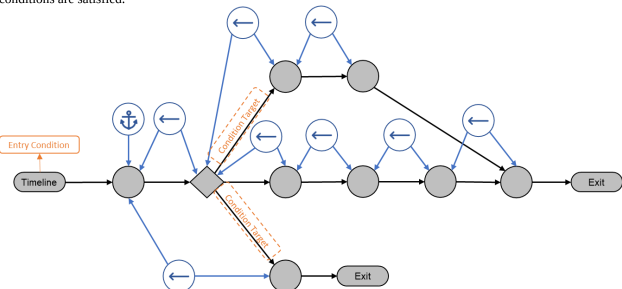
Planned timings are stored in the value attribute of the Timing class and are expected to be formatted according to ISO 8601. A corresponding window can be identified using the window attributes. The windowLower and windowUpper attributes are also expected to be formatted according to ISO 8601. Textual representations of these values can be stored in the valueLabel and windowLabel attributes, respectively.

Note that timings can be defined between each consecutive scheduled instance or all or part of the timings can be related to a fixed (anchor) timepoint:

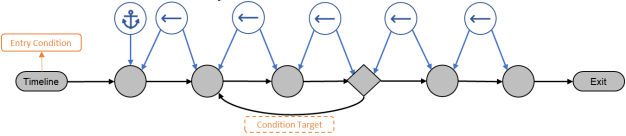


7.18.3 Decisions and Branching

Decisions and branching are handled using instances of the ScheduledDecisionInstance class within a timeline as shown in the following figure. Each decision point can handle multiple conditions; for example, simple yes/no decisions as well as a complex switch with multiple paths. Each possible route is set up with an associated destination. For switches, there should be a "default" condition specified for the case when none of the other conditions are satisfied.



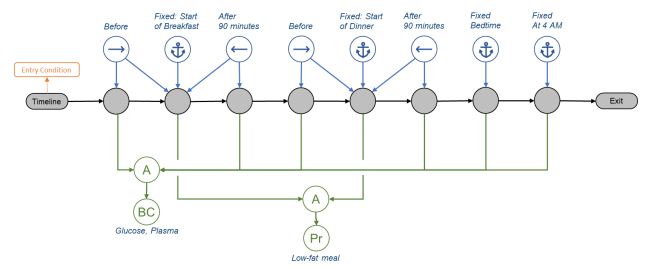
The decision can also be used to create cycles:



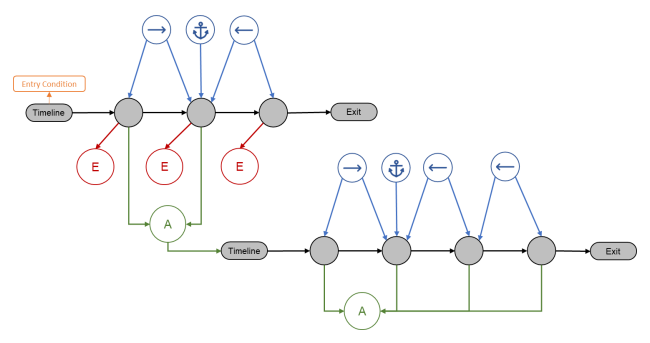
Descriptions of the decision and pointer are defined using the conditionAssignment class. This class includes 2 attributes: a description of a condition and the reference to the target instance of the scheduledActivityInstance class that it points to once this condition is met—for example: "not reached cycle 12 and fulfilling eligibility to enter next cycle", "ScheduledActivityInstance_2".

7.18.4 Profiles

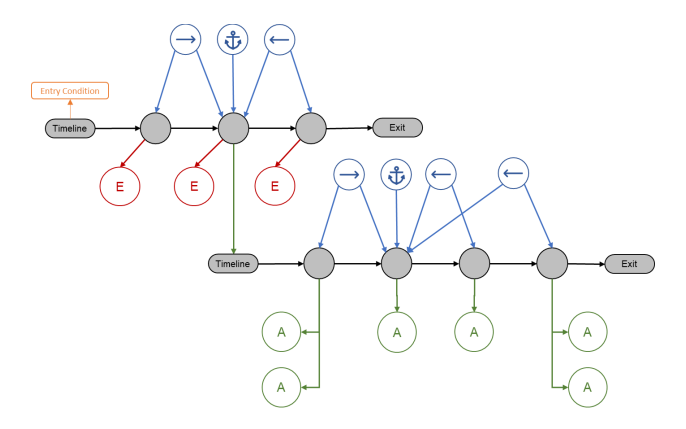
Profiles can be created using the various classes, as depicted in the following figure. A profile is another use of the timeline pattern and may reflect a sub-timeline within an encounter. A condition for entry can be defined but need not be. In this example, anchors are used to fix meal times over a single day and the associated observations scheduled in relation to the fixed meal times. The activities are shared across the steps within the profile.



The profile can be "attached" to an activity using the ActivityTimeLineId attribute so that it is executed as part of that activity, as illustrated in the following figure. This is useful for a sequence of repeated measures within the same activity.

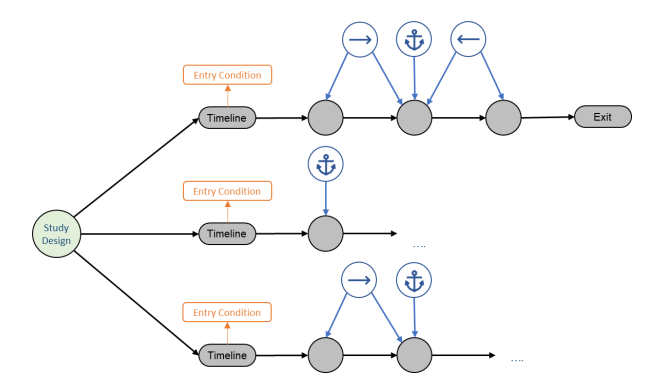


The timeline can also be attached to a ScheduledActivityInstance from another timeline using the timeline reference, thus allowing timepoints within a visit to be constructed, as shown in the following figure.



7.18.5 Unscheduled Visits

Unscheduled visits within a study are handled by creating separate timelines for each unscheduled "event" that needs to be handled within the study design. A study design would typically have 1 "main" timeline with a condition such as "subject identified". Further timelines can be created and linked to the StudyDesign instance with the timeline having an appropriate condition (e.g., "Adverse event", "Lost contact with subject"). Each timeline is then free to detail the steps taken under the respective circumstances.



7.18.6 Timeline Exit

It should be noted that the ScheduledTimelineExit instance does not perform any role other than marking the end of a timeline. It is linked from the last ScheduledActivityInstance instances in the timeline.

7.19 Indications

The indication for a study design can be placed into the Indication class. Each indication has a textual description plus the ability to define 1 or more codes from external code systems (including a sponsor's own terminology) that define the indication.

The attribute isRareDisease can be utilized to indicate whether an indication is regarded as a rare disease according to applicable rare disease registries (e.g., NIH GARD, <u>Genetic and Rare Diseases Information Center</u>).

7.20 Study Interventions

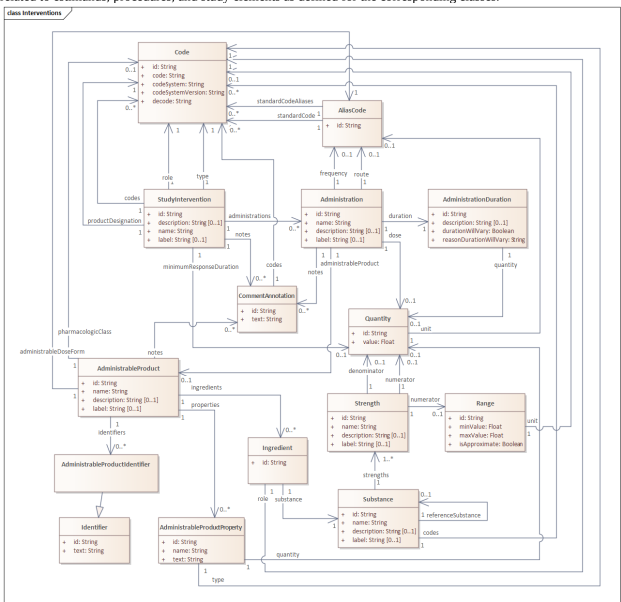
The interventions for a study can be placed into the StudyIntervention class. Each intervention needs to be defined by role, type and productDesignation. Optionally, information on 1 or more codes from external coding systems and the expected duration to minimum response can be added. Corresponding administration details can be specified in the Administration class. The frequency, dose, route, administrable product and duration can be specified for each administration.

For each administrable product optionally, information on the pharmacological class, and 1 or more identifiers, properties and ingredients may be specified. Each ingredient specified by its substance may have a reference substance. The corresponding reference strength represents the strength (quantitative composition) of the active moiety of the active substance or of another substance used to express the strength of the product. There are situations when the active substance and active moiety are different resulting in different expression of the strength. The strength of each substance is specified in the strength class using a numerator and preferably a denominator. In case the strength is not exact but estimated to be within a range, the numerator can be expressed as a range using

minValue and maxValue attributes instead of the quantity value attribute. For IDMP, the strength value or minValue and corresponding denominator value refers to the IDMP strength lower limit, while, if applicable, the strength maxValue and corresponding denominator value, refers to the strength upper limit.

Note that the internal sponsor code or compound number for the administrable product can be stored as the administrable product identifier.

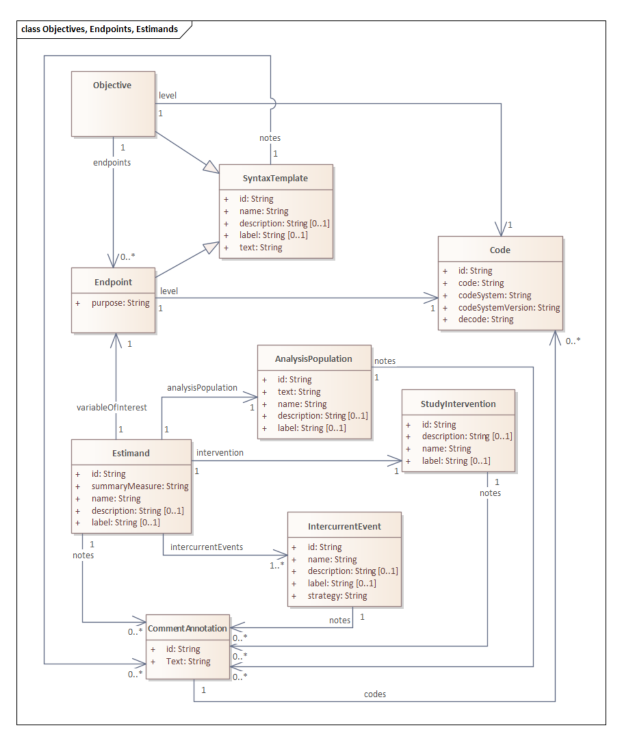
Study interventions need to be directly referred to from the Study Design class. In addition, they can be directly related to estimands, procedures, and study elements as defined for the corresponding classes.



7.21 Study Objectives and Endpoints

The study design objectives and endpoints can be defined within the Objective class and the Endpoint class. The Objective class allows for the textual description of the objective and its level (e.g., primary, secondary, exploratory) and a link to 1 or more associated endpoints containing the endpoint definition in textual form. Both the objective

and endpoint class inherit from the syntax template (see <u>Section 4.21</u>), allowing for references to information stored elsewhere in the data model. The endpoint may be a variable of interest for the study estimand (see <u>Section 4.18</u>).



7.22 Study Estimands

Aligning to the ICH guideline E9 (R1) addendum,[5] study estimands and the definition of the treatments to be investigated, the population, the variable, and the handling of intercurrent events (ICEs) are handled within the Estimand, IntercurrentEvent, and AnalysisPopulation classes along with the relationships to endpoints (for the variable of interest; see Section 4.17) and study intervention (see Section 4.16) for the treatment.

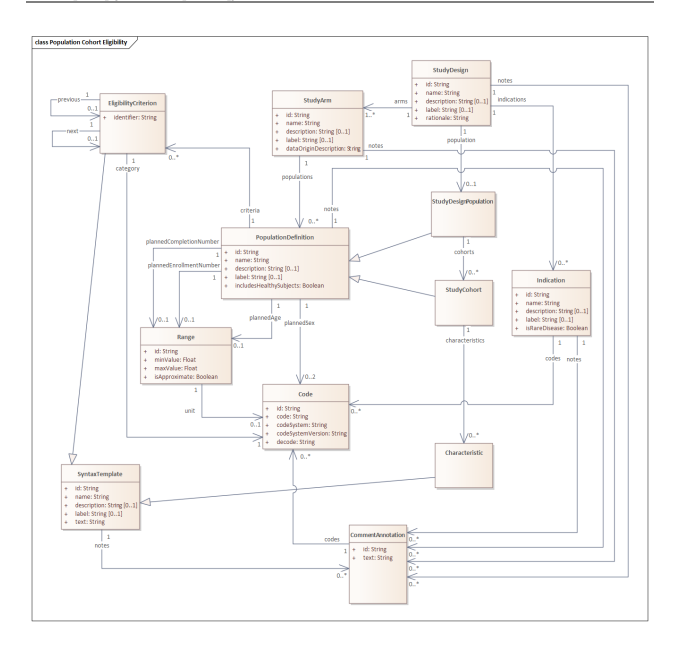
7.23 Populations, Cohorts, and Eligibility Criteria

Population and cohort definitions define a (sub-)group of subjects that take part in the study. The parent class PopulationDefinition is used to define a group of patients in general. This class includes references to the eligibility criteria that are applicable to this population. All the elements of the PopulationDefinition class are inherited by both the StudyDesignPopulation class, which stores the population details for a specific study design, and the StudyCohort class, which stores the details of subpopulations that, based on their characteristics, may deviate in how they are treated, assessed, or analyzed.

In addition to the inherited attributes from the PopulationDefinition class, the StudyDesignPopulation class may refer to the corresponding subgroups stored as study cohorts. The standard PopulationDefinition attributes criteria, PlannedCompletionNumber and/or plannedEnrollmentNumber, plannedAge, and plannedSex are either defined at the StudyDesignPopulation level or at the StudyCohort level. The allowed coded values for plannedSex are 'male' or 'female'. Either one, or both can be specified for a study design population or for a study cohort.

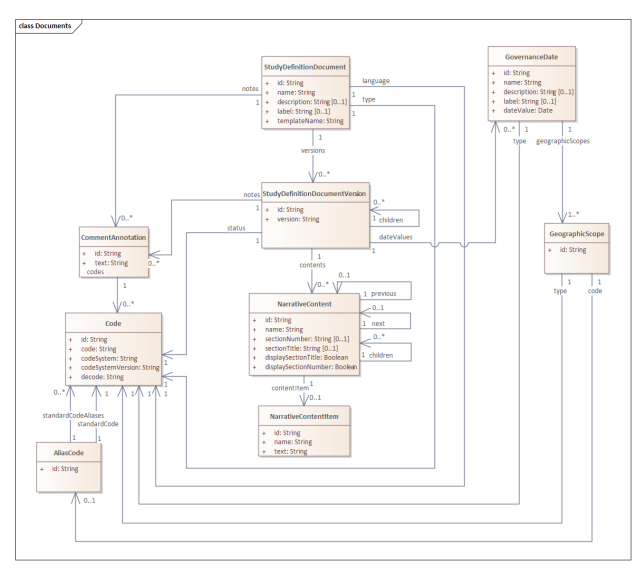
The StudyCohort class may refer to additional characteristics not defined by any of the other attributes in the PopulationDefinition class. These characteristics are stored in the Characteristic class, which inherits its attributes from the Syntax Template class (see Section 4.21) and can thus refer to any item stored elsewhere in the USDM. Eligibility criteria inherit from the Syntax Template class as well, allowing for referencing any item stored in the USDM, such as assessments stored as BCs or an indication stored in the Indication class. They are defined within a study version which allows reuse within different study designs and different cohorts. The previous and next attributes define the presentation ordering within an eligibility criterion category or overall. The identifier attribute may be used to store the short name used for mapping to SDTM TI domains (see Creation of SDTM Trial Design Domains).

In case needed, specific notes for example for grouping, mapping or providing additional information can be added to the items in a class. Corresponding codes can optionally be added to these notes aligning with internal or external standards that are applicable to the notes.



7.24 Unstructured Content

Study protocols and other study definition documents include content that is best described as "unstructured content", granting the author considerable flexibility in determining what information to include, the level of detail it will contain, the order in which it is introduced and discussed, and how it will be presented. Blocks of unstructured content can range from short text statements to many paragraphs which may also contain figures and tables. The Narrative Content class in the UML is modeled to contain such blocks of user-defined unstructured content using HTML format. The recursive nature of this class with its attribute "children" provides the user the ability to add multiple named blocks of unstructured content, allowing for a hierarchy of related information to be built up and ordered by the section number and/or the "previous" and "next" attributes. The actual blocks of unstructured content are stored in the NarrativeContentItem class allowing for reuse within and between documents. The HTML format of the "text" attribute and the section ordering provides the capability for organizing the information in a way that is compatible with any required document structure such as ICH M11,[4] the



7.25 Addressing Footnotes

Information represented by footnotes in a schedule of activities (SOA) can be stored structurally in the USDM and as such can be parsed and presented as footnotes when feasible. By using this computer-readable format, the often complex and extensive footnote information is more usable for downstream processes. This section describes the following different types of footnotes that may be identified in SOAs and how they can be stored in the USDM:

- Footnotes representing sub-timelines
- Footnotes representing timing and/or order of activities
- Footnotes representing alternative visit schedules
- Footnotes representing conditional activities, assessments, and procedures
- Repeated activities not presented in the SOA

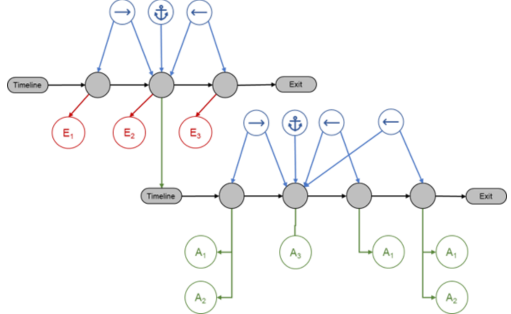
- Footnotes representing optional alternative encounter methods
- Footnotes representing measurements to be done for a specified activity
- Footnotes representing optional alternative measurement methods
- Additional instructions for procedures and/or performing assessments
- Visit and timing window information
- Eligibility requirements
- Complex combinations

7.26 Footnotes Representing Sub-timelines

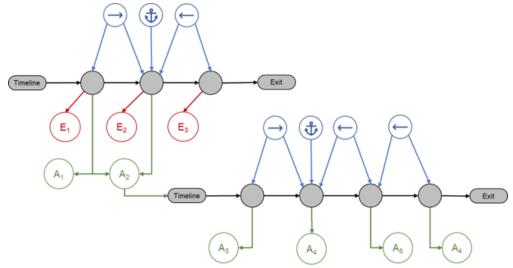
These footnotes indicate at what exact timepoints activities not presented in the SOA should be performed, for example:

- 1. Blood samples for ... predose, 1h, 24 h, ...
- 2. X assessment to be performed predose and at 40 minutes and 1.5h postdose
- 3. Measurement after 5 minutes in supine position and after 3 minutes in standing position

In case of assessments relating to dosing (examples 1 and 2), individual timepoints can be stored as ScheduledActivityInstances forming together a sub-timeline (see following diagram). This sub-timeline is referred to from a ScheduledActivityInstance on the main timeline. The time relationships (->, <- in the diagram) of these instances will be defined using the corresponding Timing classes. The timing related to the instance for the dosing activity (A_3) is defined as the anchor. Activities such as pharmacokinetic samples (A_1) and vital signs measurements (A_2) can then be added as needed, reflecting the correct timings related to dosing. Sub-timelines can be reused across multiple ScheduledActivityInstances on the main timeline.



In case of an assessment sequence relating to 1 activity (e.g., repeated blood pressure measurements in different positions), a sub-timeline can be directly referenced from the corresponding activity using the timeline relationship in this class (see following diagram). The activity A_2 (e.g., vital signs), refers to the sub-timeline indicating the corresponding positioning and assessment actions. For example, put subject in supine position (A_3), assess blood pressure (A_4); put subject in standing position (A_5) and repeat the blood pressure assessments (A_4). The timings in between are defined by the information in the corresponding Timing class.



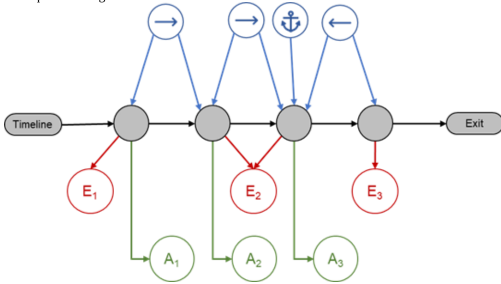
See Section 4.14, Study Timing, for more information on timelines.

7.27 Footnotes Representing Timing and/or Order of Activities

These footnotes indicate an order of activities and what should be done first, for example:

- 1. Informed consent must be obtained prior to any study-related procedure
- 2. Assessment X should be done before all other
- 3. Assessments to be done on day of admission

A simple sequence of 1 activity or groups of activities can be represented by separate instances of the scheduledActivityInstance class in the main timeline pointing to the same encounter. For example, in the following diagram, encounter E2 includes 2 scheduledactivityInstances. The first links to activities that need to be done prior to any other activity (e.g., informed consent) and the second scheduledActivityInstance relates to all other activities that are required during that encounter.



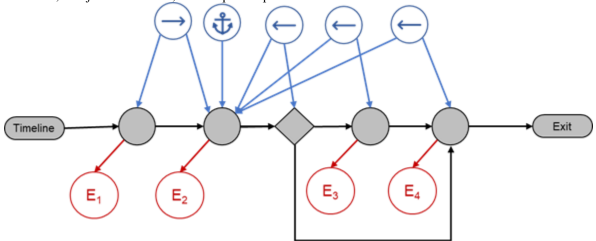
7.28 Footnotes Representing Alternative Visit Schedules

These footnotes indicate optional alternative visits based on conditions, for example:

1. Visits in case of events, inability to continue, or withdrawal (early-withdrawal visit)

- 2. An additional optional period of up to 3 weeks is permitted
- 3. Visits can occur on same day if no additional period is needed

To optionally add a visit, a scheduledDecisionInstance needs to be added to the timeline. Apart from the default next step in the timeline (defined by a defaultCondition), this scheduledDecisionInstance includes a condition and corresponding alternative next step that can be defined. In the following diagram, encounter E₃ is skipped when the condition is met. This condition as defined in the attribute conditionAssignments could then be "inability to continue", "subject withdrawn", or "no optional period of 3 weeks".



Example 3, visits occurring on the same day, is more complex. Visits can optionally be combined; the ScheduledDecisionInstance needs to be set to "no additional period needed?" If yes, then the next visit (E_3) can be skipped. In cases where activities were planned at this skipped visit E_3 (and not at the previous visit E_2), these should be added to the previous visit E_2 with the conditionality that they only need to be done when the next visit is skipped.

7.29 Footnotes Representing Conditional Activities, Assessments, and Procedures

These footnotes indicate conditions for a specified activity to be performed (or not), such as:

- 1. Assessments only for women with childbearing potential
- 2. At the discretion of the investigator
- 3. Assessments only if within x days after y
- 4. Only in case of extra wash-out needed; all others to perform assessment at end of week x
- 5. Discharge after criteria for discharge are met
- 6. Only if dipstick urinalysis is positive
- 7. Assessment to be done every 3 cycles
- 8. Only for subjects electing to participate in the additional substudy
- 9. If needed

These footnotes can be stored in the Condition class. The footnote text is stored in the text attribute and can optionally link to other elements stored in the USDM as described for syntax templates (see Section 4.21). Each specified condition in this class applies to the whole activity, a BC, a BC category, a BC surrogate, or a procedure. The context indicates to what part of the SOA it applies. This relates to where the footnote indicator is placed in the SOA. A footnote directly linked to the activity description is applicable for all occasions of that activity and should therefore have the context related to that activity. If the condition holds for a specific timepoint of that activity, then the context should be set to the corresponding scheduledActivityInstance to indicate when it is applicable. See Section 4.11, Activities, for more information.

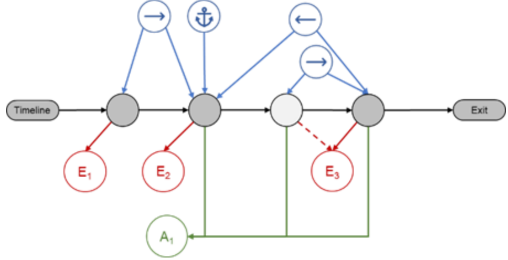
7.30 Repeated Activities Not Presented in the SOA

These footnotes specify activities that are not directly presented in the SOA because they need to be done in between regular visits, for example:

1. Questionnaire will be filled in every 2 weeks until ...

2. During run-in period, patients will perform XX measurements and inhale placebo medication at approximately 12-hour intervals for a minimum of 14 days and maximum of 21 days.

The first step in mapping these activities is to identify instances where they do not match the regular encounters represented in the SOA. These instances need to be added as ScheduledActivityInstances to the timeline with the corresponding timing information. The implementer can choose to create a separate encounter for them or to link them to the last or next encounter as required by the implementation and downstream processes (e.g., EDC setup).



7.31 Footnotes Representing Optional Alternative Encounter Methods

These footnotes specify potential encounter methods, such as:

- 1. Performed by telephone by qualified staff
- 2. If regularly allowed, visits may take place at home

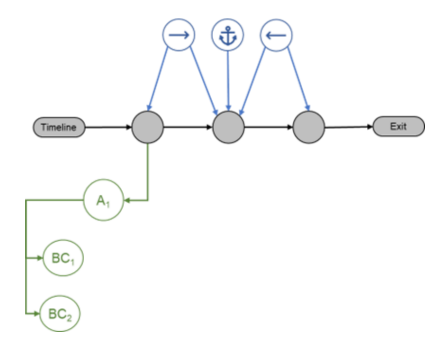
The encounter methods are specified by the attributes environmental Setting and contact Modes in the Encounter class. More than 1 contact Mode may be entered if optional alternative encounter methods are allowed.

7.32 Footnotes Representing Measurements to Be Done for a Specified Activity

In most protocols the exact assessments to be done are specified in dedicated paragraphs. However, in some cases, they are specified in the footnotes of the SOA, for example:

- 1. Hematology must include CBC with differential including but not limited to
- 2. T/B/NK cell count (i.e. CD3, CD4, CD8, CD19, CD16/56)

These assessments can be specified as BCs and linked to the corresponding SOA activity as shown in the following diagram.



7.33 Footnotes Representing Optional Alternative Measurement Methods

These footnotes indicate more than 1 alternative for an assessment, for example:

- 1. Diagnosis confirmed with either chest x-ray or CT scan
- 2. Urine or plasma pregnancy test

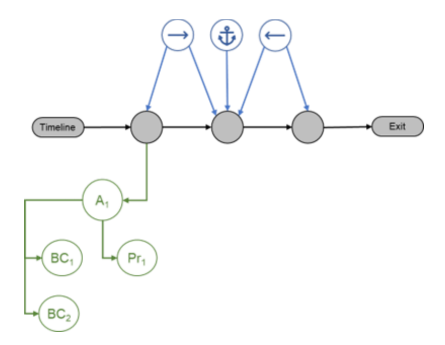
As with conditional footnotes, these footnotes can be handled using the Condition class. The text can then be stored in the corresponding text attribute. Both assessments need to be specified as a BC, procedure, or BC surrogate. The specified condition then can be related to both using the appliesTo relationship.

7.34 Additional Instructions for Procedures and/or Performing Assessments

These footnotes give details on how assessments need to be done, for example:

- 1. A ruler will be provided to assess ...
- 2. Samples will be sent to ...
- 3. Subjects should adhere to low-fat diet on day of sample collection
- 4. In order to assess y, the add-on medication should be continued for at least x weeks
- 5. X will be assessed by a blinded assessor
- 6. Patients should be instructed to use the inhaler in the morning at approximately the same time

Depending on the nature and level of instruction, this can be included in the BC when directly related to a specific assessment or added as a procedure (Pr_1) to the same activity as illustrated in the following diagram.



7.35 Visit and Timing Window Information

Visit window information is often shown in the column header of the corresponding visit, but in some cases may be added as footnotes; for example:

- 1. Assessments need to be done within 10 minutes after dosing
- 2. Visits need to take place between 5 and 10 days after dosing

As explained in Section 4.14, <u>Study Timing</u>, all specific groups of activities that occur at a specific timepoint are stored as separate scheduledActivityInstances and are linked to the corresponding timing. This timing class has attributes that can be used to specify the timing window. The window attribute is used to store the textual value of the window (e.g., "within 10 minutes after dosing") whereas the windowLower and windowUpper attributes are used for the computer readable version in ISO 8601 format (e.g., "T0M", "T10M").

7.36 Eligibility Requirements

Eligibility criteria are stored in the Eligibility Criteria class (see Section 4.19, <u>Populations, Cohorts, and Eligibility Criteria</u>). In some cases they are repeated in the SOA; for example:

- 1. Screening spirometry must demonstrate a value of In the morning of the first day of treatment value must also be in range
- 2. Patients must demonstrate >= 15% reversibility of FEV1 within .. following inhalation of ...

The EligibilityCriteria class uses text templates for the specifications of the criteria. Using these text templates, criteria can refer to the corresponding activity or assessment (BC) in the SOA. If required, these cross-references could be used by an implementation to link the criteria to the SOA and present them with the corresponding activities in the SOA.

7.37 Complex Combinations

Footnotes are often complex, long text that includes different kinds of requirements (e.g., a combination of timing, duration, conditionality, and/or methods), such as:

1. All subjects will perform a X profile for any 3 days (not required to be consecutive) during week (-2) to week (01), week 11-12, week 23-24 and week 51-52. Blood glucose readings will consist of 3 preprandial measurements (1-15 minutes before breakfast, 1-15 minutes before lunch, and 1-15 minutes before dinner) AND 3 postprandial measurements (1~1-2 hours after breakfast, 1~1-2 hours after lunch, and 1~1-2 hours

- after dinner).) The initial preprandial 6-point glucose measurement on the x day should be a fasting plasma glucose reading.
- 2. SpO2 before activity (baseline), during activity until the end of anaesthesia, and during postoperative recovery

For the purpose of comprehensibility of the SOA and for consistency throughout the study process, it is helpful to deduct the separate requirements from these footnotes and digitize them according to the solutions presented in this section.

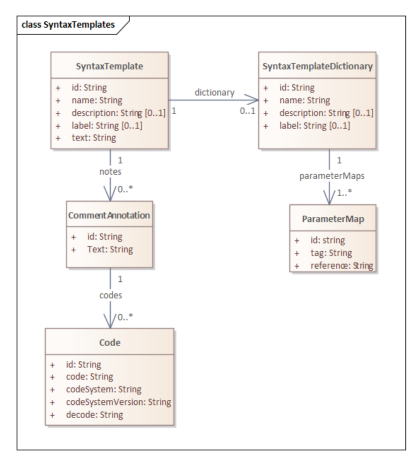
7.38 Syntax Templates

With syntax templates, human-interpretable plain text sentences are structured and linked to structured items held elsewhere in the USDM. Examples of items typically represented in the protocol as plain text that might be structured include:

- Endpoints that can be linked to a corresponding assessment and timing
- Objectives that can be linked to corresponding interventions and indications
- Eligibility criteria referring to an indication, a population, minimum and maximum age, and/or 1 or more assessments
- Conditions that can be linked to a corresponding BC or indication
- Cohort characteristics that can be linked to corresponding BCs or indications

The links are achieved by inserting tags into the plain text that reference structured content that is to be inserted into the text. These tags can be reused multiple times. This allows for consistency throughout the study design. In addition, the structured items can be more readily processed in downstream systems. The intent is that structured text allows for eligibility criteria, endpoints, objectives, and so on to be standardized and thus reused across studies, facilitating comparison and meta-analyses.

The syntax template classes are presented in the following UML.



The attributes and relationships of the SyntaxTemplate class are inherited by any class that is reusing its capabilities (e.g., Endpoint, EligibilityCriterion, Characteristic, termed "template instances"). The text attribute stores the structured text of the corresponding endpoint, criterion, or characteristic. The text attribute contains free text with embedded XHTML tags that refer to the mapping in the SyntaxTemplateDictionary. Within the SyntaxTemplateDictionary class, dictionaries can be defined that link the tags to the corresponding structured data references (to data stored elsewhere in the USDM data model) or to a fixed value.

The tags used within the text attribute of SyntaxTemplate are formatted as follows:

<usdm:tag name="parametername"/>

These tags are used as illustrated in the following example:

Subjects shall be between <usdm:tag name="min_age"/> and <usdm:tag name="max_age"/>

Instances of the SyntaxTemplateDirectory class are linked to 1 or more ParameterMap class instances. Each ParameterMap instance includes the tag (stored in the tag attribute) and a single reference or fixed value (stored in the reference attribute) as follows:

<usdm:ref klass="klassName" id="idValue" attribute="attributeName"/> or 'fixedValue'
in which:

klassName is the name of the class that holds the referenced structured data.

idValue is the id attribute value of the referenced instance of klassName.

attributeName is the name of the referenced data attribute within klassName.

fixedValue is a fixed string.

Some examples of ParameterMap references are (formatted here as tag: reference or fixedValue):

min_age: <usdm:ref klass="Range" id="Range_3" attribute="minValue"/>

max_age: <usdm:ref klass="Range" id="Range_3" attribute="maxValue"/>

 $Study Population: < usdm: ref~klass = "Study Design Population" id = "Study Design Population_1" \\$

attribute="description"/>

RefHbMax: "7.0"

It should be noted that instances of classes that inherit from SyntaxTemplate, the template instances, inherit the dictionary relationship to the SyntaxTemplateDictionary class. Each of these template instances references a single

dictionary but the dictionary can be shared across 1 or more of the template instances. Thus it is possible that a single dictionary instance—named, for example, StudyDictionary—containing a wide range of tags might be used by all the template instances or 1 dictionary instance could be created for the IE instances (named, for example, IE Dictionary), 1 dictionary instance for the Objectives and Endpoints template instances (named, for example, OEDictionary), or some mix thereof as required by implementors.

7.39 XHTML Attributes

The SyntaxTemplate and NarrativeContentItem classes each contain an attribute that contain XHTML formatted text: They are

- SyntaxTemplate text attribute
- NarrativeContentItem text attribute

The content held within these attributes should be treated at XHTML content and processed as such. It is recommended that a single root <div xmlns="http://www.w3.org/1999/xhtml"> element is used to wrap the content of the attribute. These attributes can also contain <usdm:ref> elements used to reference content held within the remainder of the model. These elements use 3 attributes to form a complete reference:

'<usdm:ref klass="klassName" id="idValue" attribute="attributeName"/>' where:

- *klassName* is the name of the class that holds the referenced data element.
- *idValue* is the id value of the referenced data element within *klassName*.
- attributeName is the attribute name of the referenced data element within klassName.

Further details of the use of these references can be found in Sections 4.20, <u>Unstructured Content</u>, and 4.21, <u>Syntax Template</u>.

7.40 Abbreviations

8 General

Abbreviation are often used with protocol documents. So as to allow for consistency of definitions throughout the study definition documents as well as in downstream processes, the USDM allows for abbreviations to be defined at the study version level. This is shown in the UML in paragraph Study, Protocols, and Amendments.

Abbreviations can be reused (i.e. referenced) both from within unstructured document content as well as from within syntax template text (e.g. for eligibility criteria or assessment conditions). In addition, the full list of abbreviations can be easily used to automatically create the full list of abbreviations in the corresponding protocol document section.

9 Abbreviations

Abbreviations consist of two parts, the abbreviated text and the expanded text. Several examples of Abbreviation instances are shown below:

Abbrevi	iation	
id	abbreviated Text	expandedText
Abbr_1	AD	Alzheimer Disease
Abbr_2	MMSE	Mini-Mental State Examination
Abbr_3	CDR	Clinical Dementia Rating Scale
Abbr_4	FCSRT	Free and Cued Selective Recall Reminding Test
Abbr_5	AChE-Is	acetylcholinesterase inhibitors
Abbr_6	DAT	Dementia of Alzheimer Type

10 Referencing From Unstructured Text

Unstructured text (held within NarrativeContentItem instances) can directly reference an abbreviation (abbreviatedText) and/or the expanded text (expandedText) using XHTML referencing (see XHTML Attributes). An example of a text item concerning the rationale is shown below. Note the example references the above example abbreviations:

Narrativ	eContentIte	em
id	Name	text
Item001	Rationale1	<pre><div xmlns="http://www.w3.org/1999/xhtml">Currently approved <usdm:ref< pre=""></usdm:ref<></div></pre>
		klass="Abbreviation" id="Abbr_1" attribute="abbreviatedText"/> treatment is purely
		symptomatic. Registered symptomatic treatment consists of <usdm:ref <="" klass="Abbreviation" td=""></usdm:ref>
		id="Abbr_5" attribute="expandedText"/> (<usdm:ref <="" id="Abbr_5" klass="Abbreviation" td=""></usdm:ref>
		attribute="abbreviatedText"/>) and memantine. <usdm:ref <="" klass="Abbreviation" td=""></usdm:ref>
		id="Abbr_5" attribute="abbreviatedText"/> in general and donepezil in particular can be
		currently regarded as gold standard for treatment of mild-to moderate <usdm:ref< td=""></usdm:ref<>
		klass="Abbreviation" id="Abbr_6" attribute="abbreviatedText"/> and is considered as
		reference drug.

11 Referencing From Syntax Templates

Abbreviations can also be referenced from <u>syntax templates</u>. Two examples are given in the following sections. Note the examples reference the above example abbreviations.

11.1 Objective

An objective is defined for Alzheimer's Disease which is abbreviated to AD. The objective class is based on syntax templates and therefore we can tag attributes stored with the associated dictionary and parameter maps. Instead of using the AD as text it is replaced by a corresponding tag as follows:

Objective.text= '<div>To assess the efficacy, safety and tolerability of different doses of Study Drug compared to placebo in treatment of prodromal **<usdm:tag name=" AD"**/><div>'

11.2 Inclusion Criterion

The inclusion criterion for the same study is defining the diagnosis and the corresponding definition. The EligibilityCriterion class which stores these criteria is also based on syntax templates and therefore we can also replace all the abbreviations by the corresponding tags as follows.

EligibilityCriterion.text= '<div>Patients with a confirmed diagnosis of prodromal **<usdm:tag name=''_AD''/>** on neuropsychological testing defined as: - Mini-Mental State Examination **<usdm:tag name=''_MMSE''/>** score: ≥ 24 and - a global **<usdm:tag name=''_CDR''/>** -score of 0 or 0.5 and - Free and Cued Selective Recall Reminding Test (**<usdm:tag name=''_FCSRT''/>**) score: o free recall test: ≤ 20 (out of 48) and o total recall test: ≤ 42 (out of 48).<div>'

The reference from the tag used in the syntax template texts of Objective and EligibilityCriterion to the specific instance in the Abbreviation class is specified in the SyntaxTemplateDictionary and the ParameterMap instances specified within the dictionary as follows:

Parameter	rMap	
id	tag	reference
Param001	_AD	<pre><usdm:ref attribute="abbreviatedText" id="Abbr_1" klass="Abbreviation"></usdm:ref></pre>
Param002	_MMSE	<pre><usdm:ref attribute="abbreviatedText" id="Abbr_2" klass="Abbreviation"></usdm:ref></pre>
Param003	_CDR	<pre><usdm:ref attribute="abbreviatedText" id="Abbr_3" klass="Abbreviation"></usdm:ref></pre>
Param004	_FCSRT	<pre><usdm:ref attribute="abbreviatedText" id="Abbr_4" klass="Abbreviation"></usdm:ref></pre>

12 USDM Data Dictionary

Note: Properties without a description in the following table are either relationships or instance identifiers and were deemed to be out of scope for terminology development. Please see Section 4.4, <u>Internal Identifiers Within the Model</u>, for additional information on the use of identifier variables in the model.

Class Name Attribute Data Type NC Card Preferr **Definition** Codel Inherited Name I Cinalit ed ist From Co Term Ref y de C42 Abbrevi A set of letters that Abbreviation 610 ation are drawn from a word or from a sequence of words and that are used for brevity in place of the full word or phrase. (CDISC Glossary) string abbreviatedTe C42 Abbrevi A set of letters that string 610 are drawn from a хt ation word or from a sequence of words and that are used for brevity in place of the full word or phrase. (CDISC Glossary) expandedText CN Abbrevi The full literal string EW representation of ation the abbreviation. Long Name 0..* A USDM notes CommentAnnot ation relationship between the Abbreviation and Comment Annotation classes which provides the set of notes related to the abbreviation. Activity C71 Study An action, 473 undertaking, or Activity event, which is anticipated to be performed or observed, or was performed or observed, according to the study protocol during the execution of the study. id string

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	name	string	C18 884 2		Study Activity Name	The literal identifier (i.e., distinctive designation) of the study activity.		
	description	string	C70 960		Study Activity Descript ion	A narrative representation of the study activity.		
	label	string	C20 745 8		Study Activity Label	The short descriptive designation for the study activity.		
	definedProced ures	Procedure		0*		A USDM relationship between the Activity and Procedure classes which identifies the set of defined procedures associated with the activity.		
	biomedicalCo ncepts	BiomedicalCon cept		0*		A USDM relationship between the Activity and BiomedicalConcept classes which identifies the set of biomedical concepts associated with the activity.		
	next	Activity		01		A USDM relationship within the Activity class which identifies the activity that follows the current activity in the display order.		
	notes	CommentAnnot ation		0*		A USDM relationship between the Activity and CommentAnnotatio n classes which provides the set of notes related to the activity.		
	timeline	ScheduleTimeli ne		01		A USDM relationship		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						between the Activity and ScheduleTimeline classes which provides the details associated with an instance of the scheduled timeline related to the		
	children	Activity		0*		activity. A USDM relationship within the Activity class which identifies the set of child activities associated with an activity.		
	previous	Activity		01		A USDM relationship within the Activity class which identifies the activity that precedes the current activity in the display order.		
	bcSurrogates	BiomedicalCon ceptSurrogate		0*		A USDM relationship between the Activity and BiomedicalConcept Surrogate classes which identifies the set of biomedical concept surrogates associated with the activity.		
	bcCategories	BiomedicalCon ceptCategory		0*		A USDM relationship between the Activity and BiomedicalConcept Category classes which identifies the set of biomedical concept categories associated with the activity.		
Address			C25 407		Address	A standardized representation of the location of a person, business,		

Class Name	Attribute Name	Data Type	Co	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
			de			building, or		
						organization. (NCI)		
	id	string				organization (1(01)		
	text	string	C20		Address	A standardized		
			131		Full	representation of		
			1		Text	the complete set of		
						components		
						denoting the		
						physical address of		
						the person, business, building,		
						or organization.		
	line	string	C25		Address	The street name		
		311118	690		Line	and number,		
						building number,		
						apartment or unit		
						number, or post		
						office box number		
						where an entity is		
	11.4.1.4		C17		District	physically located. An administrative		
	district	string	C17 622		District	or territorial		
			9			division of a city,		
						town, county,		
						parish, state,		
						country, or other		
						locality based on a		
						shared		
			G2.5		G!	characteristic.		
	city	string	C25		City	A relatively large		
			160			and/or densely populated area of		
						human habitation		
						with administrative		
						or legal status that		
						may be specified as		
						a component of a		
						postal address.		
	postalCode	string	C25		Postal	An alphanumeric		
			621		Code	code assigned to a		
	stato	atrina	C07		State	mail delivery area. A sub-division of a		
	state	string	C87 194		State	country that forms		
			194			part of a federal		
						union. States are		
						usually, but not		
						always, more		
						autonomous than		
						provinces and may		
						have different laws		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition from the central	Codel ist Ref	Inherited From
	country	Code	C25 464	01	Country	government. A sovereign nation occupying a distinct territory and ruled by an autonomous government.	(Point out to ISO 3166-1 Alpha -3 Count ry code)	
AdministrableP roduct			CN EW		Adminis trable Product	Any study product that is formulated and presented in the form that is suitable for administration to a study participant.	code	
	name	string string	CN EW		Adminis trable Product Definiti on Name	The literal identifier (i.e., distinctive designation) of the administrable product.		
	description	string	CN EW		Adminis trable Product Definiti on Descript ion	A narrative representation of the administrable product.		
	label	string	CN EW		Adminis trable Product Definiti on Label	The short descriptive designation for the administrable product.		
	administrable DoseForm	AliasCode	CN EW	1	Adminis trable Product Dose Form	The physical form in which formulated ingredient(s) are presented in the administrable product.	SDT M Termi nolog y Codeli st C6672 6	
	notes	CommentAnnot ation		0*		A USDM relationship between the		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						AdministrableProd uctn and CommentAnnotatio n classes which provides the set of notes related to the administrable product.		
	pharmacologic Class	Code	CN EW	01	Adminis trable Product Pharmac ologic Class	pharmacological class of the	(Point s to extern al codeli sts such as UNII, MED-RT)	
	identifiers	AdministrableP roductIdentifier		0*		A USDM relationship between the AdministrableProd uct and AdministrableProd uctIdentifier classes which provides the set of identifiers related to the administrable product.	KI)	
	properties	AdministrableP roductProperty		0*		A USDM relationship between the AdministrableProd uct and AdministrableProd uctProperty classes which provides the set of properties related to the administrable product.		
	ingredients	Ingredient		0*		A USDM relationship between the AdministrableProd uct and Ingredient classes which provides the set of ingredients related		

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Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
					Property Type		le Produ ct Proper ty Type	
	quantity	Quantity	CN EW	01	Adminis trable Product Property Quantity Value	The numeric value associated with an administrable product property.		
Administration			C25 409		Adminis tration	The act of dispensing, applying, or tendering a product, agent, or therapy.		
	id	string						
	name	string	C20 746 5		Adminis tration Name	The literal identifier (i.e., distinctive designation) for the administration of a product, agent, or therapy.		
	description	string	C20 746 3		Adminis tration Descript ion	A narrative representation for the administration of a product, agent, or therapy.		
	label	string	C20 746 4		Adminis tration Label	The short descriptive designation for the administration of a product, agent, or therapy.		
	administrableP roduct	AdministrableP roduct		01		A USDM relationship between the Administration and AdministrableProd uctDefinition classes which identifies the administrable product associated with the administration of the product, agent, or therapy.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	route	AliasCode	C38 114	01	Route of Adminis tration	The pathway by which a substance is administered in order to reach the site of action in the body.	SDT M Termi nolog y Codeli st C6672 9	
	dose	Quantity	C16 719 0	01	Adminis tration Dose	The value representing the amount of an agent given to an individual at one time.		
	frequency	AliasCode	C89 081	01	Dosing Frequen cy	The number of doses administered per a specific interval.	SDT M Termi nolog y Codeli st C7111	
	notes	CommentAnnot ation		0*		A USDM relationship between the Administration and CommentAnnotatio n classes which provides the set of notes related to the administration of the product, agent, or therapy.		
	duration	Administration Duration		1		A USDM relationship between the Administration and AdministrationDur ation classes which provides the duration of an instance of product, agent, or therapy administration.		
Administration Duration	id	string	C69 282		Adminis tration Duratio n			

Class Name	Attribute	Data Type	NC		Preferr	Definition	Codel	
	Name		I C- Co de	inalit y	ed Term		ist Ref	From
	description	string	C20 745 9		Adminis tration Duratio n Descript ion	A narrative representation of the agent administration duration.		
	durationWillV ary	Boolean	C20 746 1		Adminis tration Duratio n Will Vary Indicato r	An indication as to whether the agent administration duration is planned to vary within and/or across subjects.		
	reasonDuratio nWillVary	string	C20 746 2		Adminis tration Duratio n Reason Duratio n Will Vary	The explanation for why the agent administration duration will vary within and/or across subjects.		
	quantity	Quantity	C20 746 0	01	Adminis tration Duratio n Quantity Value	The value representing the amount of time over which the administration of an agent occurs.		
AliasCode	id	otring	C20 134 4		Alias Code	An alternative symbol or combination of symbols which is assigned to the members of a collection.		
	standardCode	string Code		1		A USDM relationship between the AliasCode and Code classes which provides the details of the standard code.		
	standardCode Aliases	Code		0*		A USDM relationship between the AliasCode and Code classes which identifies the set of standard code		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
			uc			aliases associated		
						with the alias code.		
AnalysisPopula			C18		Analysis	A target study		
tion			881		Populati	population on		
			4		on	which an analysis		
						is performed. These		
						may be represented		
						by the entire study		
						population, a		
						subgroup defined		
						by a particular		
						characteristic		
						measured at		
						baseline, or a		
						principal stratum		
						defined by the		
						occurrence (or non-		
						occurrence,		
						depending on		
						context) of a		
						specific		
						intercurrent event. (ICH E9 R1		
						Addendum)		
	id	string				Addendum)		
	text	string	C20		Analysis	An instance of		
	text	String	746		Populati	unstructured text		
			8		on Text	that represents the		
					on reac	analysis population.		
	name	string	C20		Analysis			
	1101110	Sumg	746		Populati	identifier (i.e.,		
			7		on	distinctive		
					Name	designation) of the		
						analysis population.		
	description	string	C18		Analysis	A narrative		
			885		Populati	representation of		
			4		on	the analysis		
					Descript	population.		
					ion			
	label	string	C20		Analysis			
			746		Populati	descriptive		
			6		on	designation for the		
					Label	analysis population.	ļ	
	notes	CommentAnnot		0*		A USDM		
		ation				relationship		
						between the		
						AnalysisPopulation		
						and CommentAnnotatio		
						n classes which		
						provides the set of		
	1		1	1	<u> </u>	provides the set of	I	L

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						notes related to the		
			~			analysis population.		
AssignedPerson			CN		Assigne	An individual		
			EW		d Person	person who is		
						allotted or		
						appointed to a		
						particular role,		
						function, or other		
			_			entity.		
	id	string						
	name	string	CN		Assigne	The literal		
			EW		d Person	identifier (i.e.,		
					Name	distinctive		
						designation) of the		
						assigned person.		
	description	string	CN		Assigne	A narrative		
			EW		d Person	representation of		
					Descript	the assigned		
					ion	person.		
	label	string	CN		Assigne	The short		
			EW		d Person	descriptive		
					Label	designation for the		
						assigned person.		
	jobTitle	string	CN		Assigne	An identifying		
			EW		d Person	designation related		
					Job	to the assigned		
					Title	person's		
						occupation.		
	organization	Organization		01		A USDM		
						relationship		
						between the		
						AssignedPerson		
						and Organization		
						classes that		
						identifies that		
						organization to		
						which the assigned		
						person belongs.		
BiomedicalCon			C20		Biomedi	A unit of		
cept			134		cal	biomedical		
			5		Concept	knowledge created		
						from a unique		
						combination of		
						characteristics that		
						include		
						implementation		
						details like		
						variables and		
						terminologies, used		
						as building blocks		
						for standardized,		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						hierarchically structured clinical research information.		
	id	string						
	name	string	C20 131 2		Biomedi cal Concept Name	The literal identifier (i.e., distinctive designation) of the biomedical concept.		
	label	string	C20 747 0		Biomedi cal Concept Label	The short descriptive designation for the biomedical concept.		
	synonyms	string	C20 131 4		Biomedi cal Concept Synony m	A word or an expression that serves as a figurative, symbolic, or exact substitute for a biomedical concept, and which has the same meaning.		
	reference	string	C20 131 3		Biomedi cal Concept Referen ce	A citation to an authoritative source		
	code	AliasCode	C20 746 9	1	Biomedi cal Concept Concept Code	A concept unique identifier assigned to a biomedical concept that points to the meaning of that biomedical concept.		
	notes	CommentAnnot ation		0*		A USDM relationship between the BiomedicalConcept and CommentAnnotatio n classes which provides the set of notes related to the biomedical concept.		
	properties	BiomedicalCon ceptProperty		0*		A USDM relationship		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
BiomedicalCon ceptCategory			C20 134 6		Biomedi cal Concept	between the BiomedicalConcept and BiomedicalConcept Property classes which identifies the set of properties associated with the biomedical concept. A grouping of biomedical concepts based on		
					Categor y	some commonality or by user defined characteristics.		
	id	string					1	
	name	string	C20 131 7		Biomedi cal Concept Categor y Name	The literal identifier (i.e., distinctive designation) of the biomedical concept category.		
	description	string	C20 131 6		Biomedi cal Concept Categor y Descript ion	A narrative representation of the biomedical concept category.		
	label	string	C20 747 1		Biomedi cal Concept Categor y Label	The short descriptive designation for the biomedical concept category.		
	code	AliasCode	C20 131 5	01	Biomedi cal Concept Categor y Code	A symbol or combination of symbols which is assigned to the biomedical concept category.		
	members	BiomedicalCon cept		0*		A USDM relationship between the BiomedicalConcept Category and BiomedicalConcept classes which identifies the set of biomedical concept members		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						associated with the biomedical concept category.		
	children	BiomedicalCon ceptCategory		0*		A USDM relationship within the BiomedicalConcept Category class which identifies the set of child categories of a biomedical concept.		
	notes	CommentAnnot ation		0*		A USDM relationship between the BiomedicalConcept and CommentAnnotation classes which provides the set of notes related to the biomedical concept category.		
BiomedicalCon ceptProperty			C20 249 3		Biomedi cal Concept Property	A characteristic from a set of characteristics used to define a biomedical concept.		
	id	string						
	name	string	C20 249 4		Biomedi cal Concept Property Name	The literal identifier (i.e., distinctive designation) of the biomedical concept property.		
	label	string	C20 747 2		Biomedi cal Concept Property Label	The short descriptive designation for the biomedical concept property.		
	isRequired	Boolean	C20 249 5		Biomedi cal Concept Property Require d Indicato r			

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name		I C-	inalit			ist	From
			Co de	y	Term		Ref	
	isEnabled	Boolean	C20		Biomedi	An indication as to		
			249		cal	whether the		
			6		Concept	biomedical concept		
					Property	property is		
					Enabled	activated for use		
					Indicato	within a given		
					r	usage context for a biomedical		
	datatype	string	C20		Biomedi	concept. The structural		
	datatype	sumg	131		cal	format of the		
			9		Concept	biomedical concept		
					Property	property response		
					Respons	value. The datatype		
					e Data	is carried in the		
					Type	attribute and		
						influences the set		
						of allowable values		
						the attribute may		
						assume. (After		
	1-	Alias Cada	C20	1	D:1:	HL7)		
	code	AliasCode	C20 131	1	Biomedi cal	A concept unique identifier assigned		
			8		Concept	to a biomedical		
			0		Property	concept property		
					Concept	that points to the		
					Code	meaning of that		
						biomedical concept		
						property.		
	responseCodes	ResponseCode		0*		A USDM		
						relationship		
						between the		
						BiomedicalConcept		
						Property and		
						ResponseCode classes which		
						identifies the set of		
						response codes		
						associated with the		
						biomedical concept		
						property.		
	notes	CommentAnnot		0*		A USDM		
		ation				relationship		
						between the		
						BiomedicalConcept		
						and CommentAnnotatio		
						n classes which		
						provides the set of		
						notes related to the		
						biomedical concept		
						property.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
BiomedicalCon ceptSurrogate			C20 759 0		Biomedi cal Concept Surrogat e	A concept that substitutes for a standard biomedical concept from the designated source.		
	id	string						
	name	string	C20 747 4		Biomedi cal Concept Surrogat e Name			
	description	string	C20 132 0		Biomedi cal Concept Surrogat e Descript ion	A narrative representation of the biomedical		
	label	string	C20 747 3		Biomedi cal Concept Surrogat e Label	descriptive designation for the		
	reference	string	C20 132 1			A citation to an authoritative source		
	notes	CommentAnnot ation		0*		A USDM relationship between the BiomedicalConcept and CommentAnnotatio n classes which provides the set of notes related to the biomedical concept surrogate.		
Characteristic			C25 447		Charact eristic	The distinguishing qualities or prominent aspects of an entity.		
	id	string						SyntaxTe mplate

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	name	string	C20 747 7		Charact eristic Name	The literal identifier (i.e., distinctive designation) of the characteristic.		SyntaxTe mplate
	description	string	C20 747 5		Charact eristic Descript ion	A narrative representation of the characteristic.		SyntaxTe mplate
	label	string	C20 747 6		Charact eristic Label	The short descriptive designation for the characteristic.		SyntaxTe mplate
	text	string	C20 747 8		Charact eristic Text	An instance of structured text that represents the characteristic.		SyntaxTe mplate
	notes	CommentAnnot ation		0*		A USDM relationship between the Characteristic and CommentAnnotatio n classes which provides the set of notes related to the characteristic.		SyntaxTe mplate
	dictionary	SyntaxTemplate Dictionary		01		A USDM relationship between the Characteristic and SyntaxTemplateDic tionary classes which provides the set of dictionary entries related to characteristics.		SyntaxTe mplate
Code			C25 162		Code	A symbol or combination of symbols which is assigned to the members of a collection.		
	id	string						
	code	string	C18 885 8		Code Value	The literal value of a code.		
	codeSystem	string	C18 885 9		Code System Name	The literal identifier (i.e., distinctive designation) of the system used to		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						assign and/or manage codes.		
	codeSystemVe rsion	string	C18 886 8		Code System Version	The version of the code system.		
	decode	string	C18 886 1		Decode	Standardized or dictionary-derived human readable text associated with a code.		
CommentAnnot ation			C44 272		Comme nt Annotati on	An explanatory or critical comment, or other in-context information (e.g., pattern, motif, link), that has been associated with data or other types of information.		
	id	string						
	text	string	CN EW		Comme nt Annotati on Text	An instance of unstructured text that represents the comment annotation.		
	codes	Code	CN EW	0*	Comme nt Annotati on Code	•		
Condition			C25 457		Conditio n	A state of being.		
	id	string	137					SyntaxTe mplate
	name	string	C20 748 3		Conditio n Name	The literal identifier (i.e., distinctive designation) of the condition.		SyntaxTe mplate
	description	string	C20 748 1		Conditio n Descript ion	representation of		SyntaxTe mplate
	label	string	C20 748 2			The short descriptive designation for the condition.		SyntaxTe mplate
	text	string	C20 748 4		Conditio n Text			SyntaxTe mplate

Class Name	Attribute	Data Type	NC I C-		Preferr	Definition	Codel	
	Name		Co de	inalit y	ed Term		ist Ref	From
						represents the		
		G		0 11		condition.		G . TD
	notes	CommentAnnot		0*		A USDM		SyntaxTe
		ation				relationship between the		mplate
						Condition and		
						CommentAnnotatio		
						n classes which		
						provides the set of		
						notes related to the		
						condition.		
	dictionary	SyntaxTemplate		01		A USDM		SyntaxTe
	dictionary	Dictionary		01		relationship		mplate
		Dictionary				between the		implate
						Condition and		
						SyntaxTemplateDic		
						tionary classes		
						which provides the		
						set of dictionary		
						entries related to		
						conditions.		
	context	Activity,		0*		A USDM		
		ScheduledActiv				relationship		
		ityInstance				between the		
						Condition and the		
						ScheduledActivityI		
						nstance or Activity		
						classes which		
						identifies the		
						scheduled activity		
						instance or activity		
						to which the		
		1				condition belongs.		
	appliesTo	Activity,		0*		A USDM		
		BiomedicalCon				relationship		
		cept,				between the		
		BiomedicalCon				Condition and the		
		ceptCategory, BiomedicalCon				Activity, Procedure,		
		ceptSurrogate,				BiomedicalConcept		
		Procedure				, BiomedicalConcept		
						Surrogate, or		
						BiomedicalConcept		
						Category classes		
						which identifies the		
						procedure, activity,		
						biomedical		
						concept,		
						biomedical concept		
						surrogate, or		
						biomedical concept		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						category that applies to the condition.		
ConditionAssig nment			C20 133 5		Conditio n Assign ment	An allotting or appointment to a condition or set of conditions that are to be met in order to make a logical decision.		
	id	string						
	condition	string	C47 953		Logical Conditio n	An assumption on which rests the validity or effect of something else.		
EligibilityCriter ion	conditionTarg	ScheduledInsta nce	C16 112	1	Study Eligibili	A USDM relationship between the ConditionAssignme nt and ScheduledInstance classes which identifies the scheduled instance associated with the condition assignment. Characteristics which are		
					ty Criterio n	necessary to allow a subject to participate in a clinical study, as outlined in the study protocol. The concept covers inclusion and exclusion criteria.		
	id	string						SyntaxTe
	name	string	C20 748 8		Study Eligibili ty Criterio n Name	The literal identifier (i.e., distinctive designation) of the study eligibility criterion.		mplate SyntaxTe mplate
	description	string	C20 748 6		Study Eligibili ty Criterio n	A narrative representation of the study eligibility criterion.		SyntaxTe mplate

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
					Descript ion			
	label	string	C20 748 7		Study Eligibili ty Criterio n Label	The short descriptive designation for the study eligibility criterion.		SyntaxTe mplate
	text	string	C20 748 5		Study Eligibili ty Criterio n Text	An instance of structured text that represents the study eligibility criterion.		SyntaxTe mplate
	notes	CommentAnnot ation		0*		A USDM relationship between the EligibilityCriterion and CommentAnnotatio n classes which provides the set of notes related to the eligibility criterion.		SyntaxTe mplate
	dictionary	SyntaxTemplate Dictionary		01		A USDM relationship between the EligibilityCriterion and SyntxTemplateDict ionary classes which provides the set of dictionary entries related to eligibility criteria.		SyntaxTe mplate
	identifier	string	C20 748 9		Study Eligibili ty Criterio n Identifie	A sequence of characters used to identify, name, or characterize the inclusion or exclusion criterion.		
	category	Code	C83 016	1	Study Eligibili ty Criterio n Categor y	A classification of the inclusion exclusion criterion.	SDT M Termi nolog y Codeli st C6679	
	next	EligibilityCriter ion		01		A USDM relationship within	,	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						the EligibilityCriterion class which identifies the eligibility criterion that follows the current eligibility criterion in the display order.		
	previous	EligibilityCriter ion		01		A USDM relationship within the EligibilityCriterion class which identifies the eligibility criterion that precedes the current eligibility criterion in the display order.		
Encounter			CN EW		Study Encount er	Any physical or virtual contact between two or more parties involved in a study, at which an assessment or activity takes place.		
	id	string						
	name	string	C17 101 0		Study Encount er Name	The literal identifier (i.e., distinctive designation) for a protocol-defined study encounter.		
	description	string	C18 883 6		Study Encount er Descript ion	A narrative representation of the protocol-		
	label	string	C20 749 0		Study Encount er Label	The short		
	environmental Settings	Code	C18 884 0	0*	Environ mental Setting	The environment/setting where the event, intervention, or finding occurred.	SDT M Termi nolog y Codeli st	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
							C1272 62	
	contactModes	Code	C18 884 1	0*	Contact Mode	The means by which an interaction occurs between the subject/participant	SDT M Termi nolog	
						and person or entity (e.g., a device).	y Codeli st C1714 45	
	type	Code	C18 883 9	1	Study Encount er Type	A characterization or classification of the study encounter.	C1887 28	
	notes	CommentAnnot ation		0*		A USDM relationship between the Encounter and CommentAnnotatio n classes which provides the set of notes related to an encounter.		
	transitionEnd Rule	TransitionRule		01		A USDM relationship between the Encounter and TransitionRule classes which provides the details associated with a transition rule used to trigger the end of an encounter.		
	next	Encounter		01		A USDM relationship within the Encounter class which identifies the encounter that chronologically follows the current encounter.		
	transitionStart Rule	TransitionRule		01		A USDM relationship between the Encounter and TransitionRule classes which provides the details associated with a		

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name		I C-		ed		ist	From
			Co de	y	Term		Ref	
			uc			transition rule used		
						to trigger the start		
						of an encounter.		
	scheduledAt	Timing		01		A USDM		
						relationship		
						between the		
						Encounter and		
						Timing classes		
						which provides		
						information related		
						to the scheduled		
						timing of an		
		Enganatas		01		encounter. A USDM		
	previous	Encounter		01				
						relationship within the Encounter class		
						which identifies the		
						encounter that		
						chronologically		
						precedes the		
						current encounter.		
Endpoint			C25		Study	A defined variable		
Ziiopoiiit			212		Endpoin			
					t	an outcome of		
						interest that is		
						statistically		
						analyzed to address		
						a particular		
						research question.		
						NOTE: A precise		
						definition of an		
						endpoint typically		
						specifies the type		
						of assessments		
						made, the timing of		
						those assessments, the assessment		
						tools used, and		
						possibly other		
						details, as		
						applicable, such as		
						how multiple		
						assessments within		
						an individual are to		
						be combined. After		
						BEST Resource		
						(CDISC Glossary)		
	id	string						SyntaxTe
		1	Gac		G. 1	TTI 1'. 1		mplate
	name	string	C20		Study	The literal		SyntaxTe
			749		Endpoin			mplate
			2]	t Name	distinctive		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						designation) of the study endpoint.		
	description	string	C18 882 4		Study Endpoin t Descript ion	A narrative representation of the study endpoint.		SyntaxTe mplate
	label	string	C20 749 1		Study Endpoin t Label	The short descriptive designation for the study endpoint.		SyntaxTe mplate
	text	string	C20 749 3		Study Endpoin t Text	An instance of structured text that represents the study endpoint.		SyntaxTe mplate
	notes	CommentAnnot ation		0*		A USDM relationship between the Endpoint and CommentAnnotatio n classes which provides the set of notes related to the study endpoint.		SyntaxTe mplate
	dictionary	SyntaxTemplate Dictionary		01		A USDM relationship between the Endpoint and Syntax TemplateDic tionary classes which provides the set of dictionary entries related to study endpoints.		SyntaxTe mplate
	purpose	string	C18 882 5		Study Endpoin t Purpose Descript ion	The textual		
	level	Code	C18 882 6	1	Study Endpoin t Level	A characterization or classification of the study endpoint that determines its category of importance relative to other study endpoints.	C1887 26	
Estimand			C18 881 3		Estiman d	A precise description of the treatment effect		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						reflecting the clinical question posed by a given clinical trial objective. It summarises at a population level what the outcomes would be in the same patients under different treatment conditions being compared. (ICH E9 R1 Addendum)		
	id	string				,		
	summaryMeas ure	string	C18 885 3		Populati on- Level Summar y	A synopsis of the clinical endpoint of interest within the analysis target study population.		
	name	string	CN EW		Estiman d Name	The literal identifier (i.e., distinctive designation) of the estimand.		
	description	string	CN EW		Estiman d Descript ion	A narrative representation of the estimand.		
	label	string	CN EW		Estiman d Label	The short descriptive designation for the estimand.		
	analysisPopula tion	AnalysisPopula tion		1		A USDM relationship between the Estimand and AnalysisPopulation classes which provides the details associated with an instance of the analysis population used to partially define a study estimand.		
	notes	CommentAnnot ation		0*		A USDM relationship between the Estimand and CommentAnnotatio		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						n classes which provides the set of notes related to a study estimand.		
	variableOfInte	Endpoint		1		A USDM relationship between the Estimand and Endpoint classes which provides the details associated with an instance of the variable of interest within a study endpoint used to partially define a study estimand.		
	intercurrentEv ents	IntercurrentEve nt		1*		A USDM relationship between the Estimand and IntercurrentEvent classes which identifies the set of intercurrent events associated with a study estimand.		
	intervention	StudyInterventi		1		A USDM relationship between the Estimand and StudyIntervention classes which provides the details associated with an instance of the intervention used to partially define a study estimand.		
GeographicSco pe			C20 759 1		Geograp hic Scope	The extent or range related to the physical location of an entity.		
	id code	string AliasCode	C20 749 4	01	Geograp hic Scope Code	A symbol or combination of symbols which is assigned to the geographic scope.	(Point out to extern al dictio naries: Standa	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
							rd code is ISO-3166; Alias codes drawn from GENC, UN Regio n Codes , etc.)	
	type	Code	C20 749 5	1	Geograp hic Scope Type	A characterization or classification of the geographic scope.	C2074 12	
GovernanceDat e			C20 759 5		Study Governa nce Date	Any of the dates		
	id	string						
	name	string	C20 749 9		Study Governa nce Date Name	The literal identifier (i.e., distinctive designation) of the study governance date		
	description	string	C20 749 7		Study Governa nce Date Descript ion	A narrative representation of the study governance date.		
	label	string	C20 749 8		Study Governa nce Date Label	The short descriptive designation for the study governance date.		
	dateValue	Date	C20 750 0		Study Governa nce Date Value	The information		

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name		I C-	inalit	ed		ist	From
			Co de	y	Term		Ref	
	type	Code	C20	1	Study	A characterization	C2074	
			749		Governa	or classification of	13	
			6		nce	the study		
					Date	governance date.		
					Type			
	geographicSco	GeographicSco		1*		A USDM		
	pes	pe				relationship		
						between the		
						GovernanceDate		
						and		
						GeographicScope classes which		
						identifies the set of		
						geographic scopes		
						associated with the		
						governance date.		
Identifier			C25		Identifie	One or more		
Identifier			364		r	characters used to		
			30.		1	identify, name, or		
						characterize the		
						nature, properties,		
						or contents of a		
						thing.		
	id	string						
	text	string	CN		Identifie	An instance of		
			EW		r Text	structured text that		
						represents the		
						identifier.		
	scope	Organization		1		A USDM		
						relationship		
						between the		
						Identifier and		
						Organization classes which		
						provides the details		
						associated with		
						each organization		
						that has assigned		
						the identifier.		
Indication			C41		Disease/	The disease or		
			184		Conditio	condition the		
					n	intervention will	1	
					Indicati	diagnose, treat,		
					on	prevent, cure, or	1	
						mitigate.		
	id	string						
	name	string	C20		Disease/	The literal	1	
			750		Conditio		1	
			3		n	distinctive	1	
]		Indicati	designation) of the	1	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
					on	disease/condition		
	description	string	C11 203 8		Name Disease/ Conditio n Indicati	A narrative representation of the condition, disease or disorder		
					on Descript ion	that the clinical		
	label	string	C20 750 2		Disease/ Conditio n Indicati on Label	The short descriptive designation for the disease/condition indication.		
	isRareDisease	Boolean	C20 750 1		Disease/ Conditio n Indicati on Is Rare Disease Indicato r			
	codes	Code	C18 882 2		Disease/ Conditio n Indicati on Code	represents the disease/condition indication.	(Point out to multip le Biome dical coding dictio naries such as SNO MED CT (for FDA), MedD RA, NCIt, ICD's, etc.)	
	notes	CommentAnnot ation		0*		A USDM relationship between the Indication and CommentAnnotatio n classes which		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						provides the set of notes related to the disease/condition indication.		
Ingredient			C51 981		Ingredie nt	Any component that constitutes a part of a compounded substance or mixture.		
	id	string						
	role	Code	CN EW	1	Ingredie nt Role	The intended use of the ingredient within the context of the compounded substance or mixture.	(Point to FHIR value set: Ingred ient Role)	
	substance	Substance		1		A USDM relationship between the Ingredient and Substance classes that identifies the substance associated with the ingredient.		
IntercurrentEve	id	string	C18 881 5		Intercurr ent Event	An event(s) occurring after treatment initiation that affects either the interpretation or the existence of the measurements associated with the clinical question of interest. (ICH E9 Addendum on Estimands)		
	name	string	C18 885 5		Intercurr ent Event Name	The literal identifier (i.e., distinctive designation) of the intercurrent event.		
	description	string	C18 885 6		Intercurr ent Event Descript ion	A narrative representation of the intercurrent event.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	label	string	C20 750 4		Intercurr ent Event Label	The short descriptive designation for the intercurrent event.		
	strategy	string	C18 885 7		Intercurr ent Event Strategy	A textual description of the planned strategy to manage and/or mitigate intercurrent events.		
	notes	CommentAnnot ation		0*		A USDM relationship between the IntercurrentEvent and CommentAnnotation classes which provides the set of notes related to the intercurrent event.		
Masking			C19 127 8		Masking	The mechanism used to obscure the distinctive characteristics of the study intervention or procedure to make it indistinguishable from a comparator. (CDISC Glossary)		
	id description	string string	C20 750 5		Masking Descript ion	A narrative representation of the study masking strategy, based on a person's role within the study.		
			C20 759 2		Narrativ e Content	The container that holds an instance of unstructured text and which may include objects such as tables, figures, and images.		
	id name	string string	C20 750 7		Narrativ e Content Name	The literal identifier (i.e., distinctive designation) of the narrative content.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	sectionNumbe r	string	C20 750 9		Narrativ e Content Section Number	The numeric identifier assigned to a particular document section containing narrative content.		
	sectionTitle	string	C20 751 0		Narrativ e Content Section Title	An identifying designation for the document section containing narrative content.		
	displaySection Title	Boolean	CN EW		Narrativ e Content Section Title Display Indicato r	An indication as to whether the section title is to be displayed in the document containing narrative content.		
	displaySection Number	Boolean	CN EW		Narrativ e Content Section Number Display Indicato r	An indication as to whether the section number is to be displayed in the document containing narrative content.		
	contentItem	NarrativeConte ntItem		01		A USDM relationship between the NarrativeContent and NarrativeContentIte m classes which identifies the content item associated with the narrative content.		
	previous	NarrativeConte nt		01		A USDM relationship within the NarrativeContent class which identifies the narrative content that precedes the current narrative content in the display order.		
	next	NarrativeConte nt		01		A USDM relationship within		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						the NarrativeContent class which identifies the narrative content that follows the current narrative content in the display order.		
	children	NarrativeConte nt		0*		A USDM relationship within the NarrativeContent class which identifies the set of child content associated with an instance of narrative content.		
NarrativeConte ntItem			CN EW		Narrativ e Content Item	An individual item within the container that holds an instance of unstructured text and which may include objects such as tables, figures, and images.		
	id	string				images.		
	name	string	CN EW		Narrativ e Content Item Name	The literal identifier (i.e., distinctive designation) of the narrative content item.		
	text	string	CN EW		Narrativ e Content Item Text	An instance of unstructured text that represents the narrative content item.		
Objective	:4	atrica	C14 245 0		Study Objectiv e	The reason for performing a study in terms of the scientific questions to be answered by the analysis of data collected during the study.		Cunto
	id	string						SyntaxTe mplate

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	name	string	C20 751 2		Study Objectiv e Name	The literal identifier (i.e., distinctive designation) of the study objective.		SyntaxTe mplate
	description	string	C94 090		Study Objectiv e Descript ion	A narrative representation of the study objective. (BRIDG)		SyntaxTe mplate
	label	string	C20 751 1		Study Objectiv e Label	The short descriptive designation for the study objective.		SyntaxTe mplate
	text	string	C20 751 3		Study Objectiv e Text	An instance of structured text that represents the study objective.		SyntaxTe mplate
	notes	CommentAnnot ation		0*		A USDM relationship between the Objective and CommentAnnotatio n classes which provides the set of notes related to the study objective.		SyntaxTe mplate
	dictionary	SyntaxTemplate Dictionary		01		A USDM relationship between the Objective and SyntaxTemplateDic tionary classes which provides the set of dictionary entries related to study objectives.		SyntaxTe mplate
	level	Code	C18 882 3	1	Study Objectiv e Level	A characterization or classification of the study objective that determines its category of importance relative to other study objectives.	C1887 25	
	endpoints	Endpoint		0*		A USDM relationship between the Objective and Endpoint classes which identifies the		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						set of endpoints associated with the study objective.		
Organization			C19 711		Organiz ation	A formalized group of persons or other organizations collected together for a common purpose (such as administrative, legal, political) and the infrastructure to carry out that purpose. (BRIDG)		
	id	string						
	name	string	C93 874		Organiz ation Name	The literal identifier (i.e., distinctive designation) of the organization.		
	label	string	C20 751 4		Organiz ation Label	The short descriptive designation for the organization.		
	identifier	string	C93 401		Organiz ation Identifie r	A unique symbol that establishes identity of the organization. (BRIDG)		
	identifierSche me	string	C18 881 9		Identifie r Provider Organiz ation Name	The name of the organization that provides the identifier for the entity.		
	legalAddress	Address		01		A USDM relationship between the Organization and Address classes which provides the legal address for an organization.		
	type	Code	C18 882 0	1	Organiz ation Type	A characterization or classification of the formalized group of persons or other organizations collected together for a common purpose (such as	C1887 24	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						administrative, legal, political) and the infrastructure to carry out that purpose.		
	managedSites	StudySite		0*		A USDM relationship between the Organization and StudySite classes which identifies the set of study sites managed by the organization.		
ParameterMap			C20 745 6		Paramet er Map	The paired name and value for a given parameter.		
	id	string						
	tag	string	C20 751 5		Program ming Tag	Character strings bounded by angle brackets that act as containers for programming language elements.		
	reference	string	C20 751 6		Program ming Tag Referen ce			
PopulationDefi nition			C20 759 3		Populati on Definiti on	A concise explanation of the meaning of a population.		
	id	string						
	name	string	C20 752 0		Populati on Definiti on Name	The literal identifier (i.e., distinctive designation) of the population definition.		
	description	string	C20 751 7		Populati on Definiti on Descript ion	A narrative representation of the population definition.		

Class Name	Attribute	Data Type	NC		Preferr	Definition		Inherited
	Name		I C- Co de	inalit y	ed Term		ist Ref	From
	label	string	C20 751 9		Populati on Definiti on Label	The short descriptive designation for the population definition.		
	includesHealth ySubjects	Boolean	C20 751 8		Populati on Definiti on Includes Healthy Subjects Indicato r	An indication as to whether the population definition includes healthy subjects, that is, subjects without the disease or condition under study.		
	plannedSex	Code	C20 752 3	02	Populati on Definiti on Planned Sex	The protocoldefined sex within the population definition.	SDT M Termi nolog y Codeli st C6673 2	
	notes	CommentAnnot ation		0*		A USDM relationship between the PopulationDefinition and CommentAnnotation classes which provides the set of notes related to the population definition.		
	criteria	EligibilityCriter ion		0*		A USDM relationship between the PopulationDefinitio n and EligibilityCriterion classes which identifies the set of eligibility criteria associated with the population definition.		
	plannedAge	Range	C20 770 1	01	Populati on Definiti on Planned Age	The anticipated age of subjects within the population definition.		

Class Name	Attribute	Data Type	NC		Preferr	Definition		Inherited
	Name		I C-	inalit	ed		ist	From
			Co de	y	Term		Ref	
	plannedEnroll	Range	C20	01	Populati			
	mentNumber		752		on	representing the		
			2		Definiti	planned number of		
					on	subjects to be		
					Planned	entered in a clinical		
					Enrollm	trial, within the		
					ent Number	population definition.		
	plannedCompl	Range	C20	01	Populati			
	etionNumber	runge	752	01	on	representing the		
			1		Definiti	planned number of		
					on	subjects that must		
					Planned	complete the study		
					Complet			
					ion	objectives and		
					Number	endpoints of the		
						study, within the		
						population definition.		
Procedure			C98		Procedu	Any activity		
Tioccdure			769		re	performed by		
			100			manual and/or		
						instrumental means		
						for the purpose of		
						diagnosis,		
						assessment,		
						therapy,		
						prevention, or		
	id	string				palliative care.		
	name	string	C20		Procedu	The literal		
	Tidillo	Sums	132		re Name			
			5			distinctive		
						designation) of the		
						procedure.		
	description	string	C20		Procedu	A narrative		
			132		re	representation of		
			4		Descript	the procedure.		
	label	string	C20		ion Procedu	The short		
	14001	sumg	752		re Label	descriptive		
			4		Lucci	designation for the		
						procedure.		
	procedureTyp	string	C18		Procedu	A characterization		
	e		884		re Type	or classification of		
			8			the study		
			1			procedure.		
	code	Code	C15	1	Procedu	A symbol or	(Point	
			462		re Code	combination of	out to	
			6			symbols which is	extern	
L							al	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						assigned to medical procedure.	dictio nary like CPT, MedD RA, SNO MED CT, etc.)	
	notes	CommentAnnot ation		0*		A USDM relationship between the Procedure and CommentAnnotatio n classes which provides the set of notes related to a procedure.		
	studyIntervent	StudyInterventi on		01		A USDM relationship between the Procedure and StudyInterventioncl asses which provides the details associated with an instance of an intervention performed during the conduct of a procedure.		
Quantity	id		C25 256		Quantity			
	value	string Float	C25 712		Quantity Value	A numerical quantity measured or assigned or computed.		
	unit	AliasCode	C44 258	01	Quantity Unit		SDT M Termi nolog y Codeli st C7162 0	

Class Name	Attribute Name	Data Type	NC I C- Co	Card inalit	Preferr ed Term	Definition	Codel ist Ref	Inherited From
			de	J			ICI	
Range			C38 013		Range	The difference between the lowest and highest numerical values; the limits or scale of variation.		
	id	string				or variation.		
	minValue	Float	C25 570		Minimu m Value	The smallest value in quantity or degree in a set of values.		
	maxValue	Float	C25 564		Maximu m Value	The largest value in quantity or degree in a set of values.		
	isApproximate	Boolean	C20 752 5		Value Range is Approxi mate Indicato	An indication as to whether the value range is almost, but not quite, exact.		
	unit	Code	C25 709	01	Unit of Measure	A named quantity in terms of which other quantities are measured or specified, used as a standard measurement of like kinds.	SDT M Termi nolog y Codeli st C7162 0	
ReferenceIdenti fier			CN EW		Referen ce Identifie r	A sequence of characters used to identify, name, or characterize the reference.		
	id	string						Identifier
	text	string	CN EW		Referen ce Identifie r Text	An instance of structured text that represents the reference identifier.		Identifier
	scope	Organization		1		A USDM relationship between the ReferenceIdentifier and Organization classes which provides the details associated with each organization that has assigned the reference identifier.		Identifier

Class Name	Attribute Name	Data Type	NC I C- Co de	y	Preferr ed Term	Definition	ist Ref	Inherited From
	type	Code	CN EW	1	Referen ce Identifie r Type	A characterization or classification of the reference identifier.	CNE W Refere nce Identif ier Type	
ResponseCode			C20 134 7		Respons e Code	A symbol or combination of symbols representing the response to the question.		
	isEnabled	string Boolean	C20 133 0		Respons e Code Enabled Indicato	An indication as to whether the response code is activated for use within a given usage context.		
	code	Code	C25 162	1	Code	A symbol or combination of symbols which is assigned to the members of a collection.		
ScheduleTimeli ne			C20 134 8		Schedul e Timelin e	A chronological schedule of planned temporal events.		
	id	string						
	name	string	C20 133 4		Schedul e Timelin e Name	The literal identifier (i.e., distinctive designation) of the schedule timeline.		
	description	string	C20 133 2		Schedul e Timelin e Descript ion	A narrative representation of the schedule timeline.		
	label	string	C20 753 0		Schedul e Timelin e Label	The short descriptive designation for the schedule timeline.		
	entryConditio n	string	C20 133 3		Schedul e Timelin e Entry	A logical evaluation on which rests the validity of entry		

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name	J.F.	I C-		ed		ist	From
			Co de	y	Term		Ref	
			uc		Conditio	into a schedule		
					n	timeline.		
	mainTimeline	Boolean	C20		Main	An indication as to		
			133		Timelin	whether the		
			1		e	timeline or timeline		
					Indicato	component is part		
					r	of the central or		
		~				principal timeline.		
	instances	ScheduledInsta		0*		A USDM		
		nce				relationship		
						between the ScheduleTimeline		
						and		
						ScheduledInstance		
						classes which		
						identifies the set of		
						scheduled instances		
						(e.g., scheduled		
						activity instances or		
						scheduled decision		
						instances)		
						associated with the		
						scheduled timeline.		
	entry	ScheduledInsta		1		A USDM		
		nce				relationship		
						between the		
						ScheduleTimeline		
						and		
						ScheduledInstance		
						classes which		
						defines the entry		
						into a scheduled		
						instance (e.g., scheduled activity		
						instances or		
						scheduled decision		
						instances) for a		
						timeline.		
	exits	ScheduleTimeli		0*		A USDM		
		neExit				relationship		
						between the		
						ScheduleTimeline		
						and		
						ScheduleTimelineE		
						xit classes which		
						identifies the set of		
						exits from the		
		TD: :	-	0 **		scheduled timeline.		
	timings	Timing		0*		A USDM		
						relationship		
						between the ScheduleTimeline		
		<u>I</u>	<u> </u>	<u> </u>		Schedule I imenne		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						and Timing classes which identifies the set of timings associated with the scheduled timeline.		
ScheduleTimeli neExit			C20 134 9		Schedul e Timelin e Exit	To go out of or leave the schedule timeline.		
ScheduledActiv ityInstance	id	string	C20 135 0		Schedul ed Activity Instance	A scheduled occurrence of an activity event.		
	id	string						Schedule dInstance
	name	string	C20 753 3		Schedul ed Activity Instance Name	The literal identifier (i.e., distinctive designation) of the scheduled activity instance.		Schedule dInstance
	description	string	C20 753 1		Schedul ed Activity Instance Descript ion	A narrative representation of the scheduled activity instance.		Schedule dInstance
	label	string	C20 753 2		Schedul ed Activity Instance Label	The short descriptive designation for the scheduled activity instance.		Schedule dInstance
	defaultConditi on	ScheduledInsta nce		01		A USDM relationship within the ScheduledActivityI nstance class which identifies the default condition within a scheduled activity instance.		Schedule dInstance
	epoch	StudyEpoch		01		A USDM relationship between the ScheduledActivityI nstance and StudyEpoch classes which identifies the study epoch associated with a		Schedule dInstance

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						scheduled activity		
	activities	Activity		0*		instance. A USDM relationship between the ScheduledActivityI nstance and		
						Activity classes which identifies the set of activities associated with a scheduled activity		
						instance.		
	encounter	Encounter		01		A USDM relationship between the ScheduledActivityI nstance and Encounter classes which defines the subject encounter associated with the ScheduleActivityIn stance.		
	timeline	ScheduleTimeli		01		A USDM relationship between the ScheduledActivityI nstance and ScheduleTimeline classes which provides the details associated with an instance of a scheduled timeline related to a scheduled activity instance.		
	timelineExit	ScheduleTimeli neExit		01		A USDM relationship between the ScheduledActivityI nstance and ScheduleTimelineE xit classes which provides the details associated with the exit from a timeline related to a scheduled activity instance.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
ScheduledDecis ionInstance			C20 135 1		Schedul ed Decisio n Instance	A scheduled occurrence of a decision event.		
	id	string						Schedule dInstance
	name	string	C20 753 6		Schedul ed Decisio n Instance Name	The literal identifier (i.e., distinctive designation) of the scheduled Decision instance.		Schedule dInstance
	description	string	C20 753 4		Schedul ed Decisio n Instance Descript ion	A narrative representation of the scheduled Decision instance.		Schedule dInstance
	label	string	C20 753 5		Schedul ed Decisio n Instance Label	The short descriptive designation for the scheduled Decision instance.		Schedule dInstance
	defaultConditi on	ScheduledInsta nce		01		A USDM relationship within the ScheduledDecision Instance class which identifies the default condition within a scheduled decision instance.		Schedule dInstance
	epoch	StudyEpoch		01		A USDM relationship between the ScheduledDecision Instance and StudyEpoch classes which identifies the study epoch associated with a scheduled decision		Schedule dInstance
	conditionAssi gnments	ConditionAssig nment		1*		instance. A USDM relationship between the ScheduledDecision Instance and		

Class Name	Attribute Name	Data Type	NC I C-	inalit	Preferr ed	Definition	Codel	Inherited From
			Co de	y	Term		Ref	
						ConditionAssignme		
						nt classes which		
						identifies the set of		
						condition		
						assignments associated with a		
						scheduled decision		
						instance.		
ScheduledInsta			C20		Schedul	A scheduled		
nce			129		ed	occurrence of a		
			9		Instance	temporal event.		
	id	string						
	name	string	C20		Schedul	The literal		
			745		ed	identifier (i.e.,		
			5		Instance Name	distinctive		
					Name	designation) of the scheduled instance.		
	description	string	C20		Schedul	A narrative		
	description	Sums	745		ed	representation of		
			3		Instance	the scheduled		
					Descript	instance.		
					ion			
	label	string	C20		Schedul	The short		
			745		ed	descriptive		
			4		Instance	designation for the		
	defaultConditi	ScheduledInsta		01	Label	scheduled instance. A USDM		
	on	nce		01		relationship within		
						the		
						ScheduledInstance		
						class which		
						identifies the		
						default condition		
						within a scheduled		
	epoch	StudyEpoch	1	01		instance. A USDM		
	СРОСП	Diddy Epoch		01		relationship		
						between the		
						ScheduledInstance		
						and StudyEpoch		
						classes which		
						identifies the study		
						epoch associated with a scheduled		
						instance.		
Strength			CN		Substan	The content of an		
Sacingui			EW		ce	substance		
					Strength			
						quantitatively per		
						dosage unit, per		
						unit of volume, or		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						per unit of weight, according to the pharmaceutical dose form of the product.		
	id	string	G) I		0.1			
	name	string	CN EW		Substan ce Strength Name	The literal identifier (i.e., distinctive designation) of the substance strength.		
	description	string	CN EW		Substan ce Strength Descript ion	A narrative representation of the substance strength.		
	label	string	CN EW		Substan ce Strength Label	The short descriptive designation for the substance strength.		
	numerator	Quantity, Range		01		A USDM relationship between the Strength and the Quantity and Range classes that identifies the numerator's value or range of values associated with the substance strength.		
	denominator	Quantity		01		A USDM relationship between the Strength and Quantity classes that identifies the denominator associated with the substance strength.		
Study			C15 206		Clinical Study	A clinical study involves research using human volunteers (also called subjects or participants) that is intended to add to medical knowledge. There are two main types of clinical studies:		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						clinical trials (also called interventional studies) and observational studies. [http://ClinicalTrials.gov](CDI SC Glossary)		
	id	string						
	name	string	C68 631		Clinical Study Name	The literal identifier (i.e., distinctive designation) of the clinical study.		
	description	string	C14 270 4		Clinical Study Descript ion	A narrative representation of the clinical study.		
	label	string	C20 747 9		Clinical Study Label	The short descriptive designation for the clinical study.		
	versions	StudyVersion		0*		A USDM relationship between the Study and Study Version classes which identifies the set of versions associated with the study.		
	documentedB y	StudyDefinition Document		0*		A USDM relationship between the Study and StudyDefinitionDo cument classes signifying that the study is documented in a study definition document.		
StudyAmendme nt			C20 759 4		Study Amend ment	A written description of a change(s) to, or formal clarification of, a study.		
	id	string						
	number	string	C20 753 7		Study Amend ment Number	A string of numerals that uniquely identifies		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						a protocol		
	summary	string	C11 562 7		Study Amend ment Summar y	amendment. A short narrative representation describing the changes introduced in the current		
						version of the protocol.		
	substantialImp act	Boolean	C20 753 8		Study Amend ment Substant ial Impact Indicato r	An indication as to whether the amendment is likely to have a substantial impact on the safety or rights of study subjects/participant s.		
	enrollments	SubjectEnrollm ent		1*		A USDM relationship between the StudyAmendment and SubjectEnrollment classes which provides the set of subject enrollments associated with the study amendment.		
	secondaryReas ons	StudyAmendme ntReason		0*		A USDM relationship between the StudyAmendment and StudyAmendmentR eason classes which identifies the set of secondary reasons for issuing the study amendment.		
	previous	StudyAmendme nt		01		A USDM relationship within the StudyAmendment class which identifies the study amendment that chronologically precedes the current study amendment.		

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name		I C-	inalit	ed		ist	From
			Co de	y	Term		Ref	
	primaryReaso	StudyAmendme		1		A USDM		
	n	ntReason				relationship		
						between the		
						StudyAmendment		
						and		
						StudyAmendmentR eason classes which		
						identifies the		
						primary reason for		
						issuing the study		
						amendment.		
StudyAmendme			C20		Study	The rationale for		
ntReason			745		Amend	the change(s) to, or		
			7		ment	formal clarification		
					Reason	of, a protocol.		
	id	string						
	otherReason	string	C20		Other	The rationale for		
			753		Reason	the change(s) to, or		
			9		for	formal clarification		
					Study	of, a protocol that		
					Amend	is not otherwise		
	_		~-		ment	specified.	~	
	code	Code	C20	1	Study	A symbol or	C2074	
			754		Amend	combination of	15	
			0		ment Reason	symbols which is		
					Code	assigned to the study amendment		
					Code	reason.		
StudyArm			C17		Study	A planned pathway		
Study/Mili			444		Arm	assigned to the		
			7			subject as they		
						progress through		
						the study, usually		
						referred to by a		
						name that reflects		
						one or more		
						treatments,		
						exposures, and/or		
						controls included in		
	: 4	atuin a				the path.		
	id	string	C17		Ctuder	The literal		
	name	string	098		Study Arm	identifier (i.e.,		
			4		Name	distinctive		
			'		1 (4111)	designation) of the		
						study arm.		
	description	string	C93		Study	A narrative		
	F	5	728		Arm	representation of		
					Descript	the study arm.		
					ion			

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	label	string	C17 245 6		Study Arm Label	The short descriptive designation for the study arm.		
	dataOriginDes cription	string	C18 882 8		Study Arm Data Origin Descript ion	The textual representation of the study arm data origin.		
	dataOriginTyp e	Code	C18 882 9	1	Study Arm Data Origin Type	A characterization or classification of the study arm with respect to where the study arm data originates.	C1887 27	
	type	Code	C18 882 7	1	Study Arm Type	A characterization or classification of the study arm.	Protoc ol Termi nolog y Codeli st C1742 22	
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyArm and CommentAnnotatio n classes which provides the set of notes related to the study arm.		
	populations	PopulationDefi nition		0*		A USDM relationship between the StudyArm and PopulationDefinition classes which identifies the set of populations associated with the study arm.		
StudyCell			C18 881 0		Study Design Cell	A partitioning of a study arm into individual pieces, which are associated with an epoch and any number of		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						sequential elements within that epoch.		
	id	string		_				
	arm	StudyArm		1		A USDM relationship between the StudyCell and StudyArm classes which identifies the study arm associated with a study cell.		
	epoch	StudyEpoch		1		A USDM relationship between the StudyCell and StudyEpoch classes which identifies the study epoch associated with a study cell.		
	elements	StudyElement		0*		A USDM relationship between the StudyCell and StudyElement classes which identifies the set of study elements associated with the study cell.		
StudyCohort			C61 512		Study Cohort	A group of individuals who share a set of characteristics (e.g., exposures, experiences, attributes), which logically defines a population under study.		
	id	string				study.		Populatio nDefiniti on
	name	string	C20 754 4		Study Cohort Name	The literal identifier (i.e., distinctive designation) of the study cohort.		Populatio nDefiniti on

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	description	string	C20 754 2		Study Cohort Descript ion	A narrative representation of the study cohort.		Populatio nDefiniti on
	label	string	C20 754 3		Study Cohort Label	The short descriptive designation for the study cohort.		Populatio nDefiniti on
	includesHealth ySubjects	Boolean	C20 748 0		Study Cohort Includes Healthy Subjects Indicato	An indication as to whether the study cohort includes healthy subjects, that is, subjects without the disease or condition under study.		Populatio nDefiniti on
	plannedSex	Code	C20 754 1	02	Study Cohort Planned Sex	The protocoldefined sex within the study cohort.	SDT M Termi nolog y Codeli st C6673	Populatio nDefiniti on
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyCohort and CommentAnnotatio n classes which provides the set of notes related to the study cohort.		Populatio nDefiniti on
	criteria	EligibilityCriter ion		0*		A USDM relationship between the StudyCohort and EligibilityCriterion classes which identifies the set of eligibility criteria associated with the study cohort.		Populatio nDefiniti on
	plannedAge	Range	C20 754 5	01	Study Cohort Planned Age	The anticipated age of subjects within the study cohort.		Populatio nDefiniti on
	plannedEnroll mentNumber	Range	C20 770 2	01	Study Cohort Planned	The value representing the planned number of		Populatio nDefiniti on

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
					Enrollm ent Number	subjects to be entered in a clinical trial, within the study cohort.		
	plannedCompl etionNumber	Range	C20 754 6	01	Study Cohort Planned Complet ion Number	The value representing the planned number of subjects that must complete the study in order to meet the objectives and endpoints of the study, within the study cohort.		Populatio nDefiniti on
	characteristics	Characteristic		0*		A USDM relationship between the StudyCohort and Characteristic classes which identifies the set of subject characteristics associated with the study cohort.		
StudyDefinition Document			CN EW		Study Definiti on Docume nt	Any physical or electronic document that is related to defining a study or part of a study.		
	id name	string string	CN		Study	The literal		
			EW		Definiti on Docume nt Name	identifier (i.e., distinctive designation) of the study definition document.		
	description	string	CN EW		Study Definiti on Docume nt Descript ion	A narrative representation of the study definition document.		
	label	string	CN EW		Study Definiti on Docume nt Label	The short descriptive designation for the study definition document.		

Class Name	Attribute	Data Type	NC		Preferr	Definition		Inherited
	Name		I C-	inalit	ed		ist	From
			Co de	y	Term		Ref	
	templateName	string	CN		Study	The literal		
			EW		Definiti	identifier (i.e.,		
					on	distinctive		
					Docume	designation) of the		
					nt	study definition		
					Templat e Name	document template.		
	language	Code	CN	1	Study	The language in	(Point	
			EW		Definiti	which the study	out to	
					on	definition	ISO	
					Docume	document is	639	
					nt	written.	langua	
					Languag e		ge value	
							list)	
	type	Code	CN	1	Study	A characterization	CNE	
	C) PC	Code	EW	1	Definiti	or classification of	W	
					on	the study definition	Study	
					Docume	document.	Defini	
					nt Type		tion	
							Docu	
							ment	
							Type	
	notes	CommentAnnot		0*		A USDM		
		ation				relationship		
						between the		
						StudyDefinitionDo		
						cument and CommentAnnotatio		
						n classes which		
						provides the set of		
						notes related to the		
						study definition		
						document.		
	versions	StudyDefinition		0*		A USDM		
		DocumentVersi				relationship		
		on				between the		
						StudyDefinitionDo		
						cument and		
						StudyDefinitionDo		
						cumentVersion		
						classes which		
						identifies the set of versions associated		
						with the study		
						definition		
						document.		
StudyDefinition			CN		Study	A representation of		
DocumentVersi			EW		Definiti	a particular edition		
on					on	or snapshot of the		
					Docume	study definition		
						document as it		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
					nt Version	exists at a particular point in time.		
	id	string						
	version	string	CN EW		Study Definiti on Docume nt Version	A representation of a particular edition or snapshot of the study definition document as it exists at a particular point in time.		
	status	Code	CN EW	1	Study Definiti on Docume nt Status	A condition of the study definition document at a point in time with respect to its state of readiness for implementation.	C1887 23	
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyDefinitionDo cumentVersion and CommentAnnotatio n classes which provides the set of notes related to the study definition document version.		
	dateValues	GovernanceDat e		0*		A USDM relationship between the StudyDefinitionDo cumentVersion and GovernanceDate classes which provides the set of governance dates associated with the study definition document version.		
	contents	NarrativeConte nt		0*		A USDM relationship between the StudyDefinitionDo cumentVersion and NarrativeContent classes which identifies the set of narrative content		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						associated with the version of the study definition document.		
	children	StudyDefinition DocumentVersi on		0*		A USDM relationship within the StudyDefinitionDo cumentVersion class which identifies the set of child documents of a study definition document version.		
StudyDesign			C15 320		Study Design	A plan detailing how a study will be performed in order to represent the phenomenon under examination, to answer the research questions that have been asked, and informing the statistical approach.		
	id	string				• •		
	name	string	C20 133 8		Study Design Name	The literal identifier (i.e., distinctive designation) of the study design.		
	description	string	C14 713 9		Study Design Descript ion	A narrative representation of the study design.		
	label	string	C20 754 8		Study Design Label	The short descriptive designation for the study design.		
	rationale	string	C14 270 5		Study Design Rational e	Reason(s) for choosing the study design. This may include reasons for the choice of control or comparator, as well as the scientific rationale for the study design.		
	activities	Activity		0*		A USDM relationship		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						between the StudyDesign and Activity classes which identifies the set of activities associated with the study design.		
	trialIntentType s	Code	C49 652	0*	Trial Intent Type	The planned purpose of the therapy, device, or agent under study in the clinical trial.	SDT M Termi nolog y Codeli st C6673	
	blindingSche ma	Code	C49 658	01	Trial Blinding Schema	The type of experimental design used to describe the level of awareness of the study subjects and/ or study personnel as it relates to the respective intervention(s) or assessments being observed, received or administered.	SDT M Termi nolog y Codeli st C6673	
	therapeuticAre as	Code	C10 130 2	0*	Therape utic Areas	A categorization of a disease, disorder, or other condition based on common characteristics and often associated with a medical specialty focusing on research and development of specific therapeutic interventions for the purpose of treatment and prevention.	(Point out to extern al dictio naries)	
	characteristics	Code	C20 754 7	0*	Study Design Charact eristic	The distinguishing qualities or prominent aspect of a study design.	C2074 16	
	trialTypes	Code	C49 660	0*	Trial Type	The nature of the interventional study for which	SDT M Termi nolog	

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						information is being collected.	y Codeli st C6673	
	interventionM odel	Code	C98 746	1	Interven tion Model Type	The general design of the strategy for assigning interventions to subjects in a clinical study. (clinicaltrials.gov)	SDT M Termi nolog y Codeli st C9907	
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyDesign and CommentAnnotation classes which provides the set of notes related to the study design.		
	encounters	Encounter		0*		A USDM relationship between the StudyDesign and Encounter classes which identifies the set of encounters associated with the study design.		
	estimands	Estimand		0*		A USDM relationship between the StudyDesign and Estimand classes which identifies the set of estimands associated with the study design.		
	indications	Indication		0*		A USDM relationship between the StudyDesign and Indication classes which identifies the set of indications associated with the study design.		

Class Name	Attribute	Data Type	NC		Preferr	Definition	Codel	Inherited
	Name		I C-	inalit	ed		ist	From
			Co de	y	Term		Ref	
	objectives	Objective		0*		A USDM		
						relationship		
						between the		
						StudyDesign and		
						Objective classes		
						which identifies the		
						set of objectives		
						associated with the		
	scheduleTimel	ScheduleTimeli		0*		study design.		
	ines	ne schedule i imen		0*		A USDM relationship		
	illes	ne				between the		
						StudyDesign and		
						ScheduleTimeline		
						classes which		
						identifies the set of		
						scheduled timelines		
						associated with the		
						study design.		
	arms	StudyArm		1*		A USDM		
						relationship		
						between the		
						StudyDesign and		
						StudyArm classes		
						which identifies the		
						set of study arms		
						associated with the		
						study design.		
	studyCells	StudyCell		1*		A USDM		
						relationship		
						between the		
						StudyDesign and		
						StudyCell classes which identifies the		
						set of study cells		
						associated with the		
						study design.		
	documentVers	StudyDefinition		01		A USDM		
	ion	DocumentVersi				relationship		
		on				between the		
						StudyDesign and		
						StudyDefinitionDo		
						cumentVersion		
						classes which		
						identifies the		
						version of the study		
						definition		
						document		
						associated with the		
	.1	G. 1 El .		0 4		study design.		
	elements	StudyElement		0*		A USDM		
			1	Ì		relationship		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						between the StudyDesign and StudyElement classes which identifies the set of study elements associated with the		
	studyIntervent ions	StudyInterventi		0*		study design. A USDM relationship between the StudyDesign and StudyIntervention classes which identifies the set of study interventions associated with study design.		
	epochs	StudyEpoch		1*		A USDM relationship between the StudyDesign and StudyEpoch classes which identifies the set of study epochs associated with the study design.		
	population	StudyDesignPo pulation		01		A USDM relationship between the StudyDesign and StudyDesignPopula tion classes which identifies the population associated with the study design.		
StudyDesignPo pulation			C14 272 8		Study Design Populati on	The population within the general population to which the study results can be generalized.		
	id	string						Populatio nDefiniti on
	name	string	C20 755 3		Study Design Populati on Name	The literal identifier (i.e., distinctive designation) of the study design population.		Populatio nDefiniti on

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	description	string	C70 834		Study Design Populati on Descript ion	A narrative representation of the study design population.		Populatio nDefiniti on
	label	string	C20 755 0		Study Design Populati on Label	The short descriptive designation for the study design population.		Populatio nDefiniti on
	includesHealth ySubjects	Boolean	C20 754 9		Study Design Populati on Includes Healthy Subjects Indicato r	An indication as to whether the study design population includes healthy subjects, that is, subjects without the disease or condition under study.		Populatio nDefiniti on
	plannedSex	Code	C20 755 1	02	Study Design Populati on Planned Sex	The protocoldefined sex within the study design population.	SDT M Termi nolog y Codeli st C6673	Populatio nDefiniti on
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyDesignPopula tion and CommentAnnotatio n classes which provides the set of notes related to the study design population.		Populatio nDefiniti on
	criteria	EligibilityCriter ion		0*		A USDM relationship between the StudyDesignPopula tion and EligibilityCriterion classes which identifies the set of eligibility criteria associated with the study design population.		Populatio nDefiniti on

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
	plannedAge	Range	C20 745 0	01	Study Design Populati on Planned Age	The anticipated age of subjects within the study design population.		Populatio nDefiniti on
	plannedEnroll mentNumber	Range	C20 745 2	01	Study Design Populati on Planned Enrollm ent Number	The value representing the planned number of subjects to be entered in a clinical trial, within the study design population.		Populatio nDefiniti on
	plannedCompl etionNumber	Range	C20 745 1	01	Study Design Populati on Planned Complet ion Number	The value representing the planned number of subjects that must complete the study		Populatio nDefiniti on
	cohorts	StudyCohort		0*		A USDM relationship between the StudyDesignPopula tion and StudyCohort classes which identifies the set of study cohorts associated with the study design population.		
StudyElement	id	string	C14 273 5		Study Design Element	A basic building block for time within a clinical study comprising the following characteristics: a description of what happens to the subject during the element; a definition of the start of the element; a rule for ending the element.		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	name	string	C18 883 3		Study Design Element Name	The literal identifier (i.e., distinctive designation) of the study design element.		
	description	string	C18 883 4		Study Design Element Descript ion			
	label	string	C20 755 4		Study Design Element Label	The short descriptive designation for the study design element.		
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyElement and CommentAnnotatio n classes which provides the set of notes related to the study element.		
	transitionEnd Rule	TransitionRule		01		A USDM relationship between the StudyElement and TransitionRule classes which provides the details associated with a transition rule used to trigger the end of a study element.		
	studyIntervent ions	StudyInterventi on		0*		A USDM relationship between the StudyElement and StudyIntervention classes which identifies the set of study interventions associated with the study element.		
	transitionStart Rule	TransitionRule		01		A USDM relationship between the StudyElement and TransitionRule		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						classes which provides the details associated with a transition rule used to trigger the start of a study element.		
StudyEpoch			C71 738		Study Epoch	A named time period defined in the protocol, wherein a study activity is specified and unchanging throughout the interval, to support a study-specific purpose.		
	id	string						
	description	string	C93 825		Study Epoch Name	The literal identifier (i.e., distinctive designation) of the study epoch, i.e., the named time period defined in the protocol, wherein a study activity is specified and unchanging throughout the interval, to support a study-specific purpose. A narrative		
	1		824		Epoch Descript ion	representation of the study epoch.		
	label	string	C20 755 5		Study Epoch Label	The short descriptive designation for the study epoch.		
	type	Code	C18 883 0	1	Study Epoch Type	A characterization or classification of the study epoch, i.e., the named time period defined in the protocol, wherein a study activity is specified and	SDT M Termi nolog y Codeli st C9907	

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						unchanging throughout the interval, to support a study-specific purpose.		
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyEpoch and CommentAnnotatio n classes which provides the set of notes related to the study epoch.		
	previous	StudyEpoch		01		A USDM relationship within the StudyEpoch class which identifies the study epoch that chronologically precedes the current study epoch.		
	next	StudyEpoch		01		A USDM relationship within the StudyEpoch class which identifies the study epoch that chronologically follows the current study epoch.		
StudyIdentifier			C83 082		Study Identifie r	A sequence of characters used to identify, name, or characterize the study.		
	id	string				·		Identifier
text	1	string	CN EW		Study Identifie r Text	An instance of structured text that represents the study identifier.		Identifier
	scope	Organization		1		A USDM relationship between the StudyIdentifier and Organization classes which provides the details associated with		Identifier

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						each organization that has assigned the study identifier.		
StudyInterventi on			C20 764 9		Study Interven tion	Any agent, device, or procedure being tested or used as a reference or comparator in the conduct of a clinical trial.		
	id	string						
	description	string	C20 764 7		Study Interven tion Descript ion	A narrative representation of the study intervention.		
	name	string	C20 755 8		Study Interven tion Name	The literal identifier (i.e., distinctive designation) of the study intervention.		
	label	string	C20 755 6		Study Interven tion Label	The short descriptive designation for the study intervention.		
	administration s	Administration		0*		A USDM relationship between the StudyIntervention and AgentAdministratio n classes which identifies the set of agent administrations associated with the study intervention.		
	type	Code	C98 747	1	Study Interven tion Type	The kind of product or procedure studied in a trial.	SDT M Termi nolog y Codeli st C9907	
	role	Code	C20 756 0	1	Study Interven tion Role	The intended use of the trial intervention within the context of the study design.	C2074 17	

Class Name	Attribute Name	Data Type	NC I C- Co de	inalit y	Preferr ed Term	Definition	ist Ref	Inherited From
	productDesign ation	Code	C20 755 9	1	Study Interven tion Product Designa tion	An indication as to whether the investigational intervention is an investigational medicinal product or an auxiliary medicinal product.	C2074 18	
	codes	Code	C20 764 8	0*	Study Interven tion Code	A symbol or combination of symbols which is assigned to the study intervention.	(Point out to multip le Biome dical coding dictio naries such as WHO Drug, ATC, UNII, etc.)	
	notes	CommentAnnot ation		0*		A USDM relationship between the StudyIntervention and CommentAnnotation classes which provides the set of notes related to the study intervention.	cic.)	
	minimumResp onseDuration	Quantity	C20 755 7	01	Study Interven tion Minimu m Respons e Duratio n	The value representing the minimum amount of time required to meet the criteria for response to study intervention.		
StudyRole			CN EW		Study Role	A designation that identifies the function of study personnel within the context of the study.		
	id	string						

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Term	Definition	Codel ist Ref	Inherited From
	name	string	CN EW		Study Role Name	The literal identifier (i.e., distinctive designation) of the study role.		
	label	string	CN EW		Study Role Label	The short descriptive designation for the study role.		
	description	string	CN EW		Study Role Descript ion	A narrative representation of the study role.		
	assignedPerso ns	AssignedPerson		0*		A USDM relationship between the StudyRole and AssignedPerson classes that identifies the set of individuals that are assigned to fill a particular role within the study.		
	code	Code	CN EW	1	Study Role Code	A symbol or combination of symbols which is assigned to the study role.	CNE W Study Role Code	
	masking	Masking		01		A USDM relationship between the StudyRole and Masking classes which describes the masking associated with the study role.	Couc	
	organizations	Organization		0*		A USDM relationship between the StudyRole and Organization classes which identifies the set of organizations associated with the study role.		
	appliesTo	StudyDesign, StudyVersion		0*		A USDM relationship between the StudyRole and		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						either StudyVersion or StudyDesign classes that identifies the study version or study design to which the study role applies.		
StudySite			C80 403		Study Site	The location at which a study investigator conducts study activities.		
	id	string						
	name	string	C20 756 6		Study Site Name	The literal identifier (i.e., distinctive designation) of the study site.		
	description	string	C20 756 4		Study Site Descript ion	A narrative representation of the study site.		
	label	string	C20 756 5		Study Site Label	The short descriptive designation for the study site.		
StudyTitle			C49 802		Study Title	The sponsor-defined name of the clinical study.		
	id	string	G20		C ₁ 1	A :		
	text	string	C20 756 7		Study Title Text	An instance of unstructured text that represents the study title.		
	type	Code	C20 756 8	1	Study Title Type	A characterization or classification of the study title.	C2074 19	
StudyVersion			C18 881 6		Study Version	A plan at a particular point in time for a study.		
	id	string	G20		G ₄ 1	A		
	versionIdentifi er	string	C20 757 0		Study Version Identifie r	A sequence of characters used to identify, name, or characterize the study version.		
	rationale	string	C94 122		Study Rational e	A statement describing the overall rationale of the study. This field		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						describes the contribution of this study to product development, i.e., what knowledge is being contributed from the conduct of this study.		
	abbreviations	Abbreviation		0*		A USDM relationship between the StudyVersion and Abbreviation classes which provides the set of abbreviations associated with the study version.		
	studyPhase	AliasCode	C48 281	01	Trial Phase	A step in the clinical research and development of a therapy from initial clinical trials to post-approval studies. NOTE: Clinical trials are generally categorized into four (sometimes five) phases. A therapeutic intervention may be evaluated in two or more phases simultaneously in different trials, and some trials may overlap two different phases. 21 CFR section 312.21; After ICH Topic E8 NOTE FOR GUIDANCE ON GENERAL CONSIDERATIO NS FOR CLINICAL TRIALS, CPMP/ICH/291/95	SDT M Termi nolog y Codeli st C6673 7	

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	Inherited
	Name	Z mm z y pc	I C-		ed	2 444444	ist	From
			Co	y	Term		Ref	
			de					
	businessThera	Code	C20	0*	Busines	A therapeutic area	(Point	
	peuticAreas		132		S	classification based	out to	
			2		Therape	on the structure and	extern	
					utic	operations of the	al	
					Areas	business unit.	dictio	
							naries)	
	studyType	Code	C14	01	Study	The nature of the	SDT	
			217		Type	investigation for	M	
			5		Classific	which study	Termi	
					ation	information is	nolog	
						being collected.	y C - 1-1:	
						(After clinicaltrials.	Codeli	
						gov)	st C9907	
	notes	Comment A mast		0*		A USDM	7	
	notes	CommentAnnot		0*				
		ation				relationship between the		
						StudyVersion and		
						CommentAnnotatio		
						n classes which		
						provides the set of		
						notes related to the		
						study version.		
	dateValues	GovernanceDat		0*		A USDM		
	date varues	e		0		relationship		
						between the		
						StudyVersion and		
						GovernanceDate		
						classes which		
						provides the set of		
						governance dates		
						associated with the		
						study version.		
	referenceIdent	ReferenceIdenti		0*		A USDM		
	ifiers	fier				relationship		
						between the		
						StudyVersion and		
						ReferenceIdentifier		
						classes which		
						identifies the set of		
						reference		
						identifiers		
						associated with the		
		C4 4 A 1		0 *		study version.		
	amendments	StudyAmendme		0*		A USDM		
		nt				relationship		
						between the		
						Study Amendment		
						StudyAmendment classes which		
						identifies the set of		
				<u> </u>]	ruenumes me set of		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						study amendments associated with the study version.		
	documentVers ions	StudyDefinition DocumentVersi on		0*		A USDM relationship between the StudyVersion and StudyDefinitionDo cumentVersion classes which identifies the version of the study definition document associated with the study version.		
	studyDesigns	StudyDesign		0*		A USDM relationship between the StudyVersion and StudyDesign classes which identifies the set of study designs associated with the study version.		
	studyIdentifier s	StudyIdentifier		1*		A USDM relationship between the StudyVersion and StudyIdentifier classes which identifies the set of study identifiers associated with the study version.		
	titles	StudyTitle		1*		A USDM relationship between the StudyVersion and StudyTitle classes which identifies the set of study titles associated with the study version.		
SubjectEnrollm ent			C37 948		Subject Enrollm ent	The act of enrolling subjects into a study. The subject will have met the inclusion/exclusion criteria to participate in the		

Class Name	Attribute Name	Data Type	NC I C- Co de	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
						trial and will have signed an informed consent form. (CDISC Glossary)		
	id	string						Geograph icScope
	code	AliasCode	C20 757 1	01	Subject Enrollm ent Code	A symbol or combination of symbols which is assigned to the subject enrollment.		Geograph icScope
	type	Code	C20 757 4	1	Subject Enrollm ent Type	A characterization or classification of the subject enrollment.		Geograph icScope
	quantity	Quantity	C20 757 3	1	Subject Enrollm ent Quantity Value	The value representing the number of individuals enrolled in a study.		
	appliesTo	StudySite		01		A USDM relationship between the SubjectEnrollment and StudySite classes which identifies the study site that applies to the subject enrollments.		
Substance			C45 306		Substan ce	Any matter of defined composition that has discrete existence, whose origin may be biological, mineral or chemical.		
	id	string						
	name	string	CN EW		Substan ce Name	The literal identifier (i.e., distinctive designation) of the substance.		
	description	string	CN EW		Substan ce Descript ion	A narrative representation of the substance.		
	label	string	CN EW		Substan ce Label	The short descriptive		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						designation for the substance.		
	codes	Code	CN EW	0*	Substan ce Code	A symbol or combination of symbols which is assigned to the substance.	(Point out to multip le Biome dical coding dictio naries such as WHO Drug, ATC, UNII,	
	strengths	Strength		1*		A USDM relationship between the Substance and Strength class which provides the values of the strengths of the substance.	etc.)	
	referenceSubst ance	Substance		01		A USDM relationship within the Substance class that identifies the association between two substances, one of which is used as a reference for the other.		
SyntaxTemplate			C20 759 6		Syntax Templat e	A standardized pattern used for the arrangement of words and phrases to create well-formed, structured sentences.		
	id name	string string	C20 757 7		Syntax Templat e Name	The literal identifier (i.e., distinctive designation) of the syntax template.		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
	description	string	C20 757 5		Syntax Templat e Descript ion	A narrative representation of the syntax template.		
	label	string	C20 757 6		Syntax Templat e Label	The short descriptive designation for the syntax template.		
	text	string	C20 757 8		Syntax Templat e Text	A structured text string containing prescribed text interspersed with user-defined parameter values.		
	notes	CommentAnnot ation		0*		A USDM relationship between the SyntaxTemplate and CommentAnnotation classes which provides the set of notes related to the syntax template.		
	dictionary	SyntaxTemplate Dictionary		01		A USDM relationship between the SyntaxTemplate and SyntaxTemplateDic tionary classes which provides the dictionary entry associated with a syntax template.		
SyntaxTemplate Dictionary			C20 759 7		Syntax Templat e Dictiona ry	A reference source that provides a listing of valid		
	id name	string string	C20 758 1		Syntax Templat e Dictiona ry Name	The literal identifier (i.e., distinctive designation) of the		

Class Name	Attribute Name	Data Type	NC I C- Co	Card inalit y	Preferr ed Term	Definition	Codel ist Ref	Inherited From
	description	string	de C20 757 9		Syntax Templat e Dictiona ry Descript	the syntax template		
	label	string	C20 758 0		Syntax Templat e Dictiona ry Label	The short descriptive designation for the syntax template dictionary.		
	parameterMap s	ParameterMap		1*	ty Euro	A USDM relationship between the SyntaxTemplateDic tionary and ParameterMap classes which identifies the set of parameter maps (parameter map entries) associated with a syntax template dictionary.		
Timing			C80 484		Timing	The chronological relationship between temporal events.		
	id	string				e vents.		
	name	string	C20 758 4		Timing Name	The literal identifier (i.e., distinctive designation) of the timing.		
	description	string	C16 464 8		Timing Descript ion	A narrative		
	label	string	C20 758 3		Timing Label	The short descriptive designation for the timing.		
	value	string	C20 134 1		Timing Value	The temporal value of the chronological relationship between temporal events.		

Class Name	Attribute	Data Type	NC	Card	Preferr	Definition	Codel	
	Name		I C- Co de	inalit y	ed Term		ist Ref	From
	valueLabel	string	C20 758 5		Timing Value Label	The short descriptive designation for the		
	windowLabel	string	C20 758 6		Timing Window Label	timing value. The short descriptive designation for a time period, or other type of interval, during which a temporal event may be achieved, obtained, or observed.		
	windowLower	string	C20 134 2		Timing Window , Lower	The earliest chronological value of an allowable period of time during which a temporal event takes place.		
	windowUpper	string	C20 134 3		Timing Window , Upper	The latest chronological value of an allowable period of time during which a temporal event takes place.		
	relativeToFro m	Code	C20 129 7	1	Timing Relative To From	The name of the reference event used to define the temporal relationship with another event.	C2012 65	
	type	Code	C20 129 8	1	Timing Type	A characterization or classification of the chronological relationship between temporal events.	C2012 64	
	relativeToSch eduledInstance	ScheduledInsta nce		01		A USDM relationship between the Timing and ScheduledInstance classes which identifies the scheduled instance (e.g., scheduled activity instances or scheduled decision		

Class Name	Attribute Name	Data Type	NC I C- Co de		Preferr ed Term	Definition	Codel ist Ref	Inherited From
						instances) to which		
						the timing is		
	relativeFromS	ScheduledInsta		1		relative to. A USDM		
	cheduledInsta	nce		1		relationship		
	nce	nec				between the Timing		
	nec					and		
						ScheduledInstance		
						classes which		
						identifies the		
						scheduled instance		
						(e.g., scheduled		
						activity instances or		
						scheduled decision		
						instances) to which		
			~~~			the timing applies.		
TransitionRule			C82		Transiti	A guide that		
			567		on Rule	governs the		
						allocation of subjects to		
						operational options		
						at a discrete		
						decision point or		
						branch (e.g.,		
						assignment to a		
						particular arm,		
						discontinuation)		
						within a clinical		
						trial plan.		
	id	string	<u> </u>					
	name	string	C20		Transiti	The literal		
			758		on Rule	identifier (i.e.,		
			8		Name	distinctive		
						designation) of the transition rule.		
	description	string	C18		Transiti	A narrative		
	description	Sumg	883		on Rule	representation of		
			5		Descript			
					ion			
	label	string	C20		Transiti	The short		
			758		on Rule	descriptive		
			7		Label	designation for the		
						transition rule.		
	text	string	C20		Transiti	An instance of		
			758		on Rule	unstructured text		
			9		Text	that represents the		
						transition rule.		

### 13 USDM API

### 13.1 General

The reference architecture API is designed as a mechanism for bulk transfer to allow for the creation of a study within the SDR, the reading of such a study, and the update of a study. No other API features are defined, nor is a granular API defined at this time. The API has been defined using the <a href="OpenApi Specification">OpenApi Specification</a>. The various routes, rules, and constraints for the use of the API are contained within the API specification itself. If further routes, rules, and constraints are required, these will be added to the machine-readable specification.

### 13.2 Serialization

When expressing USDM data in a monolithic, hierarchical document format (e.g., JSON, XML), the same element will appear multiple times because the model uses only class references for USDM entities. This is not optimal for an API and, so as not to repeat the same information within the JSON structure, the API has been designed to include an instance once and only once and allow for zero, 1, or more references to it as dictated by the USDM and the relationships therein. This mechanism relies on the unique identifiers of each class.

To ensure no duplication of content in the API JSON format, the following series of steps are taken to translate the logical USDM into the JSON format:

- 1. Where content is shared (referenced from 2 or more places), the "natural parent" relationship is identified. An example is the Endpoint class that is referenced from both the Objective and Estimand classes. Objective is considered the natural parent.
- 2. If a natural parent can be identified in the API, then the content of the child is included in the corresponding item of the natural parent (attribute names remain unchanged) and other relationships are added as cross-references, with the attribute names modified with a suffix of "Id" (singular) or "Ids" (plural) relationships. The datatype is modified to string so as to accommodate the cross-references and the corresponding identifiers.
- 3. If the natural parent cannot be identified, then a "collection" from a logical higher level class is formed and all relationships to this class in the logical model are added as cross-references in the API with the corresponding naming modifications as specified in step 2. This results in an additional relationship in the API for the higher level class to the collection. An example is for the class BiomedicalConcepts, where a collection is placed within the StudyDesign class.

#### 13.3 API Additional Attributes

A number of additional attributes have been added to the API to aid processing. These attributes are API-only artifacts and, as such, are not present within the UML specification or defined within the CT. The additional attributes are:

- 1. An **instanceType** attribute, included within all classes and used to state the class name.
- 2. Three attributes, included within the root node of the API:
  - **a. usdmVersion**: The version of the USDM to which the data transported have been generated from and conform to. This is a required attribute.
  - **b. systemName**: The name of the system that generated the data. This is an optional attribute.
  - **c. systemVersion**: The version of the system that generated the data. This is an optional attribute.

## **13.4 Required Content**

When sending data using the API it is recommended that the data include the following:

- 1. There is only 1 StudyVersion.
- There is 1 StudyIdentifier within the StudyVersion, scoped by an Organization of type Clinical Study Sponsor (C70793).
- 3. There is at least 1 StudyDesign within the StudyVersion.

# 14 Mapping to Other Standards and Formats

- <u>Creation of SDTM Trial Design Domains</u>
- Informing ClinicalTrials.gov Registry
- <u>Use of USDM for Populating Protocol Content</u>

## **14.1 Creation of SDTM Trial Design Domains**

Alignment between the USDM and SDTM Trial Design domains and controlled terminology elements related to study design enables the (automated) creation of the SDTM Trial Design Domains. The <u>SDTM Implementation Guide</u> (SDTMIG) includes a section related to Trial Design datasets. The corresponding trial design concepts include:

- Trial design
- Epoch
- Arm
- Study cell
- Element
- Branch
- Treatments
- Visit
- Criteria

These concepts are used for the following Trial Design Domains:

- Trial Arms (TA)
- Trial Elements (TE)
- Trial Visits (TV)
- Trial Inclusion/Exclusion Criteria (TI)
- Trial Summary (TS)

Other trials design domains like Trial Disease Assessments (TD) and Trial Disease Milestones (TM) that are described in the SDTMIG contain information that is stored in the USDM as well; these, however, are not explicitly discussed in this section.

The USDM structure that informs the TA, TE, and TV domains is described in Section 4.10, <u>Arms and Epochs</u>. The following table provides an overview of the mapping of USDM to the **SDTM TA domain**.

Variab Varia Ty Role C **USDM Path and Attribute Required USDM Selection / Derivation** le ble relationships or Name Label e **STUD** Study Ch Ident Re Study/@versions study.studyVersion.studyI YID Identifi ifier /StudyVersion/@studyIdentif dentifier.organization. type.code=C188724 er /StudyIdentifier/@studyIdent (Clinical StudySponsor) ifier DOMA Domai Ch Ident Re Set to "TA" ΙN ifier ar q Abbre viation Ch Topi ARMC Study/@versions Planne Re D d Arm /StudyVersion/@studyDesign ar q Code /StudyDesign/@arms /StudyArm/@name ARM Ch Study/@versions Descri Syno Re /StudyVersion/@studyDesign nym ption ar q of

Variab	Varia	Ту	Role	C	USDM Path and Attribute	Required USDM	Selection / Derivation
le	ble	pe	Kole	or	OSDM Path and Attribute	relationships	Selection / Derivation
Name	Label	pc		e		Telationships	
- 100	Planne		Qual	-	/StudyDesign/@arms		
	d Arm		ifier		/StudyArm/@description		
TAET	Planne	N	Timi	Re		/StudyCell/@arm	Link epochs via StudyCell
ORD	d	u	ng	q	/StudyVersion/@studyDesign	/StudyCell/@elem	to the corresponding study
	Order	m			s	ents	elements. Order epochs
	of				/StudyDesign/@studyCells		and their related elements
	Eleme				/StudyCell/@epoch		based on previous
	nt within				/StudyEpoch/@previous   @next		StudyEpoch and next
	Arm				whext		StudyEpoch attributes and derive a corresponding
	Ailli						ordering number.
ETCD	Eleme	Ch	Reco	Re	Study/@versions	/StudyCell/@arm	ordering number.
	nt	ar	rd	q	/StudyVersion/@studyDesign	,	
	Code		Qual		s		
			ifier		/StudyDesign/@studyCells		
					/StudyCell/@elements		
ELEM	D	CI.	C	D.	/StudyElement/@name	/0. 1 0.11/8	
ELEM ENT	Descri ption		Syno nym	Pe rm	Study/@versions /StudyVersion/@studyDesign	/StudyCell/@arm	
LINI	of	ar	Qual	1111	study Version/ @studyDesign		
	Eleme		ifier		/StudyDesign/@studyCells		
	nt				/StudyCell/@elements		
					/StudyElement/@description		
TABR	Branch	Ch	Rule	Ex	Study/@versions	/StudyCell/@epoc	ScheduledInstances in a
ANCH		ar		p	/StudyVersion/@studyDesign		timeline point to a
					S	/StudyCell/@arm	StudyEpoch (see Section
					/StudyDesign/@scheduleTim elines		4.14, <u>Study Timing</u> ). Branching information can
					/ScheduleTimeline/@instanc		be stored as
					es		scheduledDecisionInstanc
					/ScheduledDecisionInstance/		es using the
					@conditionAssignments		ConditionAssignment that
							points to the first instance
T + T T		G1	<b>D</b> 1	_		(0.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1	related to the next epoch.
TATR			Rule		Study/@versions	/ScheduledActivity	
ANS	ion Rule	ar		p	/StudyVersion/@studyDesign	/StudyCell/@epoch	timeline point to a StudyEpoch (see Section
	Kuic				/StudyDesign/@scheduleTim	/StudyCell/@arm	4.14, Study Timing).
					elines	/Study Con/ Curin	Transition rule
					/ScheduleTimeline/@instanc		information is stored as
					es		scheduledDecisionInstanc
					/ScheduledDecisionInstance/		es using the
					@conditionAssignments		ConditionAssignment that
							points to an instance not being the default next
							instance on the timeline.
EPOC	Epoch	Ch	Timi	Re	Study/@versions	/StudyCell/@arm	
Н	1	ar	ng	q	/StudyVersion/@studyDesign	_	
					s		
					/StudyDesign/@studyCells		
					/StudyCell/@epoch		
	<u>I</u>	<u> </u>			/StudyEpoch/@name		

The following table provides an overview of the mapping of USDM to the **SDTM TE domain** 

					erview of the mapping of		
Varia ble Name	Variab le Label	Ty pe	Role	Co re	USDM Path and Attribute	Required USDM relationships	Selection / Derivation
STUD YID	Study Identifi er	Ch ar	Identi fier	Re q	Study/@versions /StudyVersion/@study Identifiers /StudyIdentifier/@stud yIdentifier		study.studyVersion.studyIden tifier.organization. type.code=C188724 (Clinical StudySponsor)
DOM AIN	Domain Abbrev iation	Ch ar	Identi fier	Re q			Set to "TE"
ETCD	Elemen t Code	Ch ar	Topic	Re q	Study/@versions /StudyVersion/@study Designs /StudyDesign/@eleme nts /StudyElement/@name		
ELEM ENT	Descrip tion of Elemen t	Ch ar	Syno nym Quali fier	Re q	Study/@versions /StudyVersion/@study Designs /StudyDesign/@eleme nts /StudyElement/@descr iption		
TEST RL	Rule for Start of Elemen t	Ch ar	Rule	Re q	Study/@versions /StudyVersion/@study Designs /StudyDesign/@eleme nts /StudyElement/@transi tionStartRule /TransitionRule/@text		
TEEN RL	Rule for End of Elemen t	Ch ar	Rule	Pe rm	Study/@versions /StudyVersion/@study Designs /StudyDesign/@eleme nts /StudyElement/@transi tionEndRule /TransitionRule/@text		
TEDU R	Planned Duratio n of Elemen t	Ch	Timi ng	Perm	Study/@versions /StudyVersion/@study Designs /StudyDesign/@sched uleTimelines /ScheduleTimeline/@i nstances /ScheduledActivityInst ance/@timings /Timing/@value	/ScheduledActivityIn stance/@epoch /StudyCell/@epoch /StudyCell/@elemen ts	Select scheduleInstances that relate to start of the associated StudyEpoch associated with the corresponding study Element via StudyCell. Select the scheduleInstance associated with the start of the next studyEpoch. Use Timing.values of all related timings that specify the period in between for

Varia ble Name	Variab le Label	Ty pe	Role	USDM Path and Attribute	Required USDM relationships	Selection / Derivation
						calculation of the total element duration.

The following table provides an overview of the mapping of USDM to the SDTM TV domain

						of USDM to the SDTM T	
Variab	Variab	Ty	Role	Co	USDM Path and	Required USDM	Selection / Derivation
le	le	pe		re	Attribute	relationships	
Name	Label						
STUD	Study	Ch	Identi	Re			study.studyVersion.studyIden
YID	Identifi	ar	fier	q	/StudyVersion/@stu		tifier.organization.
	er				dyIdentifiers		type.code=C188724 (Clinical
					/StudyIdentifier/@st		StudySponsor)
					udyIdentifier		
DOM	Domain	Ch	Identi	Re			Set to "TV"
AIN	Abbrev	ar	fier	q			
	iation						
VISIT	Visit	Nu	Topic	Re	Study/@versions		Order encounters based
NUM	Numbe	m		q	/StudyVersion/@stu		previous and next attributes
	r				dyDesigns		and derive the visit order
					/StudyDesign/@enc		number correspondingly.
					ounter		Assign numbers based on
					/Encounter/@previo		applicable standard visit
					us   @next		numbering rules.
VISIT	Visit	Ch	Syno	Re			
	Name	ar	nym	q	/StudyVersion/@stu		
			Quali		dyDesigns		
			fier		/StudyDesign/@enc		
					ounter		
				_	/Encounter/@name		
VISIT	Planned		Timi	Pe	Study/@versions		
DY	Study	m	ng	rm	/StudyVersion/@stu		
	Day of				dyDesigns		
	Visit				/StudyDesign/@enc		
					ounter		
					/Encounter/@timing		
					/Timing/@timingVa lue		
ARMC	Planned	Ch	Reco	Ex	Study/@versions	/StudyCell/@epoch	In case visits differ by arm,
D	Arm	ar	rd		/Study/@version/@stu	/ScheduledActivityInst	the corresponding arm can be
ט	Code	aı	Quali	p	dyDesigns	ance/@epoch	derived via the
	Couc		fier		/StudyDesign/@stu	/ScheduledActivityInst	ScheduledActivityInstance
	1		1101		dyCells	ance/@encounter	relating the encounter via
	1				/StudyCell/@arm	ance/ @ encounter	StudyEpoch and StudyCell to
	1				/StudyCen/@arm /StudyArm/@name		the corresponding StudyArm.
ARM	Descrip	Ch	Syno	Pe	Study/@versions	/StudyCell/@epoch	the corresponding budy/11111.
7 11 (17)	tion of	ar	nym	rm	•	/ScheduledActivityInst	
	Planned	u1	Quali	1111	dyDesigns	ance/@epoch	
	Arm		fier		/StudyDesign/@stu	/ScheduledActivityInst	
			1101		dyCells	ance/@encounter	
	1				/StudyCell/@arm		
	1				/StudyArm/@descri		
	1				ption		
	1	·	1	1	1	I	l

Variab	Variab	Ty	Role	Co	USDM Path and	Required USDM	Selection / Derivation
le	le	pe		re	Attribute	relationships	
Name	Label						
TVST	Visit	Ch	Rule	Re	Study/@versions		
RL	Start	ar		q	/StudyVersion/@stu		
	Rule				dyDesigns		
					/StudyDesign/@enc		
					ounter		
					/Encounter/@transit		
					ionStartRule		
					/TransitionRule/@te		
					xt		
TVEN	Visit	Ch	Rule	Pe	Study/@versions		
RL	End	ar		rm	/StudyVersion/@stu		
	Rule				dyDesigns		
					/StudyDesign/@enc		
					ounter		
					/Encounter/@transit		
					ionEndRule		
					/TransitionRule/@te		
					xt		

The following table provides an overview of the mapping of USDM to the SDTM TI domain.

Varia	Variable	Ty	Role		USDM Path and Attribute		Selection / Derivation
ble	Label	pe		re		d	
Name						USDM	
						relation	
						ships	
STUD	Study	Ch	Identi	Re	Study/@versions		study.studyVersion.studyIdent
YID	Identifier	ar	fier	q	/StudyVersion/@studyIdentifiers		ifier.organization.
					/StudyIdentifier/@studyIdentifie		type.code=C188724 (Clinical
					r		StudySponsor)
DOM	Domain	Ch	Identi	Re			Set to "TI"
AIN	Abbreviatio	ar	fier	q			
	n						
IETES	Incl/Excl	Ch	Topic	Re	Study/@versions		Eligibility criteria might be
TCD	Criterion	ar		q	/StudyVersion/@studyDesigns		directly linked to a study
	Short Name				/StudyDesign/@population		Population or via one of the
					(/StudyDesignPopulation/@coho		corresponding cohorts.
					rts)		Therefore an alternative path
					/StudyDesignPopulation StudyC		is specified via the
					ohort/@criteria		StudyCohort class.
		~*	~	_	/EligibilityCriteria/@identifier		
IETES	Inclusion/E	Ch	Syno	Re	Study/@versions		The eligibility criteria are
T	xclusion	ar	nym	q	/StudyVersion/@studyDesigns		based on the SyntaxTemplate
	Criterion		Quali		/StudyDesign/@population		class (see <u>Section 4.21</u> ).
			fier		(/StudyDesignPopulation/@coho		Referenced values need to be
					rts)		replaced by actual values
					/StudyDesignPopulation StudyC		before creation of IETEST.
					ohort/@criteria		
TECATE	T 1 /C	CL	C	D .	/EligibilityCriteria/@text		
IECAT	Inclusion/E	Ch	Grou	Re	Study/@versions		
	xclusion	ar	ping	q	/StudyVersion/@studyDesigns		
	Category		Quali		/StudyDesign/@population		
			fier		(/StudyDesignPopulation/@coho		

Varia ble Name	Variable Label	Ty pe	Role	Co re	USDM Path and Attribute	Require d USDM relation ships	Selection / Derivation
					rts) /StudyDesignPopulation StudyC ohort/@criteria /EligibilityCriteria/@category /code/@decode		
IESCA T	Inclusion/E xclusion Subcategory	Ch ar	Grou ping Quali fier	Per m			Permitted value. Not available in USDM. Can be applied according to user preference.
TIRL	Inclusion/E xclusion Criterion Rule	Ch ar	Rule	Per m	/StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyC ohort/@criteria /EligibilityCriteria/@text		The eligibility criteria are based on the SyntaxTemplate class (see Section 4.21), which enhances computer readability. References values should <b>not</b> be replaced by actual values for TIRL.
TIVER S	Protocol Criteria Versions	Ch ar	Recor d Quali fier	Per m	Study/@versions /StudyVersion/@documentVersi on /StudyProtocolDocumentVersio n/@protocolVersion		

The following table provides an overview of the mapping of USDM to the SDTM TS domain.

Variable			Role	Co	USDM Path and	Require	Selection / Derivation
Name	Label	pe		re	Attribute	d USDM	
						relations	
						hips	
STUDYI	Study	Ch	Identif	Re	Study/@versions		study.studyVersion.studyIdentifie
D	Identifier	ar	ier	q	/StudyVersion/@studyIdent		r.organization.
					ifiers		type.code=C188724 (Clinical
					/StudyIdentifier/@studyIde		StudySponsor)
					ntifier		
DOMAI	Domain	Ch	Identif	Re			Set to "TS"
N	Abbrevia	ar	ier	q			
	tion						
TSSEQ	Sequenc	Nu	Identif	Re	See TSPARM mapping		
	e	m	ier	q	table below		
	Number						
TSGRPI	Group	Ch	Identif	Per	See TSPARM mapping		
D	ID	ar	ier	m	table below		
TSPAR	Trial	Ch	Topic	Re	See TSPARM mapping		
MCD	Summar	ar		q	table below		
	y						
	Paramete						
	r Short						
	Name						
TSPAR	Trial	Ch	Synon	Re	See TSPARM mapping		
M	Summar	ar	ym	q	table below		
	y						

Variable Name	Variable Label	Ty pe	Role	Co re	USDM Path and Attribute	Require d USDM relations	Selection / Derivation
						hips	
	Paramete		Qualif				
	r		ier				
TSVAL	Paramete	Ch	Result	Ex	See TSPARM mapping		
	r Value	ar	Qualif	p	table below.If not otherwise		
			ier		specified:Code/@decode		
TSVAL	Paramete	Ch	Result	Per	Fill in case of missing		
NF	r Value	ar	Qualif	m	values with expected data		
	Null		ier		as described in the		
	Flavor				<u>SDTMIG</u>		
TSVAL	Paramete	Ch	Result	Ex	See TSPARM mapping		
CD	r Value	ar	Qualif	p	table below.If not otherwise		
	Code		ier		specified:Code/@decode		
TSVCD	Name of	Ch	Result		See TSPARM mapping		
REF	Referenc	ar	Qualif	p	table below.		
	e		ier		If not otherwise		
	Terminol				specified:Code/@codeSy		
	ogy				stem		
TSVCD	Version	Ch	Result		See TSPARM mapping		
VER	of the	ar	Qualif	p	table below.		
	Referenc		ier		If not otherwise		
	e				specified:Code/@codeSy		
	Terminol				stemVersion		
	ogy						

The following table provides a list of published Trial Summary parameters (TSPARM) and their mapping to USDM elements (i.e., entities, attributes, valid values). The table includes only those parameters for which there is a mapping. Frequently used and required parameters are included.

The table is based on the SDTM Controlled Terminology codelist C66738, from SDTM Terminology Version 2023-

09-29. For all synonyms and definitions, please see the corresponding terminology file.

<b>TSPAR</b>	TSPA	Cod	Cod	TSVAL	Selection / Derivations	TSS	TSGR
M	RMCD	e	elist	USDM Path and Attribute		EQ	PID
			Cod				
			e				
Adaptive	ADAP	C14	C66	Study/@versions	If characteristics include		
Design	T	6995	738	/StudyVersion/@studyDesigns	"ADAPTIVE" then		
				/StudyDesign/@characteristics	TSVAL="Y" and		
				/code/@decode	TSVALCD="C49488"		
					Otherwise TSVAL="N"		
					and		
					TSVALCD="C49487"		
Planned	AGEM	C49	C66	Study/@versions	Use minimum of		
Minimum	IN	693	738	/StudyVersion/@studyDesigns	minimum age values of all		
Age of				/StudyDesign/@population	populations included		
Subjects				(/StudyDesignPopulation/@cohorts)	(studyDesignPopulations		
				/StudyDesignPopulation	and Cohorts). Transform		
				StudyCohort/@plannedAge	according to ISO 8601		
				/Range/@minValue + @unit	standards. If one ore more		
					populations have a null		
					minValue then TSVAL		
					should be set to null and		
					TSVALNF should be		

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
Planned Minimum	AGEM AX	C49 694	C66	Study/@versions /StudyVersion/@studyDesigns	filled instead according to ISO 21090. Use maximum of maximum age values of all		
Age of Subjects				/StudyDesign/@population (/StudyDesignPopulation/@cohorts) /StudyDesignPopulation  StudyCohort/@plannedAge /Range/@maxValue + @unit	populations included (studyDesignPopulations and Cohorts). Transform according to ISO 8601 standards. If one ore more populations have a null maxValue then TSVAL should be set to null and TSVALNF should be filled instead according to ISO 21090.		
Comparat ive Treatment Name	COMP	C68 612	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@name	StudyIntervention/@role/Code/@Code<>"C41161"  (not "Experimental Intervention") andStudyIntervention/@prod uctDesignation/ Code/@decode="IMP"	Add Uni que num ber if mor e than	If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Current Therapy or Treatment	CURT RT	C85 582	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@name	StudyIntervention/@role/Code/@Code="C165822" ("Background Treatment")	Add Uni que num ber if mor e than	If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Dose Level; Dose per Administr ation	DOSE	C25 488	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@administrations /AgentAdministration/@dose /Quantity/@value			If applica ble, combin e with the corresp

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
							onding interve ntion variabl es by a commo n
Dosing Frequenc y	DOSF RQ	C89 081	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@administrations /AgentAdministration/@frequency			If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Dose Units	DOSU	C73 558	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@administrations /AgentAdministration/@dose /Quantity/@unit			If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Extension Trial Indicator	EXTTI ND	C13 9274	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@characteristics /code/@decode	If characteristics include "Extension" then TSVAL="Y" and TSVALCD="C49488" Otherwise TSVAL="N" and TSVALCD="C49487"		
Planned Country of Investigat ional Sites	FCNT RY	C98 770	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@appliesTo /StudySite/@currentEnrollment /SubjectEnrollment/@code /AliasCode/@StandardCode	SubjectEnrollment/@type /code/@code=C25464 ("Country")	Add Uni que num ber if mor	

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
						e than 1	
Healthy Subject Indicator	HLTS UBJI	C98 737	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@cohorts) /StudyDesignPopulation  StudyCohort/@includesHealthySubjects	If True then TSVAL="Y" and TSVALCD="C49488" If False then TSVAL="N" and TSVALCD="C49487"		
Trial Disease/C ondition Indication ; Trial Disease/C ondition Indication Descripti on	INDIC	C11 2038	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@indications /Indication/@name or @description			
Interventi on Model	INTM ODEL	C98 746	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@interventionModel			
Interventi on Type	INTTY PE	C98 747	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@type			If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Trial Length	LENG TH	C49 697	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@scheduleTimelines /ScheduleTimeline/@instances /ScheduledActivityInstance/@timing s /Timing/@value	Select scheduleInstances that relate to start of the study. Select the scheduleInstance associated with the end of the study. Use Timing.values of all related timings that specify the period in between for calculation of the total trial length.		
Planned Number of Arms	NARM S	C98 771	C66 738	Study/@versions /StudyVersion/@studyDesigns	Count number of instances (each instance is an arm) defined in StudyArm class		

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
				/StudyDesign/@arms /StudyArm			
Number of Groups/C ohorts	NCOH ORT	C12 6063	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@cohorts /StudyCohort	Count number of instances (each instance is an cohort) defined in StudyCohort class		
Trial Explorato ry Objective	OBJEX P	C16 3559	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@text	Objective/@level /code/@Code = C163559 ("Exploratory Objective") Objectives are based on the SyntaxTemplate class (see Section 4.20). References values need to be replaced by actual values before creation of OBJEXP.	Add Uni que num ber	combin e with the corresp onding outcom e measur es by a commo n tsgrpid
Study Primary Objective ; Trial Primary Objective	OBJPR IM	C85 826	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@text	Objective/@level /code/@Code = C85826 ("Study Primary Objective") Objectives are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of OBJPRIM.	Add Uni que num ber	combin e with the corresp onding outcom e measur es by a commo n tsgrpid
Study Secondar y Objective ; Trial Secondar y Objective	OBJSE C	C85 827	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@text	Objective/@level /code/@Code = C85827 ("Study Secondary Objective") Objectives are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of OBJSEC.	Add Uni que num ber	combin e with the corresp onding outcom e measur es by a commo n tsgrpid
Explorato ry Outcome Measure	OUTM SEXP	C98 724	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text	Endpoint/@level /code/@Code = C170559 ("Exploratory Endpoint") Endpoints are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of OUTMSEXP.	Add Uni que num ber	combin e with the corresp onding objecti ve by a commo

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
					Alternatively, the referenced biomedical concept can be used for OUTMSEXP.		n tsgrpid
Primary Outcome Measure	OUTM SPRI	C98 772	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text	Endpoint/@level /code/@Code = C94496 ("Primary Endpoint") Endpoints are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of OUTMSPRI. Alternatively, the referenced biomedical concept can be used for OUTMSPRI.	Add Uni que num ber	combin e with the corresp onding objecti ve by a commo n tsgrpid
Secondar y Outcome Measure	OUTM SSEC	C98 781	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text	Endpoint/@level /code/@Code = C139173 ("Secondary Endpoint") Endpoints are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of OUTMSSEC. Alternatively, the referenced biomedical concept can be used for OUTMSSEC.	Add Uni que num ber	combin e with the corresp onding objecti ve by a commo n tsgrpid
Pharmaco logic Class	PCLAS	C98 768	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/ @pharmacologicClass	Corresponding @productDesignation should correspond to IMP		If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Anticipat ed Enrollme nt; Planned Enrollme nt;	PLANS UB	C49 692	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/ @plannedEnrollmentNumber /Range/@MinValue + @MaxValue	Combine MinValue and MaxValue. If equal or only 1 available then only show once.		- G- F - S

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
Planned Number of Subjects; Target Enrollme nt							
Planned Treatment Duration	PTRTD UR	C13 9276	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@administrations /AgentADministration/@duration /AdministrationDuration/@quantity /Quantity/@value + @unit			If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Trial is Randomiz ed	RAND OM	C25 196	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@characteristics /code/@decode	If characteristics include "RANDOMIZED" then TSVAL="Y" and TSVALCD="C49488" Otherwise TSVAL="N" and TSVALCD="C49487"		tog:pio
Rare Disease Indicator	RDIND	C12 6070	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@indications /Indication/@isRareDisease	If True then TSVAL="Y" and TSVALCD="C49488" If False then TSVAL="N" and TSVALCD="C49487"		
Registry Identifier	REGID	C98 714	C66 738	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier	StudyIdentifier/@studyId entifierScope /Organization/@type /Code/@code=C93453 ("Clinical Study Registry") Fill TSVCDREF with corresponding organization namestudyIdentifier/@studyId entifierScope /Organization/@name	Add Uni que num ber if mor e than	
Route of Administr ation	ROUT E	C38 114	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@administrations /AgentAdministration/@route	_		

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	TSS EQ	TSGR PID
Sex of Participan ts	SEXPO P	C49 696	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plannedS ex			
Clinical Study Sponsor; Sponsor; Study Sponsor	SPONS OR	C70 793	C66 738	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifierSco pe /Organization/@name	Organization/@type /Code/@code=C70793 ("Clinical Study Sponsor") TSVALCD=Organizatio n/@identifier TSVCDREF=Organizati on/@identifierScheme		
Sponsor's Study Reference ID	SPREF ID	C13 5009	C66 738	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier	StudyIdentifier/@studyId entifierScope /Organization/@type /Code/@code=C70793 ("Clinical Study Sponsor")		
Study Type; Study Type Classifica tion	STYPE	C14 2175	C66 738	Study/@versions /StudyVersion/@studyType			
Study Blinding Design; Study Blinding Schema; Study Masking Design; Trial Blinding Design; Trial Blinding Schema; Trial Blinding Schema; Trial Masking Design	TBLIN	C49 658	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@blindingSchema			
Control Type	TCNT RL	C49 647	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@role	StudyIntervention/@prod uctDesignation/ Code/@Decode="NIMP" Map valid values of @role to TCNTRL		
Therapeut ic Area	THER AREA	C10 1302	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@therapeuticAreas			

TSPAR M	TSPA RMCD	Cod e	Cod elist Cod e	TSVAL USDM Path and Attribute	Selection / Derivations	EQ	TSGR PID
Trial Intent Type	TINDT P	C49 652	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@trialIntentTypes		Add Uni que num ber if mor e than	
Official Study Title; Study Title; Trial Title	TITLE	C49 802	C66 738	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	StudyTitle/@Type/Code/ @decode="Official Study Title"		
Trial Phase; Trial Phase Classifica tion	TPHAS E	C48 281	C66 738	Study/@versions /StudyVersion/@studyPhase /AliasCode/@standardCode			
Investigat ional Therapy or Treatment	TRT	C41 161	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventions /StudyIntervention/@name	StudyIntervention/@role/Code/@Code="C41161"		If applica ble, combin e with the corresp onding interve ntion variabl es by a commo n tsgrpid
Trial Scope; Trial Type	ТТҮРЕ	C49 660	C66 738	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@trialTypes		Add Uni que num ber if mor e than	

## 14.2 Informing ClinicalTrials.gov Registry

The Clinical Trials.gov registry can largely be filled with the study design information captured in the USDM. The definitions for protocol registration data elements submitted to ClinicalTrials.gov for interventional studies (clinical trials) and observational studies are provided on the corresponding definitions site. Included topics and whether they are covered in USDM are presented in the table below.

CT.gov topic	USDM coverage
Study Identification	Yes
Study Status	No; not available at study design stage
Sponsor/Collaborators	No
Oversight	No
Study Description	No; protocol text covered by the Unstructured Content (see Section 4.20) class may be used for this.
Conditions and Keywords	No
Study Design	Yes; Interventional Study design parameters
Arms, Groups, and Interventions	Yes
Outcome Measures	Yes
Eligibility	Yes; Interventional Study design parameters
Contacts, Locations, and Investigator	Limited; not presented in this overview
Information	
IPD Sharing Statement	No
References	No

The mapping for the required data elements of topics that are covered is specified below.

The mapping to Study Identification is presented below. See Section 4.7, Study Identifiers and Titles, for a description of the related features in the USDM.

**Selection/Derivation** 

	- 10 -	CT.gov Requireme nt	USDM path and attribute
Study	Brief	Required	Study/@versions
Identificatio	Title		/StudyVersion/@titles

Path	Variable	Requireme nt		
Study Identificatio n	Brief Title	Required	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	StudyTitle/@Type/Code/@decode="B rief Study Title" limit to 300 characters
Study Identificatio n. Brief Title	Acronym	Required, If available	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	StudyTitle/@Type/Code/@decode="S tudy Acronym" limit to 14 characters
Study Identificatio n	Official Title	Required	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	StudyTitle/@Type/Code/@decode="O fficial Study Title" limit to 600 characters
Study Identificatio n	Secondar y ID	Required, If available	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier	StudyIdentifier/@studyIdentifierScope /Organization/@type /Code/@code <> C70793 ("Clinical Study Sponsor") studyIdentifier/@studyIdentifierScope /Organization/@name <> "NCT" (or NCT alias)
Study Identificatio n. Secondary ID	Туре	Required, If secondary ID available	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier Scope /Organization/@name	Map organization name to corresponding CT.gov terminology.
Study Identificatio n. Secondary ID	Descripti on	Required, If secondary ID available	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier Scope /Organization/@name	

CT.gov	CT.gov	CT.gov	USDM path and attribute	Selection/Derivation
Path	Variable	Requireme		
		nt		
Study	Study	Required	Study/@versions	In case of "PATIENT REGISTRY" in
Identificatio	Type		/StudyVersion/@Type	USDM, map to "Observational" in
n			/code/@decode	CT.gov. Other Study types can be
				submitted as is.

The mapping to Study Design, interventional study design parameters is presented below. See Section 4.6,

Study, Protocols, and Amendments, for a description of the related features in the USDM.

			its, for a description of the related featu		
CT.gov Path	CT.gov Variable	CT.gov Require ment	USDM path and attribute	Require d USDM relation ship	Selection/Derivation
Study Design. Intervent ional Study Design	Primary Purpose	Required	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@trialTypes /code/@decode		See Primary objective:/StudyDesign/@objectives /objective/@text where /StudyDesign/@objectives /objective/@level /code/@code=C85826 Select the TrialType that relates to the primary objective. There are 2 options to do this:  • repeat of decode terminology in objective text • reference from primary objective text to corresponding trialtype instance
Study Design. Intervent ional Study Design	Study Phase	Required	Study/@versions /StudyVersion/@studyPhase /AliasCode/@standardCode /code/@decode		Remove "A" and "B" from SDTM terminology (codelist C66737) and map 1 to 1 to CT.gov terminology if possible.
Study Design. Intervent ional Study Design	Intervent ional Study Model	Required	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@interventionModel /code/@decode		Translate CROSS-OVER to CROSSOVER. Other decode values from SDTM terminology (codelist C99076) can be submitted as is.
Study Design. Intervent ional Study Design. Intervent ional Study Model	Model descripti on		study/@versions /studyVersion/@documentVersion studyProtocolDocumentVersion/@c ontents /NarrativeContent/@text		NarrativeContent/@sectionTitl e="Intervention Model" limit to 1000 characters

CT.gov Path	CT.gov Variable	CT.gov Require ment	USDM path and attribute	Require d USDM relation ship	Selection/Derivation
Study Design. Intervent ional Study Design	Number of Arms	Required	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms /StudyArm		Count number of instances (each instance is an arm) defined in StudyArm class
Study Design. Intervent ional Study Design	Masking	Required	Study/@versions / StudyVersion/@studyDesigns /StudyDesign/@maskingRoles /Masking/@role /code/@decode		If no masking roles are defined in USDM then set Masking to "No Masking".  If masking role in USDM = "Sponsor" then leave empty.  All other values can be submitted as is
Study Design. Intervent ional Study Design. Masking	Masking Descripti on		Study/@versions / StudyVersion/@studyDesigns /StudyDesign/@maskingRoles /Masking/@role /code/@decode + @description		If masking role in USDM = "Sponsor" then fill with "Sponsor" + corresponding description.
Study Design. Intervent ional Study Design	Allocatio n	Required	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms /StudyArm and Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@characteristics /code/@decode		Count number of instances (each instance is an arm) defined in StudyArm class. If 1 or less then submission value is "N/A (not applicable)". Else If characteristics include "RANDOMIZED" then submission value is "Randomized" Otherwise submission value is "Nonrandomized"
Study Design. Intervent ional Study Design	Enrollme nt	Required	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plannedE nrollmentNumber /Range/@MinValue + @MaxValue		Combine MinValue and MaxValue. If equal or only 1 of them available then only show once.

The mapping to Arms, Groups and Interventions is presented below. See Section 4.10, Arms and Epochs, and Section 4.17, Study Interventions, for descriptions of the related features in the USDM.

CT.gov Path	CT.gov Variable	CT.gov Requirem	USDM path and attribute	Required USDM relationship	Selection/Deriv ation
Arms, Groups and Interventio ns. Arm Informatio n	Arm Title	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@arms /StudyArm/@name		Limit to 100 characters.

CT.gov Path	CT.gov Variable	CT.gov Requirem ent	USDM path and attribute	Required USDM relationship	Selection/Deriv ation
Arms, Groups and Interventio ns. Arm Informatio n	Arm Type	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@arms /StudyArm/@type /code/@decode		In case USDM arm types "Control" and "Treatment" are used they may be mapped to "Other" or any of the Experimental or Comparator types. All other USDM arm types can directly be used by moving the word "arm" from the USDM arm decode value.
Arms, Groups and Interventio ns. Arm Informatio n	Arm Description	If needed	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@arms /StudyArm/@description		Limit to 999 characters.
Arms, Groups and Interventio ns. Group/Co hort Informatio n	Group/Cohort Label	For observatio nal studies only	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@population / StudyDesignPopulation/@c ohorts /StudyCohort/@label		Limit to 100 characters.
Arms, Groups and Interventio ns. Group/Co hort Informatio n	Group/Cohort Description	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@population / StudyDesignPopulation/@c ohorts /StudyCohort/@description		Limit to 999 characters.
Arms, Groups and Interventio ns. Interventio ns	Intervention Type	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@studyCells /StudyCell/@elements /StudyElement/@studyInter ventions /StudyIntervention/@type /Code/@decode	Study/@versions /StudyVersion/@study Designs /StudyDesign/@studyC ells /StudyCell/@StudyArm	StudyCell relates StudyArm with corresponding element that relates to the corresponding intervention. From

CT.gov	CT.gov	CT.gov	USDM path and attribute	Required USDM	Selection/Deriv
Path	Variable	Requirem	•	relationship	ation
		ent			ClinicalTrials.go v: "If the same intervention is associated with more than one arm or group, provide the information once and use the Arm or Group/Interventi on Cross- Reference to associate it with more than one arm or group." Text transformation is needed for 1 to 1 mapping to ClinicalTrials.go v terminology.
Arms, Groups and Interventio ns. Interventio ns	Intervention Name	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@studyCells /StudyCell/@elements /StudyElement/@studyInter ventions /StudyIntervention/@name		Limit to 200 characters.
Arms, Groups and Interventio ns. Interventio ns	Other Intervention Name	If any	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@studyCells /StudyCell/@elements /StudyElement/@studyInter ventions /StudyIntervention/@label		Upon judgement of (system) user to decide whether label should be included as other intervention name. Limit to 200 characters.
Arms, Groups and Interventio ns. Interventio ns	Intervention Description	Required	Study/@versions /StudyVersion/@studyDesi gns /StudyDesign/@studyCells /StudyCell/@elements /StudyElement/@studyInter ventions /StudyIntervention/@descri ption		Limit to 1000 characters.
Arms, Groups	Arm or Group/Intervent	Required	Study/@versions /StudyVersion/@studyDesi	Study/@versions /StudyVersion/@study	From ClinicalTrials.go

CT.gov	CT.gov	CT.gov	USDM path and attribute	Required USDM	Selection/Deriv
Path	Variable	Requirem		relationship	ation
		ent			
and	ional Cross-		gns	Designs	v: "If the same
Interventio	References		/StudyDesign/@studyCells	/StudyDesign/@studyC	intervention is
ns.			/StudyCell/@elements	ells	associated with
Interventio			/StudyElement/@studyInter	/StudyCell/@StudyArm	more than one
ns			ventions		arm or group,
			/StudyIntervention/		provide the
					information
					once and use the
					Arm or
					Group/Interventi
					on Cross-
					Reference to
					associate it with
					more than one
					arm or group."

The mapping to **Outcome Measures** is presented below. See Section 4.17, <u>Study Objectives and Endpoints</u>, for a description of the related features in the USDM.

CT.gov Path	CT.gov Variable	CT.gov Requirem ent	USDM path and attribute	Required USDM relationship	Selection/Derivati on
Outcome Measures . Primary Outcome Measure Informati on	Title	Required	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@name		 /Endpoint/@level /code/@code=C94 496 Limit to 254 characters.
Outcome Measures . Primary Outcome Measure Informati on	Descripti on	If available	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text		/Endpoint/@level /code/@code=C94 496 The endpoint text is based on the SyntaxTemplate class (see Section 4.21). Referenced values need to be replaced by actual values before submitting. Limit to 999 characters.
Outcome Measures . Primary Outcome Measure Informati on	Time Frame	Required	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@scheduleTimeli nes /ScheduleTimeline/@ScheduleI nstance /Timing/@value	/Endpoint/@level /code/@code=C94 496 In case of reference to the corresponding Timing class, check and use the

CT.gov Path	CT.gov Variable	CT.gov Requirem ent	USDM path and attribute	Required USDM relationship	Selection/Derivati on
					referenced timing for this attribute. Limit to 254 characters.
Outcome Measures . Primary Secondar y Measure Informati on	Title	If any	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@name		 /Endpoint/@level /code/@code=C13 9173 Limit to 254 characters.
Outcome Measures . Primary Secondar y Measure Informati on	Descripti on	If available	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text		/Endpoint/@level /code/@code=C13 9173 The endpoint text is based on the SyntaxTemplate class. Referenced values need to be replaced by actual values before submitting. Limit to 999 characters.
Outcome Measures . Primary Secondar y Measure Informati on	Time Frame	If any	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@scheduleTimeli nes /ScheduleTimeline/@ScheduleI nstance /Timing/@value	/Endpoint/@level /code/@code=C13 9173 In case of reference to the corresponding Timing class, check and use the referenced timing for this attribute. Limit to 254 characters.
Outcome Measures . Other Pre- specified Outcome Measures	Title	If any	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@name		/Endpoint/@level /code/@code=C17 0559 Limit to 254 characters.
Outcome Measures . Other Pre- specified Outcome Measures	Descripti on	If available	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text		 /Endpoint/@level /code/@code=C17 0559 The endpoint text is based on the SyntaxTemplate

CT.gov Path	CT.gov Variable	CT.gov Requirem ent	USDM path and attribute	Required USDM relationship	Selection/Derivati on
Outcome Measures . Other Pre- specified Outcome Measures	Time Frame	If any	Study/@versions /StudyVersion/@studyD esigns /StudyDesign/@objectiv es /objective/@endpoints /Endpoint/@text	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@scheduleTimeli nes /ScheduleTimeline/@ScheduleI nstance /Timing/@value	class. Referenced values need to be replaced by actual values before submitting. Limit to 999 characters /Endpoint/@level /code/@code=C17 0559 In case of reference to the corresponding
					Timing class, check and use the referenced timing for this attribute. Limit to 254 characters.

The mapping to **Eligibility** is presented below. See Section 4.19, <u>Populations</u>, <u>Cohorts</u>, and <u>Eligibility</u> Criteria, for a description of the related features in the USDM.

CT.go v Path	CT.go v Varia ble	CT.gov Requir ement	USDM path and attribute	Required USDM relationship	Selection/Derivatio n
Eligibi lity. Sex/G ender	Sex	Require d	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plannedSex /code/@decode		Map 1 to 1 to corresponding ct.gov terminology.
Eligibi lity. Sex/G ender	Gende r Based	If applicab le	Not in USDM v3.0		ClinicalTrials.gov: "Gender means a person's self- representation of gender identity." In general, it can be decided whether this is 'No' for all trials governed by the sponsor.
Eligibi lity. Sex/G ender	Gende r Eligibi lity Descri ption		Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@criteria/ EligibilityCriteria/@text	Study/@versions /StudyVersion/@study Designs /StudyDesign/@popula tion /StudyDesignPopulatio n/@plannedSex	The eligibility text is based on the SyntaxTemplate class (see Section 4.21). Referenced values need to be replaced by actual values before submitting. Limit to 1000 characters.

CT.go v Path	CT.go v Varia ble	CT.gov Requir ement	USDM path and attribute	Required USDM relationship	Selection/Derivatio n
					Select the criterium referencing to the corresponding plannedSex value, if any.
Eligibi lity. Age Limits	Minim um Age	d	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@plannedAge / Range/@minValue		
Eligibi lity. Age Limits	Unit of Time	Require d	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@plannedAge / Range/@unit / code/@decode		Map 1 to 1 to corresponding <u>ClinicalTrials.gov</u> te rminology.
Eligibi lity. Age Limits	Maxi mum Age	Require d	RequiredStudy/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@plannedAge / Range/@maxValue		
Eligibi lity. Age Limits	Unit of Time	Require d	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@plannedAge / Range/@unit / code/@decode		Map 1 to 1 to corresponding <u>Clini</u> <u>calTrials.gov</u> terminology.
Eligibi lity	Accept s Health y Volunt eers	Require d	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@cohorts) /StudyDesignPopulation /StudyCohort/ @includesHealthySubjects		If any of the values for the StudyDesignPopulat ion or a StudyCohort is True then set to "Yes"; otherwise set to "No".
Eligibi	Eligibi lity Criteri a	Require	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyPopulation/@criteria/ EligibilityCriteria/@text		The eligibility text is based on the SyntaxTemplate class. Referenced values need to be replaced by actual values before submitting. Select limited list for submission and limit to 20000 characters.
Eligibi lity	Study Popula tion Descri ption	For observat ional studies only	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@description		Limit to 1000 characters.

CT.go v Path	v	CT.gov Requir ement	USDM path and attribute	Required USDM relationship	Selection/Derivatio n
Eligibi	Sampli	For	Not in USDM v3.0		
lity	ng	observat			
	Metho	ional			
	d	studies			
		only			

## 14.3 Use of USDM for Populating Protocol Content

A secondary aim of the USDM is to demonstrate that protocol-related content can be pulled from a reference implementation of the USDM and populated programmatically into the corresponding fields of a structured document. The TransCelerate CPT is a <u>publicly available resource</u> proposed to harmonize clinical trial protocol content in a streamlined format. The below table indicates how the USDM v3.0 (*updating to v4.0 during phase 4 of development*) content can be used to populate the structured CPT fields of CPT version v010 including the <u>CPT_BWE document</u> that is the base word template and the <u>CPT_TEE document</u> that is required to be used with the Addin.

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Page Header / Title Page	Versi on Numb er	CPT:VersionNumber	Tex t	One ToM any	Study/@versions /StudyVersion/@documentVersi on /studyProtocolDocumentVersion/ protocolVersion	Tex t, text	Sort by EffectiveDate and Version
Page Header / Title Page	Proto col ID	CPT:ProtocolID	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier		studyIdentifier/@ studyIdentifierSco pe /Organization/@or ganizationType /code/@code="C1 88724" (Clinical Study Sponsor)
Title Page	Acron ym	CPT:Acronym	Tex t	One ToO ne	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	Tex t	StudyTitle/@Typ e/Code/@decode= "Study Acronym"
Title Page	Proto col Short Title	CPT:ProtocolShortTit le	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	Tex t	StudyTitle/@Typ e/Code/@decode= "Brief Study Title"
Title Page	Proto col Title	CPT:ProtocolTitle	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@titles /StudyTitle/@Text	Tex t	StudyTitle/@Typ e/Code/@decode= "Official Study Title"

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Title Page	Amen dment Numb er	mber	Tex t	One ToO ne	Study/@versions /StudyVersion/@amendments /StudyAmendment/@number	Tex t	protocolAmendme nt: use previous attribute for sorting and take the number of last amendment
Title Page	Comp ound Numb er	CPT:CompoundNum ber	Tex t	One ToO ne	Will be added to USDM v4.0		
Title Page	Spons or Name	CPT:SponsorName	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier Scope /Organization/@name	Tex t	studyIdentifier/@ studyIdentifierSco pe /Organization/@or ganizationType /code/@code="C7 0793" (Clinical Study Sponsor)
Title Page	Spons or Legal Addre ss	CPT:SponsorLegalAd dress	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@studyIdentifier Scope /Organization/@legalAddress /Address/@text+@line+@district + @city+@postalCode+@state	Tex t	studyIdentifier/@ studyIdentifierSco pe /Organization/@or ganizationType /code/@code="C7 0793" (Clinical Study Sponsor)
Title Page	Study Phase	CPT:StudyPhase	Ch oic e	vs.C odeL ist	Study/@versions /StudyVersion/@studyPhase /AliasCode/@standardCode /code/@decode	Co ded val ue	Retrieve decode Value from standardCode element. Transform into CPT master code value
Title Page / Synopsis	Blindi ng		Tex t	ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@blindingSchema / code/@decode	Co ded val ue	
Title Page / Synopsis	Prima ry Purpo se	CPT:PrimaryPurpose	Tex t	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@trialIntentTypes / code/@decode	Co ded val ue	See CDISC SDTM extensible codelist C66736 for USDM content aligning with CPT primary purpose codes. Note that USDM and the SDTM TS domain allows for multiple values. If

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty	Map ping Typ e (CP T to	USDM Path and Attribute	US D M Fiel d Ty	Selection / Derivations
	rame		pc	USD M)		pe	
							more values are present in USDM then they need to be combined to fill Primary Purpose in CPT.
Title Page / Synopsis	Interv ention Mode 1	CPT:InterventionMod el	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@interventionMod el / code/@decode	Co ded val ue	See CDISC SDTM extensible codelist C99076 for USDM content aligning with CPT primary purpose codes.
Title Page / Synopsis	Condi tion or Disea se	CPT:ConditionDiseas e	Tex t	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@indications /indication/@name + @description	Tex t	
Title Page / Synopsis	Regul atory Agen cy ID	CPT:RegulatoryAgen cyID	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@scope /Organization/@name	Tex t	studyIdentifier/@ studyIdentifierSco pe /Organization/@or ganizationType /code/@code="C1 88863" (Regulatory Agency)
Title Page / Synopsis	Regul atory Agen cy Numb er	CPT:RegulatoryAgen cyNumber	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyIdentifiers /StudyIdentifier/@text	Tex t	StudyIdentifier/@ scope /Organization/@or ganizationType /code/@code="C1 88863" (Regulatory Agency)
Title Page / Synopsis	Pediat ric Invest igatio nal Plan Numb er	CPT:PediatricInvestig ationalPlanNumber	Tex t	One ToO ne	Study/@versions /StudyVersion/@referenceIdentifiers /ReferenceIdentifier/@text	Tex t	ReferenceIdentifi er/@type /Code/@decode=" Pediatric Investigation Plan"
Title Page / Study Populatio n	Sex of partici pants	CPT:Sexofparticipant s	Ch oic e	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plann edSex /code/@decode	Co ded val ue	Refer to CDISC codelist for Sex and corresponding eCPT mapping

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
							values in Data mapping sheet
Title Page	Proto col Appr oval Date	CPT:ApprovalDate	Tex t	One ToO ne	Study/@versions /StudyVersion/@documentVersion on /studyProtocolDocumentVersion/ @dateValues/GovernanceDate/ @dateValue	Dat e	GovernanceDate/ @type /code/@Code = C132352 ("Sponsor approval date")
List of Abbrevia tions	List of Abbre viatio ns	CPT:ListOfAbbreviat ions	Ric h Tex t	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "List of Abbreviations".
Synopsis	Ratio nale	CPT:Rationale	Ric h Tex t	One ToO ne	Study/@versions /StudyVersion/@Rationale	Tex t	
Synopsis	Numb er of Partic ipants	CPT:NumberofPartici pants	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plann edCompletionNumber /Range/@MinValue + @MaxValue	Inte ger	Combine MinValue and MaxValue. If equal then only one of both.
Synopsis	Enroll ment Targe t	CPT:EnrollmentTarg et	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population /StudyDesignPopulation/@plann edEnrollmentNumber /Range/@MinValue + @MaxValue	Inte ger	Combine MinValue and MaxValue. If equal then only one of both.
Synopsis	Numb er of Arms	CPT:NumberofArms	Tex t	Cou nt	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms		Count the number of arms defined within the study design.
Synopsis / Objective s, Endpo ints, and	Prima ry Objec tives	CPT:ObjectivesPrima ry	Ric hTe xt	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@text	Tex t	

СРТ	CPT	CPT Variable Name	CP	Map	USDM Path and Attribute	US	Selection /
Section	Varia ble Displ ay Name	(compacted)	T Va r Ty pe	ping Typ e (CP T to USD M)		D M Fiel d Ty pe	Derivations
Estimand s  Synopsis  Objective	Prima ry	CPT:EndpointsPrimar	hTe	One ToM	Study/@versions /StudyVersion/@studyDesigns	Tex t	/code/@Code =
Objective s, Endpo ints, and Estimand s	oints		xt	any	/StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text		C94496 ("Primary Endpoint") Endpoints are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of EndpointsPrimary. They can be grouped with the corresponding objective via the objective-endpoint relationship.
Synopsis / Objective s, Endpo ints, and Estimand s	Secon dary Objec tives	CPT:ObjectivesSecon dary		One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@text	Tex t	Objective/@level /code/@Code = C85827 ("Secondary Objective") Objectives are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of ObjectivesSeconda ry.
Synopsis / Objective s, Endpo ints, and	Secon dary Endp oints	CPT:EndpointsSecon dary	Ric hTe xt	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text	Tex t	

CPT	CPT	CPT Variable Name	СР	Map	USDM Path and Attribute	US	Selection /
Section	Varia ble Displ ay Name	(compacted)	T Va r Ty pe	ping Typ e (CP T to USD M)		D M Fiel d Ty pe	Derivations
Estimand							Endpoints are
S							based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of EndpointsSecondar y. They can be grouped with the corresponding objective via the objective-endpoint relationship.
Synopsis	Overa ll Desig n Syno psis	CPT:OverallDesignS ynopsis	Ric h Tex t	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s/contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or
Synopsis	Brief Sum mary	CPT:BriefSummary	Ric h Tex t	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or
Synopsis	Maski ng	CPT:Masking	Tex t	One ToM any	Study/@versions /StudyVersion/@studyDesigns/ StudyDesign/@maskingRoles /Masking/@role/code/@decode	Co ded val ue	Combine decoded role(s) if more then 1. Align CPT coded values with

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
							DDF coded values
Synopsis	Rand omly Assig ned / enroll ed	CPT:RandomlyAssig nedEnrolled	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@characteristics /code/@decode	Co ded val ue	for Masking roles.  If USDM decodes include "RANDOMIZED" then value for CPT will be randomized, otherwise depending on the study design it can be set to enrolled or assigned to investigational intervention.
Synopsis	Grou ps and Durati on	CPT:InterventionGro upsandDuration	Ric h Tex t	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Brief Summary" and it should be a child within section 1.1 with title "Study Arms and Duration". The Narrative content text may include references to the corresponding arm descriptions in the arm class and timing of the last intervention day.
Schema	Sche ma	CPT:Schema	Pict ure	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Study Rationale	Study Ratio nale	CPT:StudyRationale	Ric h Tex t	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	1.2 with title "Schema". HTML content need to include the schema as picture. Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle. For CPT the section should be 2.1 with title "Study Rationale". This may include a reference to Study/@versions /StudyVersion/@R ationale which is mapped to the rationale presented in the synopsis.
Objective s, Endpoint s, and Estimand s	Objec tives, Endp oints, and Estim ands	CPT:ObjectivesEndp ointsAndEstimands	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	and/or
Objective s, Endpoint s, and Estimand s	Tertia ry Explo ratory Objec tives	CPT:ObjectivesTertia ryExploratory	Ric hTe xt	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Objective/@text	Tex t	Objective/@level /code/@Code = C163559 ("Exploratory Objective") Objectives are based on the

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Objective s, Endpoint s, and Estimand s	Tertia ry Explo ratory Endp oints	CPT:EndpointsTertiar yExploratory	Ric hTe xt	One ToM any	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@objectives /Objective/@endpoints /Endpoint/@text	Tex t	SyntaxTemplate class. References values need to be replaced by actual values before creation of ObjectivesTertiary Exploratory.  Endpoint/@level /code/@Code = C170559 ("Exploratory Endpoint") Endpoints are based on the SyntaxTemplate class. References values need to be replaced by actual values before creation of EndpointsTertiary Exploratory. They can be grouped with the corresponding objective via the objective-endpoint relationship.
Objective s, Endpoint s, and Estimand s	ry	CPT:PrimaryEstiman		One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Estimand(s) for Primary Objective(s)" and it should be a child within section 3. The text should link to the estimands corresponding population, endpoint,

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Objective s, Endpoint s, and Estimand s	Secon dary Estim ands	CPT:SecondaryEstim ands	Ric hTe xt	One	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	intervention and intercurrent events specified in the corresponding classes.  Select content based on NarrativeContent/@sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Estimand(s) for Secondary Objective(s)" and it should be a child within section 3. The text should link to the estimands corresponding population, endpoint, intervention and intercurrent events specified in the corresponding classes.
Objective s, Endpoint s, and Estimand s	ry	CPT:TertiaryEstiman ds	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
							population, endpoint, intervention and intercurrent events specified in the corresponding classes.
Study Design	Study Desig n	CPT:StudyDesign	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	For CPT the sectionTitle should be "Study Design" with section number 4. The text may link to attributes that are stored elsewhere in the USDM and that are relevant to the study design.
Study Design	Overa II Desig n	CPT:OverallDesign	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Overall Design" with section number 4.1. The text may link to attributes that are stored elsewhere in the USDM and that are relevant to the overall design.
Study Design	Scient ific Ratio nale	CPT:ScientificRation aleforStudyDesign	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents	HT ML for mat	Select content based on NarrativeContent/ @sectionNumber

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute  /NarrativeContent/@contentItem	US D M Fiel d Ty pe	Selection / Derivations and/or
	Study Desig n				s /contentItems/@text	Tex t	@sectionTitle For CPT the sectionTitle should be "Scientific Rationale for Study Design" with section number 4.2.
Study Populatio n	Inclus ion Criter ia Age	CPT:InclusionCriteria Age	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria. The criterion text may link to Minimum and Maximum age stored in the Study Design Population or Cohort classes.
Study Populatio n	Plann ed Mini mum Age of Subje cts	CPT:PlannedMinimu mAgeofSubjects	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation  StudyCohort/@plannedAge /Range/@minValue + @unit	Tex t	Use minimum of minimum age values of all populations included (studyDesignPopul ations and Cohorts). Transform according to ISO 8601 standards. If 1 or more populations have a null minValue then TSVAL should be set to null and TSVALNF should be filled instead

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
							according to ISO 21090.
Study Populatio n	Plann ed Maxi mum Age of Subje cts	CPT:PlannedMaximu mAgeofSubjects	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation  StudyCohort/@plannedAge /Range/@maxValue + @unit	Tex t	
Study Populatio n	Inclus ion Criter ia Type of Partic ipants	CPT:InclusionCriteria TypeOfParticipant	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="I NCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Inclus ion Criter ia Weig ht	CPT:InclusionCriteria Weight	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="I NCLUSION" and /EligibilityCriterio

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
					hort/@criteria /EligibilityCriterion/@text		n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Inclus ion Criter ia Sex	CPT:InclusionCriteria Sex	t	ToO ne	Study/@versions /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	t	Select content based on EligibilityCriterion /@category/ code/@decode="I NCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria. The criterion text may link to planned Sex stored in the Study Design Population or Cohort classes.
Study Populatio n	Inclus ion Criter ia Infor med Conse nt	CPT:InclusionCriteria InformedConsent	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="I NCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Study Populatio n	Inclus ion Criter ia Other	CPT:InclusionCriteria Other	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	on/@text or a custom code may indicate the grouping of the eligibility criteria. Select content based on EligibilityCriterion /@category/ code/@decode="I NCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Exclusion Criter ia Medical Conditions	CPT:ExclusionCriteri aMedicalConditions	Tex	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	t	Select content based on EligibilityCriterion /@category/ code/@decode="E XCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Exclu sion Criter ia Liver Safety	CPT:ExclusionCriteri aLiverSafety	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="E XCLUSION" and /EligibilityCriterio

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute  hort/@criteria	US D M Fiel d Ty pe	Selection / Derivations  n/@notes
					/EligibilityCriterion/@text		The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Exclusion Criteria Prior Concomitant Thera	CPT:ExclusionCriteri aPriorConcomitantTh erapy	Tex t	One ToO ne	Study/@versions /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="E
Study Populatio n	Exclusion Criter ia Prior Concurrent Clinic al Study	CPT:ExclusionCriteri aPriorConcurrentClin icalStudy		One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="E

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Study Populatio n	Exclusion Criter ia Diagn ostic Asses sment s	CPT:ExclusionCriteri aDiagnosticAssessme nts	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	/EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Populatio n	Exclusion Criter ia Other	CPT:ExclusionCriteri aOther	Tex t	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@population (/StudyDesignPopulation/@coho rts) /StudyDesignPopulation StudyCo hort/@criteria /EligibilityCriterion/@text	HT ML for mat ted Tex t	Select content based on EligibilityCriterion /@category/ code/@decode="E XCLUSION" and /EligibilityCriterio n/@notes The CPT parameter may be indicated in the corresponding /CommentAnnotati on/@text or a custom code may indicate the grouping of the eligibility criteria.
Study Interventi ons Administ ered	Interv ention Label	CPT:InterventionLab	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@label	Tex t	
Study Interventi ons Administ ered	Interv ention Name	CPT:InterventionNa me	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@name	Tex t	

CPT Section	CPT Varia ble Displ ay	CPT Variable Name (compacted)	CP T Va r Ty	Map ping Typ e (CP	USDM Path and Attribute	US D M Fiel d	Selection / Derivations
	Name		pe	T to USD M)		Ty pe	
Study Interventi ons Administ ered	Interv ention Descr iption	CPT:InterventionDes cription	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@description	Tex t	
Study Interventi ons Administ ered	Interv ention Type		Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@type /code/@decode	Tex t	
Study Interventi ons Administ ered	ulatio n	CPT:DoseFormulatio n	Ric hTe xt		Will be added to USDM v4.0		
Study Interventi ons Administ ered	Unit Dose Stren gth	CPT:UnitDoseStrengt h	Ric hTe xt		Will be added to USDM v4.0		
Study Interventi ons Administ ered	Dosa ge Level	CPT:DosageLevel	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@administrati ons /AgentAdministration/@dose /Quantity/@value +/Quantity/@unit / code/@decode +AgentAdministration/@frequen cy/AliasCode/@standardCode/C ode/@decode	Tex t+ Co ded val ues	Combine administration strength, corresponding unit and frequency to 1 variable for CPT
Study Interventi ons Administ ered	Rout of Admi nistrat ion	CPT:RouteofAdminis tration	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@administrati ons /AgentAdministration/@route /code/@decode	Co ded val ue	
Study Interventi ons Administ ered	Use	CPT:Use	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyInterventio ns /StudyIntervention/@role / code/@decode	Co ded val ue	

<b>CPT</b> <b>Section</b>	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Study Interventi ons Administ ered	IMP and NIMP	CPT:IMPandNIMP	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@studyCells /StudyCell/@elements /StudyElement/@studyInterventi ons /StudyIntervention/@productDes ignation /code/@decode	Co ded Val ue	
Study Interventi ons Administ ered	Sourc ing	CPT:Sourcing	Ric hTe xt		Will be added to USDM v4.0		
Study Interventi ons Administ ered	Packa ging and Labeli ng	CPT:PackagingandLa beling	Ric hTe xt		Will be added to USDM v4.0		
Study Interventi ons Administ ered	Curre nt Form er Name s Aliase s	CPT:CurrentFormerN amesAliases	Ric hTe xt		Will be added to USDM v4.0		
Study Interventi ons Administ ered	Arm Name	CPT:ArmName	Ric hTe xt	One ToO ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms /StudyArm/@name	Tex t	
Study Interventi ons Administ ered	• • • • • • • • • • • • • • • • • • • •	CPT:ArmType	Ric hTe xt	ne	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms /StudyArm/@type /code/@decode	Co ded val ue	
Study Interventi ons Administ ered	Arm Descr iption	CPT:ArmDescription	Ric hTe xt	One ToO ne Man yTo One	Study/@versions /StudyVersion/@studyDesigns /StudyDesign/@arms /StudyArm/@description	Tex t	studyArmDescripti on, ArmName and Decode Value of ArmType to be sent as an arrayList in response.
Statistica 1 Consider ations	Gener al Consi derati ons	CPT:GeneralConsider ations	Ric hTe xt	One	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted	Select content based on NarrativeContent/ @sectionNumber

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
						Tex t	and/or @sectionTitle For CPT the sectionTitle should be "General Considerations" with section number 9.1.
Statistica l Consider ations	Statist ical Hypot heses	CPT:StatisticalHypot heses	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Decision Criteria/Statistical Hypotheses" with section number 9.1.1
Statistica 1 Consider ations	Popul ations for Analy ses	CPT:PopulationsFor Analyses	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Analysis Sets" with section number 9.2.
Statistica l Consider ations	Statist ical Analy ses	CPT:StatisticalAnalys	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	and/or

CPT Section	CPT Varia ble Displ ay Name	CPT Variable Name (compacted)	CP T Va r Ty pe	Map ping Typ e (CP T to USD M)	USDM Path and Attribute	US D M Fiel d Ty pe	Selection / Derivations
Statistica 1 Consider ations	Prima ry Endp oint Analy sis	tAnalysis	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Primary Endpoint(s)/Estima nd(s)" with section number 9.3.1.
Statistica 1 Consider ations	Secon dary Endp oint Analy sis	CPT:SecondaryEndp ointAnalysis	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Analyses Supporting Secondary Objective /[label]" with section number 9.4.1.
Statistica 1 Consider ations	Tertia ry Explo ratory Endp oint Analy sis	CPT:TertiaryExplorat oryEndpointAnalysis	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	and/or
Statistica 1 Consider ations	Other Safety Analy ses	CPT:OtherSafetyAnal yses	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or

CPT	CPT	<b>CPT Variable Name</b>	CP	Map	USDM Path and Attribute	US	Selection /
Section	Varia ble Displ ay Name	(compacted)	T Va r Ty pe	ping Typ e (CP T to USD M)		D M Fiel d Ty pe	Derivations
							For CPT the sectionTitle should be "/[Other] Safety Analyses" with section number 9.6.
Statistica l Consider ations	Other Analy ses	CPT:OtherAnalyses	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle For CPT the sectionTitle should be "Other Analyses" with section number 9.7.
Statistica l Consider ations	Interi m Analy ses	CPT:InterimAnalyses	Ric hTe xt	One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or
Statistica 1 Consider ations	Samp le Size Deter minat ion	CPT:SampleSizeDete rmination		One ToO ne	Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted Tex t	Select content based on NarrativeContent/ @sectionNumber and/or
Referenc es	Refer ences	CPT:References	Ric hTe xt		Study/@documentedBy /Document/@versions /DocumentVersion/@contents /NarrativeContent/@contentItem s /contentItems/@text	HT ML for mat ted	Select content based on NarrativeContent/ @sectionNumber and/or @sectionTitle

CPT	CPT	CPT Variable Name	CP	Map	USDM Path and Attribute	US	Selection /
Section	Varia	(compacted)	T	ping		D	Derivations
	ble		Va	Тур		M	
	Displ		r	e		Fiel	
	ay		Ty	(CP		d	
	Name		pe	T to		Ty	
				USD		pe	
				M)			
						Tex	For CPT the
						t	sectionTitle should
							be "References"
							with section
							number 11.

# 15 Appendices

- USDM Team
- Glossary and Abbreviations
- References
- Revision History
- Representations and Warranties, Limitations of Liability, and Disclaimers

### 15.1 USDM Team

Name	Institution/Organization
John Owen	Project Manager, CDISC
Dave Iberson-Hurst	USDM Product Owner, CDISC
Berber Snoeijer	USDM Technical Team Lead, CDISC
Erin Muhlbradt	Controlled Terminology Expert, NCI-EVS
Craig Zwickl	Controlled Terminology Expert, CDISC
Gerry Campion	Senior Software Engineer, CDISC

The USDM has been developed in partnership with TransCelerate Biopharma and Accenture. CDISC would like to acknowledge the support and input from the following groups:

- TransCelerate DDF Core Team
- TransCelerate member company subject-matter experts
- Accenture DDF development team
- CDISC DDF volunteer teams and volunteer vendor organizations

## **15.2 Glossary and Abbreviations**

The following abbreviations and terms are used in this document. Additional definitions can be found in the <u>CDISC</u> Glossary.

ADaM	Analysis Data Model
API	Application programming interface

Biomedical Research Integrated Domain Group			
Biomedical concept: A unit of biomedical knowledge created from a unique combination of			
characteristics that include implementation details like variables and terminologies, used as			
building blocks for standardized, hierarchically structured clinical research information			
Clinical Data Acquisition Standards Harmonization Project			
Clinical Data Interchange Standards Consortium			
(ICH) Clinical Electronic Structured Harmonised Protocol  "Collected" refers to information that is recorded and/or transmitted to the sponsor. This includes			
data entered by the site on CRFs/eCRFs as well as vendor data such as core lab data. This term is			
a synonym for "captured."			
(TransCelerate) Common Protocol Template			
Case report form (sometimes, case record form): A printed, optical, or electronic document			
designed to record all required information to be reported to the sponsor for each trial subject			
Controlled terminology: A finite set of values that represent the only allowed values for a data			
item. These values may be codes, text, or numeric. A codelist is a type of controlled terminology.			
Clinical Trial Registry			
Digital Data Flow (project)			
A collection of observations with a topic-specific commonality about a subject			
Electronic case report form			
Electrocardiogram			
Electronic data capture			
Electronic health record			
European Medicines Agency			
Electronic patient-reported outcome			
European Union Drug Regulating Authorities Clinical Trial Database			
(US) Food and Drug Administration			
(HL7) Fast Healthcare Interoperability Resources			
The suite of CDISC standards that describe the clinical study protocol (Protocol), design (Study			
Design), data collection (CDASH), laboratory work (Lab), analysis (ADaM), and data tabulation (SDTM and SEND)			
(NIH) Genetic and Rare Diseases Information Center			
(FDA) Geopolitical Entities, Names and Codes			
Health Level Seven International			
HyperText Markup Language			
Intercurrent events; events that occur after randomization and alter the course of the randomized			
treatment during the intended study treatment period			
International Classification of Diseases			
International Council for Harmonisation of Technical Requirements for Pharmaceuticals for			
Human Use			
JavaScript Object Notation			
Logical Observation Identifiers Names and Codes			
Medical Dictionary for Regulatory Activities. A global standard medical terminology designed to			
supersede, in regulatory submissions, other terminologies previously used in the medical product			
development process (such as COSTART and ICD9).			
Medical Subject Headings (thesaurus)			
(NIH) National Cancer Institute Enterprise Vocabulary Services  National Institutes of Health			
(NIH) National Cancer Institute Enterprise Vocabulary Services National Institutes of Health			
(NIH) National Cancer Institute Enterprise Vocabulary Services National Institutes of Health Operational Data Model			
(NIH) National Cancer Institute Enterprise Vocabulary Services  National Institutes of Health  Operational Data Model  A recipient of medical treatment			
(NIH) National Cancer Institute Enterprise Vocabulary Services  National Institutes of Health  Operational Data Model  A recipient of medical treatment  Portable data format			
(NIH) National Cancer Institute Enterprise Vocabulary Services  National Institutes of Health  Operational Data Model  A recipient of medical treatment  Portable data format  Personal health record			
(NIH) National Cancer Institute Enterprise Vocabulary Services  National Institutes of Health  Operational Data Model  A recipient of medical treatment  Portable data format			

## CDISC [Title] (Version n [Status])

PRO	Patient-reported outcome			
SDM-XML	Study/Trial Design Model in XML			
SDR	Study Definitions Repository			
SDTM	Study Data Tabulation Model			
SDTMIG	SDTM Implementation Guide (for Human Clinical Trials)			
SEND	Standard for the Exchange of Nonclinical Data			
SME	Subject-matter expert			
SNOMED	Systemized Nomenclature of Medicine			
SOA	Schedule of activities			
SSU	Study start-up			
Subject	A participant in a study			
UML	Unified modeling language			
USDM	United Study Definitions Model			
USDM-IG	USDM Implementation Guide			
UUID	Universally unique identifier			
WHO	World Health Organization			
XML	Extensible markup language			

### 15.3 References

- 1. National Cancer Institute. About BRIDG. Accessed June 22, 2023. https://bridgmodel.nci.nih.gov
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- 3. International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use. *Guideline for Industry. Structure and Content of Clinical Study Reports* (ICH E3). July 1996. Accessed June 21, 2023. https://www.fda.gov/media/71271/download
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- 5. European Medicines Agency. *ICH E9 (R1) addendum on estimands and sensitivity analysis in clinical trials to the guideline on statistical principles for clinical trials*. February 17, 2020. Accessed January 5, 2024. <a href="https://www.ema.europa.eu/en/documents/scientific-guideline/ich-e9-r1-addendum-estimands-and-sensitivity-analysis-clinical-trials-guideline-statistical-principles-clinical-trials-step-5_en.pdf">https://www.ema.europa.eu/en/documents/scientific-guideline/ich-e9-r1-addendum-estimands-and-sensitivity-analysis-clinical-trials-guideline-statistical-principles-clinical-trials-step-5_en.pdf</a>

## **15.4 Revision History**

### 15.4.1 USDM Implementation Guide

The USDM v1.0 was released as part of the DDF Reference Architecture in August 2022. Version v1.0 of the USDM has no associated implementation guide therefore there is no revision history for the Implementation Guide. The first version of the USDMIG is therefore v2.0. This section details the changes made to the USDMIG between v2.0 and v3.0.

#### 15.4.2 USDMIG Amendments between USDM v3.0 and USDM v4.0

#	Release	Overview	Notes
	#		
1	3.2	UML update for Arms and Epochs section	Name of encounter attribute environmentalSetting changed to environmentalSettings
			<ul> <li>Added notes attributes to Encounter, StudyArm, StudyElement and StudyEpoch</li> </ul>
			classes
2		UML update for <b>Study Timing</b> section	<ul> <li>Moved relationships timeline and timelineExit.</li> </ul>
			Name of encounter attribute environmentalSetting changed to environmentalSettings.
			Added description for encounter timing - scheduledAt
3		UML and text update for <b>Populations</b> , Cohorts,	Added relationship criteria from Study Version to Eligibility Criterion.
		and Eligibility Criteria section	Changed criteria cardinality from PopulationDefinition to EligibilityCriterion from
			1* to 0* in UML.

#	Release #	Overview	Notes
4		UML update for Study, Protocols, and Amendments section	<ul> <li>Added notes attributes to PopulationDefinition, SyntaxTemplate, Indication, StudyArm, StudyDesign and StudyVersion classes.</li> <li>Updated text accordingly to specify that criteria should either be referenced from Study Population or from Study Cohort.</li> <li>Updated text regarding eligibility criteria: removed reference to context attribute and specify that they are defined within a study version.</li> <li>Added explanation of previous/next criteria</li> <li>Added notes attributes to StudyVersion and StudyDesign classes.</li> </ul>
5	1	UML update for Study Identifiers and Titles	Added notes attribute to StudyVersion class.
6		UML and text update for Activities section	<ul> <li>Added notes attribute to Study Version class.</li> <li>Added notes attribute to Activity, Procedure, BiomedicalConcept, BiomedicalConceptSurrogate, BiomedicalConceptCategory, and BiomedicalConceptProperty classes.</li> <li>Added ScheduleTimeline class to the UML view</li> <li>Explained the use of timeline attribute in the Activity class</li> </ul>
7		UML update for <u>Study Interventions</u> section	<ul> <li>Added notes attributes to StudyIntervention and AgentAdministration classes.</li> </ul>
8		UML update for <u>Study Objectives and Endpoints</u> section	<ul> <li>Added notes attributes to Estimand, AnalysisPopulation, IntercurrentEvent, StudyIntervention and SyntaxTemplate classes.</li> <li>Added name, description and label to Estimand class</li> </ul>
9		UML update for Syntax Templates section	Added notes attribute to SyntaxTemplate class.
10	3.3	UML and text update for <u>Activities</u> section	<ul> <li>Added children attribute to Activity class</li> <li>Added example to explain how SoA activities are stored in the Activity class with respect to the previous, next and children attributes.</li> </ul>
11		UML and text update for for Study Timing section	<ul> <li>Changed cardinality for relativeFromScheduleInstance relationship</li> <li>Added corresponding text for anchors relativeToScheduleInstance relationship should be equal to relativeFromScheduleInstance or missing.</li> </ul>
12	3.4	Updated <u>CPT mapping</u> section for version 3,0 and further alignment	
13		Updated <u>Unstructured Content</u> section to include multiple template support	<ul> <li>Added new UML view for documents.</li> <li>Adjusted text to include new NarrativeContentItem and reusability of text across documents.</li> </ul>
14		Updated <u>Study, Protocols, and Amendments</u> section to include multiple template support	<ul><li>Updated UML.</li><li>Adjusted text to refer to the right classes.</li></ul>
15	3.5	Updated Study, Protocols, and Amendments section to include abbreviations	<ul> <li>Updated UML.</li> <li>Added text to explain the use of the new abbreviation class and corresponding attributes.</li> </ul>

#	Release #	Overview	Notes
16	3.6	Created <u>Abbreviations</u> section to give examples of how it can be used.	Created new section with examples.
17		Updated <u>Study, Protocols, and Amendments</u> section.	Created cross-reference to Abbreviations section.
18		Updated XHTML Attributes section.	Referred to NarrativeContentItem instead of NarrativeContent.
19		Updated <u>Study Identifiers and Titles</u> section.	Updated UML to include inheritance of identifier class and to add reference identifiers.  Here the standard for the stan
20		Undeted the of HCDM for Develoting Ductoral	Updated text to add explanation of reference identifiers.
20		Updated <u>Use of USDM for Populating Protocol</u> <u>Content</u> section	<ul> <li>Include mapping to pediatric investigational plan number.</li> <li>Updated mappings based on changed attribute names.</li> </ul>
21		Updated <u>Study Interventions</u> section	<ul> <li>Updated UML to include all changes for the new model version.</li> <li>Updated explanation of the model and included some references to IDMP.</li> </ul>
22		Updated <u>Controlled Terminology</u> section	Small tweak to section on AliasCode to clarify that standard value sets do not have to be CDISC code lists.
23		Updated Populations, Cohorts, and Eligibility Criteria section	<ul> <li>Updated UML to include small change on plannedSex relationship.</li> <li>Updated text to explain the use of plannedSex (use Male and/or Female).</li> </ul>
24		Updated <u>Study Roles and Organizations</u> section	<ul> <li>Changed section name from 'Organizations' to 'Study Roles and Organizations'.</li> <li>Updated UML to include significant changes in the model.</li> <li>updated text to explain this part of the model and expected use.</li> </ul>

### 15.4.3 USDMIG Amendments between USDM v2.0 and USDM v3.0

#		Overview	Notes
	#		
1	2.1	Created Naming Conventions section	1. This section details the conventions used for naming and the use of attribute datatypes
		-	2. To support model split and element renaming
2		Edits to Internal Identifiers Within the	1. To support model split and element renaming
		Model	Click here to see changes
			Versions Compared  1 Current Join Course J
			View Page History  The USEM normative form is a unified modeling language (MA) model. Each class defined within the UAL has an identification attribute that can be used to provide a unique identifier for an instance of the class. The identifier
			should be unique and self-consistent within the scope of a version of a study, No attempt is made to defer the from type, or structure of those identifiers the attributes are defend as a defined as a finite or the study for the structure of the self-interest processing or the study for the structure of the study for the st
			The boarder enhances are interned to extract a desired and the first of the state o
			for example, for the forecounter case all instances are included from the Study Resign class using the attribute
			shereas the Studypoint references the indiance using the attribute
			— The only exception is the identifier at the head of the model within the Study class. Implementations are free to allocate the value to this field using, for example, a UUID, to ensure uniqueness within the implementation.

#	Release	Overview	Notes
3		Edits to Overview	1. To support model split and element renaming Click here to see changes  Versions Compared  1. Current John Dave Bernion Hand John Compared  Ver Page Holder  Ver Page Holder  The USCM recentling was changed.  The USCM recentling was changed.  The USCM recentling was changed.
4		Edits to USDM API	1. To support model split and element renaming Click here to see changes Versions Compared  2
5		UML Split Model and Model Naming Changes	<ul> <li>Replaced all String Id references in the UML to instances of the class.</li> <li>Changed all class properties for Id, Name and Description to consistent across the model. Removed the class name prefix from these properties.</li> </ul>
6	2.3	Added <u>Unstructured Content</u> section to the USDM Features	Added new section for unstructured content  1. This section introduces the content class that is used to store unstructured narrative content.
7		Add Syntax Templates" section to the USDM Features	<ol> <li>This section introduces the classes that enable syntax text templates</li> <li>It explains the how the syntax text templates can be used in the USDM</li> <li>It explains how references can be made to data elements stored elsewhere in the data model.</li> <li>It gives examples of text templates and corresponding examples.</li> </ol>
8		Added label to Naming Conventions section.	
9	2.4	Change class name "Content" to "Narrative Content" in the <u>Unstructured</u> <u>Content</u> section of USDM Features	
10	2.8	Update to Controlled Terminology section	Added detail on standard codes and alias code

#	Release #	Overview	Notes
11	π	inserted Principles section	Added notes on principles. Needs further work
12		Update to API section	Improved text within API section and added details re the "instanceType" attribute
13		Update to Arms and Epoch section	Small updates to text, inserted UML and added links to related pages.
14		Update to Activities section	Small updates to text, inserted UML, added conditional class information and added links to related
		<u> </u>	pages.
15		Update to Study Population section	Updates to text in accordance with model changes, added UML and cohort and eligibility description.
16		Update to Intervention section	updates to text in accordance with model changes and added UML
17		Added new section Addressing Footnotes	identified 12 types of footnotes and describing how they can be included in the USDM
18		Updated section Study Timing	Added UML, updated text and timeline figures
19		Updated section Relationship to Other	Moved mapping to SDTM trial summary domains to Creation of SDTM Trial Design Domains
		CDISC Standards	
20		Updated <u>USDM Team</u>	Updated <u>USDM Team</u> page to include the latest team members for USDM v3.0
21		Added Creation of SDTM Trial Design	
		<u>Domains</u>	
22		Updated Study, Version, Identifier	Changed title to Study, Protocols, and Amendments. Added UML and description of protocol and
		section	amendment versions.
			Identifiers will be handled in new section.
23		Updated Syntax Templates	Updated content according to html reference style
24		Added Study Identifiers and Titles	Moved description of Study Identifiers here and added Titles description
25		Updated <u>Procedures</u>	Added reference to study intervention. Removed conditionality which is described more general for
			all related classes in <u>Activities</u>
26 27		Updated <u>Indications</u>	Added description of new attribute isRareDisease
27		Updated Study Objectives and Endpoints	Inserted UML and reference to syntax template class
28		Updated <u>Study Estimands</u>	updated reference names
29	2.9	Updated Fundamentals of the USDM	Added information on v3.0
30 31		Updated Arms and Epochs	Added link to Creation of SDTM Trial Design Domains
31		Updated <u>Study Timing</u>	Replaced UML based on changed relationship to timing class. Some minor textual changes.
32		Updated Study Objectives and Endpoints	Replaced UML based on changed reference name from Estimand to studyIntervention class.
33		Updated Populations, Cohorts, and	Replaced UML based on chanced name of EligibilityCriterion class and small textual updates.
		Eligibility Criteria	
34		Updated <u>Use of USDM for Populating</u>	Adapted the POC mapping to v3.0 of USDM. No additional variables are mapped based on new
		<u>Protocol Content</u>	features of USDM v3.0. This is indicated in the introduction.
35		Updated Study, Protocols, and	Removed study site information from UML and descriptions. Moved to new paragraph: <u>Study Roles</u>
		Amendments	and Organizations
36		Added Study Roles and Organizations	Added UML and description of Organization class and corresponding research Organization and sites.
	2.11	Updated Syntax Templates	Updated content requirements based on current reference strategy and JIRA comments.
38		Updated <u>Arms and Epochs</u>	Updated UML based on new version of ScheduleInstance class.

#	Release	Overview	Notes
	#		
39		Updated Study Timing	Updated UML based on new ConditionAssignment class and updates in Timing class. Updated
			corresponding text.
40		Updated <u>Study Interventions</u>	Updated UML based on Jira tickets of public review. This includes cardinality updates and adding the
			option to add alias codes for unit, route and frequency.
41		Updated Study Objectives and Endpoints	Updated UML since objective level is required. Added option of exploratory objectives in the text.
42		Updated Populations, Cohorts, and	Updated UML for plannedSex. Added requirement that plannedSex, plannedAge and
		Eligibility Criteria	plannedEnrollment or plannedCompletion number should be either filled at the
			studyDesignPopulation level or the studyCohort level.
43		Update to API section	Updated API to include initial rules for the minimum content to be included within the data sent via
			the API. Also added details with regard to the root attributes that includes the USDM version.
44		Updated Naming Conventions	Updated to reflect latest practice
45		Inserted XHTML Attributes	Inserted new section on XHTML attributes
46		Updated Biomedical Concepts	Updated to include more details on enabled and required flags
47		Updated <u>Unstructured Content</u>	Updated to refer to XHTML attributes paragraph
48		Updated Study Roles and Organizations	Updated UML - included AliasCode class

## 15.4.4 Amendments between USDM v1.0 and USDM v2.0 (UML, CT, API)

The following table lists at a high level the major changes that occurred between USDM v1.0 and USDM v2.0

		Overview	Notes
	#		
1	1	Bugfixes and review comments from DDF Phase I	1. StudyEpoch Class: Add encounters relationship, 1 -> 0*
			2. IntercurrentEvent Class: strategy attribute rename to
			"intercurrentEventStrategy" and is of type String
			3. PointInTime Class: remove from the model
			4. Encounter Class Attributes "startRule" and "endRule" should be renames
			and prefixed with "transition", so "transitionStartRule", "transitionEndRule"
			5. Workflow Class Attribute "workflowId" renamed to "uuid"
			6. Estimand Class Attribute "variableOfInterest" type should be Endpoint not
			Encounter
2	1	Addition of Therapeutic Area	Class: Study Attribute businessTherapeuticArea
			Class: StudyDesign Attribute therapeuticAreas
3	1	Allow for multiple trial types entries on the StudyDesign	Class StudyDesign Attribute trialType amended to a list
		class	
4	2	Terminology Flexibility	Code and CodeAlias classes added to the model
5	2	Addition of name and description for StudyDesign class	Class: StudyDesign Attribute studyDesignName
			Class: StudyDesign Attribute studyDesignDescription

#	Sprint #	Overview	Notes
7	3	Attribute name changes	Class: Study Attribute: studyIdentifier amended to studyIdentifiers     Class: Study Attribute: studyProtocolVersion amended to studyProtocolVersions     Class: Study Attribute: studyDesign amended to studyDesigns
9	3	Visit Contact Mode	Not sure what has changed here
10		Allow Study Phase to use the Code Alias	Class: Study Attribute studyPhase amended from Code to AliasCode
10	4	Add flag for Activity and Procedures being optional	<ol> <li>Class: Activity Attribute activityIsOptional added</li> <li>Class: Procedure Attribute procedureIsOptional added</li> <li>Also see additional change to 16 below</li> </ol>
12	5	Additional elements added in to support eCPT population	Class: Study Attribute; studyRationale added     Class: Study Attribute: studyAcronym added     Class: StudyDesignPopulation Attribute: plannedNumberOfParticipants added     Class: StudyDesignPopulation Attribute: plannedMaximumAgeOfParticipants added     Class: StudyDesignPopulation Attribute: plannedMinimumAgeOfParticipants added     Class: StudyDesignPopulation Attribute: sexOfParticipants added     Class: StudyDesignPopulation Attribute: sexOfParticipants added     Class: StudyDesign Attribute: studyDesignRationale added     Class: Organization Attribute: organizationLegalAddress added
15	6	New class for Address	Class: Address added with the following attributes  Text Line City District State Postal Code Country
16	6	Amend activityIsOptional and procedureIsOptional to conditional	Class: Activity Attribute activityIsOptional amended to activityIsConditional     Class: Procedure Attribute procedureIsOptional amended to procedureIsConditional
17	6	Addition of TBLIND/Trial Blinding Schema (valid values in codelist C66735) code to studyDesignBlindingScheme	Class: StudyDesign Attribute studyDesignBlindingScheme codelist     TBLIND added
19	7	Biomedical Concepts sub model added	See <u>Biomedical Concepts</u> section for additional information.  Addition of the following Classes (note that class StudyData was removed and replaced with the Biomedical Concept classes

#	Sprint #	Overview	Notes
	π		BiomedicalConcept
			BioemdcialConceptProperty
			ResponseCode
			BiomedicalConceptCategory
			BiomedicalConceptSurrogate
20	9	Study Timing and "Timepoints" added to the model	See Study Timing section for additional information.
			Addition of the following Classes (note that class StudyData was removed and
			replaced with the Biomedical Concept classes
			ScheduleTimeline
			Timing
			ScheduledInstance
			ScheduledDecisionInstance
			ScheduledActivityInstance
			ScheduleTimelineExit
21	11	Internal Review Sprint Changes	API only: studyStudyDesignPopulations changed to studyPopulations
			StudyEpoch.encounters type List <encounter> Amended to</encounter>
			StudyEpoch.encounter <b>Ids</b> type List< <b>String</b> >
			StudyEpoch.trialIntentType type List <code> Amended to</code>
			StudyEpoch.trialIntentTypes type List <code></code>
			Procedure.procedureName type String Added
			Procedure.procedureDescription type String Added
22	11-14	Public Review Sprint Changes	StudyEpoch.encounters type List <encounter> changed to</encounter>
			StudyEpoch.encounterIds type List <string></string>
			StudyDesign.trialIntentType type List <code> changed to</code>
			StudyDesign.trialIntentTypes type List <code></code>
			Procedure.procedureDescription type String added
			Procedure.procedureName type String added

As part of the v2.0 updates, the elements of the RA (USDM, CT, API, and IG) are stored within a <u>Github repository</u> and version managed as a series of releases corresponding to the sprints, a subsequent release for internal review, a release for public review, and a release for the final publication as v2.0.

- **Controlled Terminology:** For a complete list of controlled terminology changes between <u>USDM v1.0</u> and the public review version, see the USDM_CT_Changes.xlsx file in the <u>controlled terminology deliverable folder</u>.
- **UML:** A list of changes to the UML model between USDM v2.0 and the public review version can be found <u>here</u>. A list of model changes between Internal Review and Public Review can be found <u>here</u>. A list of changes between Public Review and Publication can be found <u>here</u>.
- **API:** For a complete list of API changes between USDM v1.0 and USDM v2.0, use a file-comparison tool to compare the API from <u>USDM v1.0</u> and the API for <u>USDM v2.0</u>. Please refer to the USDM API.yaml files in the API deliverable folder.

#### 15.4.5 Amendments between USDM v2.0 and USDM v3.0

- **Controlled Terminology:** For a complete list of controlled terminology changes between USDM v2.0 and the public review version, see the USDM_CT_Changes.xlsx file in the <u>controlled terminology deliverable folder</u>.
- UML: A list of changes to the UML model between USDM v2.0 and the public review version can be found here.
- **API:** For a complete list of API changes between USDM v2.0 and USDM v3.0, use a file-comparison tool to compare the API from <u>USDM v2.0</u>. and the API for <u>USDM v3.0</u> Please refer to the USDM API.yaml files in the API deliverable folder.

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