Centers for Medicare & Medicaid Services
Office of the National Coordinator for Health Information Technology

Quality Data Model, Version 5.4

FOR USE BY MEASURE DEVELOPERS CREATING MEASURES USING CLINICAL QUALITY LANGUAGE (CQL)

June 2018
# Record of Changes

<table>
<thead>
<tr>
<th>QDM Version</th>
<th>Date</th>
<th>Author / Owner</th>
<th>Description of Change</th>
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<tr>
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<td>The MITRE Corporation</td>
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<td>ESAC Inc.</td>
<td>Updated to align with emerging standards (HL7 FHIR) and add increase explicit capabilities.</td>
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1. Introduction

1.1 Background

In 2009, the National Quality Forum convened the Health Information Technology Expert Panel, which established the Quality Data Model (QDM) to enable electronic clinical quality expressions for measurement. The QDM was developed at the request of the American Health Information Community and the Office of the National Coordinator for Health Information Technology (ONC), with funding from the Agency for Healthcare Research and Quality (AHRQ). January 1, 2014, responsibility for maintenance and evolution of the QDM transitioned to the Centers for Medicare & Medicaid Services (CMS) and in conjunction with its federal partner, the ONC, assumed responsibility for its maintenance and evolution. ESAC, Inc. currently manages the QDM under the direction of CMS. Previously published versions of QDM included the data model (i.e., how to specify the information needed for the measure) and the logic required to compare one data element with another. Beginning with QDM version 5.0, the QDM includes only the data model. This and future versions require a separate method for expressing logic, Clinical Quality Language (CQL). Readers can access information about CQL at the eCQI Resource Center (https://ecqi.healthit.gov/).

1.2 Purpose

The QDM describes clinical concepts in a standardized format to enable electronic quality performance measurement in support of operationalizing the Meaningful Use Program of the Health Information Technology for Economic and Clinical Health Act. This model is the backbone for representing criteria used in quality measures by stakeholders involved in electronic quality measurement development and reporting. Stakeholders of the QDM include measure developers, federal agencies, Health Information Technology (HIT) vendors, standards organizations, informatics experts, providers, and researchers.

The QDM is intended to enable automation of structured data captured through routine care in electronic health records (EHR), personal health records (PHR), and other electronic clinical sources. It provides a structure for describing clinical concepts contained within quality measures in a standardized format, allowing individuals (e.g., providers, researchers, or measure developers) who monitor clinical performance and outcomes to communicate information concisely and consistently.

The QDM is one of several standards in the broader electronic Clinical Quality Improvement (eCQI) landscape and operates concurrently with changing measure concepts, tools, and other standards for electronically representing quality measures. The QDM is pivotal in the electronic Clinical Quality Measures (eCQM) ecosystem to encourage improved quality of the measure outputs. The QDM conceptual data model allows measure developers to express information for eCQMs selected for use in CMS quality reporting programs. The QDM 5.3 represents a significant change from prior production versions as it no longer contains any logic expression. Clinical Quality Language (CQL) replaces the logic contained in prior versions of QDM.1 QDM

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1 Clinical Quality Language (CQL) information is available from the eCQI Resource Center at: https://ecqi.healthit.gov/cql.
5.4 adds some clarity and removes some attributes that were inconsistent with clinical data capture. Note that the deleted attributes have not been used in CMS program measures to date. The Measure Authoring Tool\(^2\) (MAT) implements the QDM specification, thus providing a software tool for measure developers to author their measures in standard formats.

### 1.3 Vision

The QDM must adapt and evolve to facilitate the introduction of quality measurement and feedback into a provider’s daily routine. It must adapt as eCQMs advance from process- to outcomes-based measures and increasingly use patient-reported data. The QDM must evolve to align Clinical Decision Support (CDS) rules with quality measurement. As the QDM changes to enable expression of new measure types and CDS rules, its updates must be capable of incorporation into the MAT, where needed, and be compatible with quality and CDS standards. To meet this end, the QDM User Group and the HL7 Clinical Quality Information Workgroup carefully addressed each QDM datatype and its associated attributes to assess consistency with data to be expected from EHRs in interoperability messages. Most of the work mapped QDM details to the FHIR Quality Improvement Core (QI Core).\(^3\)

### 1.4 Scope

The QDM version 5.4 specification is to be used in conjunction with Clinical Quality Language (CQL) version 1 which provides the ability to express logic that is human readable yet structured enough for electronically processing a query. QDM version 5.4 aims to accomplish the following:

- Provide the method for describing individual data elements within a measure that can be expressed using CQL to specify measure criteria; and
- Provide continuity from prior published versions of the QDM.

### 1.5 Audience

The audience includes all stakeholders responsible for translating Clinical Quality Measures into electronic specifications. They include, but are not limited to, measure developers, the MAT development team, EHR vendors, providers reporting eCQMs, and the community developing health IT standards.

### 1.6 Document Organization

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<td>Section 2: Data Model</td>
<td>Describes the general data model for the QDM</td>
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<td>Section 3: Timing Considerations</td>
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\(^3\) FHIR QI Core version 3.0 is available at: [http://build.fhir.org/ig/cqframework/qi-core/index.html](http://build.fhir.org/ig/cqframework/qi-core/index.html).
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2. Data Model

2.1 QDM Basics

The QDM consists of criteria for data elements, relationships for relating data element criteria to each other, and functions for filtering criteria to the subset of data elements that are of interest. The following sections describe the different components of the QDM.

2.2 Category

A category consists of a single clinical concept identified by a value set. A category is the highest level of definition for a QDM element. The QDM currently contains 21 categories. Some examples of categories are Medication, Procedure, Condition/Diagnosis/Problem, Communication, and Encounter.

2.3 Datatype

A datatype is the context in which each category is used to describe a part of the clinical care process. Examples of datatypes include ‘Medication, Active’ and ‘Medication, Administered’ as applied to the Medication category. Each category contains from one to five datatypes.

2.4 Attribute

An attribute provides specific detail about a QDM element. QDM elements have two types of attributes, datatype-specific and data flow attributes. Healthcare interoperability standards generally use the term metadata to refer to attributes as defined in QDM. Metadata in this context is information about the data that more specifically defines the details required to calculate the measure requirements.

2.4.1 Datatype-Specific Attributes

Datatype-specific attributes provide detail about a QDM element based on its datatype. For example, Medication, Dispensed, Medication, Ordered and Medication Administered all contain information about dosage, supply, frequency and route. Medication, Dispensed and Medication Ordered include the attribute refills, but Medication, Administered does not. Because these attributes pertain to specific datatypes, they are called datatype-specific attributes.

2.4.2 Data Flow Attributes

Data flow attributes provide specific detail about the location of data represented by a QDM element. In order to identify the authoritative source of an element in a particular use case, the electronic record requires metadata about the origin, or provenance of the information.⁴ For example, a diabetes medication order may be found in medication orders, whereas allergies

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⁴ Provenance refers to attributes about the origin/source of health information at the time it is first created or tracks the uses and alterations of the information over its lifecycle. [Chua JA, ONC Privacy & Security Update: Data Provenance (DPROV) S&I Initiative, NCVHS Meeting, September 22, 2014. Available at: https://www.ncvhs.hhs.gov/wp-content/uploads/2014/09/140922p2.pdf.]
related to diabetes medication will be on the allergy list. Similarly, a clinician’s account of an allergy may be found in an EHR allergy list, but a patient’s account of an allergy will be found in a PHR allergy list. Data flow attributes allow a measure developer to clearly define the location of the data so that the intended meaning can be achieved. The following data flow attributes apply to all QDM elements.

- **Health Record Field**: The location within an electronic record where the data should be found.
- **Source**: The originator of the quality data element. The source may be an individual or a device.
- **Recorder**: The individual or device that enters the data element into a health record field. The desired recorder also may be, but is not necessarily, the source of the data.

Users should note that including data flow attributes in measures requires careful evaluation of feasibility as the required data provenance details are not necessarily available in real-world settings.

### 2.5 QDM Element

A QDM element encapsulates a certain category with an associated datatype. It is a discrete unit of information used in quality measurement to describe part of the clinical care process, including a clinical entity and its context of use. It can include criteria for any relevant metadata about a clinical or administrative concept relevant to quality measurement. A QDM element provides an unambiguous definition and enables consistent capture and use of data for quality measurement. It may be defined for any given measure and reused when the same information is required for another measure. Reuse encourages standardization of quality measures and reduces the generation of additional software requirements for every new measure. Starting with QDM 5.3 and continued through QDM 5.4, the tooling used to create an eCQM, the Measure Authoring Tool (MAT), modifies the method by which a measure developer defines components of the QDM data element. However, the QDM data model structure remains the same. Figure 1 identifies the terminology and gives examples of usage for the basic components that constitute a QDM element.

---

With QDM versions through and including version 4.3, the measure developer specified the entire QDM data element (including the attributes) directly in the measure standard called Health Quality Measure Format (HQMF). Figure 2 shows an example of a QDM data element specifying that only high density lipoprotein laboratory tests with results less than 40 mg/dL are relevant to the measure.

Starting with QDM version 5.3 and continued through QDM 5.4, the part of the QDM data element defined up front in the HQMF includes the only the category and context as represented by the datatype, and the value set or a direct referenced code\(^6\). The request to clinical software to retrieve information can include any of the potential attributes allowed by the QDM datatype as

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\(^6\) Direct referenced code is defined in Section 2.7, Value Sets.
specified in the QDM model. Requesting a retrieve of all possible attributes would be excessive and such is not the intent. Instead, the measure further specifies the QDM data element’s desired attributes (metadata) in the CQL, consistent with what the QDM 5.4 model allows. Unlike QDM logic, CQL allows expression of multiple attributes and attributes of attributes. That means the measure specifications can be shortened and more readable. Figure 3 shows the general approach to defining a QDM data element using tooling that incorporates CQL logic.

Figure 3. QDM 5.3 Data Element Definition using CQL

The retrieve request to the clinical software can include any of the potential metadata allowed by the QDM datatype as specified in the QDM model.

The two figures below show how a measure describes a QDM data element that specifies only borderline or elevated antinuclear antibody laboratory tests as part of the measure logic. Figure 4 indicates the portion of the data element defined in the HQMF portion of the measure, the datatype and value set (or direct referenced code).
Figure 4. QDM 5.4 Description of a laboratory test, specifying only the portion defined in the HQMF portion of the measure

The retrieve request to the clinical software can include any of the potential metadata allowed by the QDM datatype as specified in the QDM model.

Examples:
- LOINC — Lab tests / observable entities
- SNOMED-CT — Conditions, Procedures
- RxNorm — Medications (administered or ingredient level)

Figure 5 completes the definition of the QDM data element by specifying the required metadata in the CQL portion of the measure. In the example, the measure needs to specify only borderline or elevated results for the laboratory test, antinuclear antibody (ANA). Using CQL, the example allows specification of the two components of the ANA report (homogeneous pattern and speckled pattern). The CQL also allows the measure to indicate that one or both of the components’ results must meet the criteria (result >= 1:80). Earlier versions of QDM using QDM logic required a more complex description for the same specification.

Appendices A.1.1 and A.1.3 provide further description about using components and results in QDM 5.3. Appendix A.2 includes guidance regarding the use of result thresholds as attributes.
Figure 5. QDM 5.4 Description of a laboratory test, specifying the metadata in the CQL to assure retrieval of only borderline or elevated results of either one of two antinuclear antibody (ANA) components’ tests (homogeneous pattern and speckled pattern). Note the light blue color indicates the portion expressed in the HQMF and the peach color indicates the portion of the data element expressed with CQL.

To identify all patients with borderline or elevated Antinuclear Antibody (ANA)

- **HQMF**
  - Laboratory Test, Performed: Antinuclear Antibody (ANA)
  - Uses a LOINC value set for ANA

- **CQL**
  - Union of:
    - Component: Homogeneous Pattern (result >= 1:80)
    - Component: Speckled Pattern (result >= 1:80)
  - Uses LOINC value sets for "homogeneous pattern" and "speckled pattern"

The two figures below provide another example of a QDM data element using HQMF and the CQL portion of the measure. This example specifies the length of stay for an encounter, and also the arrival time. By convention, an Encounter, Performed start and stop times reference the admission and discharge times. These times are referenced by periods (the interval between start and stop times). To determine arrival time requires specification of the location attribute, i.e., where, within the scope of the encounter, the patient was treated. The location period defines the arrival (start) and departure (stop) times for each location specified. Figure 6 indicates the portion of the data element defined in the HQMF portion of the measure, the datatype and value set (or direct referenced code).

Appendix A.1.2 describes the use of facility location attributes in greater detail.

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7 Periods, the interval between start and stop times for a QDM data element are described in greater detail in Section 3.

8 See Section 4.1 for definitions of attributes
Figure 6. QDM 5.4 Description of and encounter (Encounter, Performed), specifying only the portion defined in the HQMF portion of the measure

The retrieve request to the clinical software can include any of the potential metadata allowed by the QDM datatype as specified in the QDM model.

Examples of all potential metadata:
- Encounter, Performed
- Relevant Period (Administrative start/stop times)
- Admission Source
- Diagnosis
- Discharge Disposition
- Location Period (Physical arrival/departure times)
- Length of Stay
- Negation Rationale
- Principal Diagnosis
- Author Date/Time
- Code
- Facility Locations (may appear 0 or many times) (Each component will have: code, facility location)

Dataflow attributes:
- Source
- Recorder
- Health Record Field

Figure 7 completes the definition of the QDM data element by specifying the required metadata in the CQL portion of the measure. In the example, the measure needs to specify two things about the Encounter, the length of stay and the arrival time. The measure defines length of stay as the difference between discharge time and admission time. These are more administrative concepts, addressing the times used to determine the number of days during which a patient receives treatment in the hospital. A measure may need to be more specific about when a patient physically arrives at or departs from an encounter setting. To determine physical arrival and departure, the measure must define the facility locations component to differentiate between the physical start and stop times from the administrative timings. Since the patient may have arrived through the emergency department (ED) or as a direct admission to the hospital in a medical-surgical or ICU location, the different locations need to be addressed as components of the Encounter, Performed. Earlier versions of QDM using QDM logic required a more complex description for the same specification. Appendix A.1 describes the use of facility locations attributes in greater detail.
Figure 7. QDM 5.4 Description of an Encounter, Performed specifying the metadata in the CQL to determine length of stay (based on relevant period – admission to discharge time) and arrival time (based on facility location components, each with its respective location period – arrival to discharge). Note the light blue color indicates the portion expressed in the HQMF and the peach color indicates the portion of the data element expressed with CQL.

### 2.6 Code System

A code system is a collection of coded concepts with definitions from a particular taxonomy, vocabulary, or classification system. Concepts from a code system are used in value sets. Specific code systems are used in applying the QDM to quality measures based on the recommendations of the HIT Standards Committee of the ONC and established certification rules for meaningful use. For example, International Classification of Diseases, Ninth Revision (ICD-9-CM), International Classification of Diseases, Tenth Revision (ICD-10), Systematized Nomenclature of Medicine – Clinical Terms (SNOMED-CT®), and Current Procedural Terminology (CPT®) are examples of code systems. The concept of diabetes may be described in the QDM with ICD-9-CM, ICD-10, and/or SNOMED-CT®. Refer to the Measures Management Blueprint for a complete review of code systems that should be used for each QDM datatype and attribute.

### 2.7 Value Sets

A value set is a list of specific values (terms and their codes) derived from single or multiple standard vocabularies (or code systems) used to define clinical concepts (e.g., patients with diabetes, clinical visits, reportable diseases) used in quality measures and to support effective health information exchange. Value sets are used to define the set of codes that can possibly be found in a patient record for a particular concept, or to which codes used locally can be mapped. In QDM elements, value sets can be used to define possible codes for the QDM element’s category or the QDM element’s attributes. The National Library of Medicine Value Set Authority Center provides measure developers with the ability to create new value sets or re-use
value sets previously developed and associated with published eCQMs. Note, beginning with QDM 5.3 and continued through QDM 5.4, measure developers will be able to specify individual codes to describe QDM elements rather than creating value sets.

Initially all data elements in eCQMs used value sets to identify the appropriate code(s) to be used. In some cases, the value set contained only a single code. The “single code value sets” are concerning because they create an alternative identifier (the value set identifier) for the code system concept. The practice of creating “single code value sets” also creates additional work for the eCQM implementer to unpack the value set rather than just use the code as specified. There are situations where a “single code value set” is appropriate, but in many cases the intent is for the measure to only consider patient data with a single specific code. The term used for such a single code in an eCQM without a value set is directly referenced code. Current uses of directly referenced codes include birth date and death date since these two concepts are used in many measures. These codes are embedded in the MAT structure so that measure developers do not need to create a direct referenced code or value set for these concepts. Beginning with implementation of QDM 5.3 and CQL, direct referenced codes will be included directly with the measure logic and included in a new terminology section of the HQMF.

In many cases, eCQMs will use directly referenced codes if there is only one concept that meets the definition of the eCQM data elements. However, some value sets that contain only one code will continue to exist for the following circumstances:

- There is only one code available at the time the value set is created; however, there is a reasonable expectation that additional codes will be created to represent the intent of the value set. Thus, rather than creating a direct referenced code for a quality data element in a measure and later changing it to a value set, the author may create a single code value set. For example, there is only one test for antibodies for a specific virus currently available, but additional tests are in development and will be equally appropriate to meet the intent of the measure clause. Because the tests are not yet available, they have not been submitted to LOINC for coding. The measure author may create a value set containing the existing code in this instance.

- A value set initially contained multiple codes, however all except one was retired by the code system(s). Since the measure may need to allow look-back, the value set remains valid with only one active code.

### 2.7.1 Value Sets and Directly Referenced Codes that Define QDM Categories

It is important to note that value sets define a QDM element’s category, not a QDM element’s datatype. Here is an example of a very common QDM element:

```
Procedure, Performed: "value set A"
```

In this example, the value set defines which procedure the criterion is looking for. The codes in this value set should only indicate the procedure, not whether the procedure was performed, ordered, or recommended, since that is represented using different datatypes. The following example shows a QDM element with an attribute:

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9 The Value Set Authority Center provides downloadable access to all official versions of the vocabulary value sets contained in published measures. For more information, visit [https://vsac.nlm.nih.gov/](https://vsac.nlm.nih.gov/). Last accessed June 2018.
Laboratory Test, Performed: "value set A" (result: "value set B")

In this example, value set A defines the category of the QDM element. Since the category is a Laboratory Test, value set A answers the question of which laboratory test. Value set B, on the other hand, defines the attribute result. Value set B should contain codes for different coded result values.

Similarly, direct referenced codes define a QDM element’s category, not a QDM element’s datatype. Here is an example of a very common QDM element using a direct referenced code:

Assessment, Performed: "Asthma Control Test using LOINC version 2.59 Code (82674.3)"

In this example, the direct referenced code defines which assessment the criterion is looking for. The codes provided should only indicate the specific assessment, and not its context (i.e., whether the procedure was performed, ordered, or recommended), since the context is represented using different datatypes. The following example shows a QDM element with an attribute:

Assessment, Performed: "Asthma Control Test using LOINC version 2.59 Code (82674.3)"

Components:
- In the past 4 weeks, how much of the time did your asthma keep you from getting as much done at work, school or at home? Using LOINC version 2.59 Code (82669-3)
- During the past 4 weeks, how often have you had shortness of breath? Using LOINC version 2.59 Code (82670-1)
- During the past 4 weeks, how often did your asthma symptoms (wheezing, coughing, shortness of breath, chest tightness or pain) wake you up at night or earlier than usual in the morning? Using LOINC version 2.59 Code (82671-9)
- During the past 4 weeks, how often have you used your rescue inhaler or nebulizer medication (such as albuterol)? Using LOINC version 2.59 Code (82672-7)
- How would you rate your asthma control during the past 4 weeks? Using LOINC version 2.59 Code (82673-5)

Total score ACT Using LOINC version 2.59 Code (82668-5)

In this example, the directly referenced code Asthma Control Test defines the category of the QDM Assessment element. Since the category is an Assessment, directly referenced code Asthma Control Test answers the question of which assessment. Six additional directly referenced codes define the attributes (in this case the individual assessment questions in the test) associated with the assessment. More information about components can be found in Section 4.

2.7.2 Value Sets that Define QDM Attributes

Some QDM attributes can be defined by value sets or direct referenced codes, similar to how QDM Categories are defined by value sets.
2.7.3 Value Set Groupings

Direct referenced codes indicate a specific code and, therefore, addressed only one code system. Each value set contains codes from one code system. However, multiple value sets can be combined into one value set called a value set grouping. A parent value set may also contain child (or nested) value sets that define the same category. The approach is consistent with the Health Level 7 (HL7) definition for a value set as “a uniquely identifiable set of valid concept representations, where any concept representation can be tested to determine whether or not it is a member of the value set … A sub-value set is a sub-set of a ‘parent’ value set … When a value set entry references another value set, the child value set is referred to as a nested value set. There is no pre-set limit to the level of nesting allowed within value sets. Value sets cannot contain themselves or any of their ancestors (i.e., they cannot be defined recursively).” With respect to value sets, a value is a specific code defined by a given taxonomy. Values are included in value sets. In the context of QDM elements, some categories (e.g., laboratory test) have an attribute of “result.” A result may be expressed as a value (numeric or alphanumeric).

---

3. Timing Considerations for QDM Data Elements

Prior versions of QDM included timing attributes of QDM data elements, basically start and stop times or author times. QDM uses author dateTime to indicate the time a data element is documented, i.e., a single point in time. To assure accurate retrieval of time-related information from clinical software, QDM 5.4 includes more explicit definition of the intent of start and stop for each datatype. For instances about events that can occur over a period of time, QDM 5.3 describes intervals between the startTime and the stopTime defined for each datatype. QDM 5.4 uses the term period to indicate these intervals to be consistent with current activities in HL7 Fast Healthcare Interoperability Resources (FHIR). The major reason for identifying periods is to enable simpler logic expression in CQL, i.e., refer to start time as the beginning of the period and end time as the end of the period. Further, to be consistent, QDM 5.4 limits the number of types of periods while assuring they can be somewhat explanatory.

3.0 Types of periods

3.0.1 Relevant period

Relevant period is the default, or general, method to describe a start to stop time for the following 14 datatypes:

- Adverse Event
- Care Goal
- Communication, Performed
- Device, Applied
- Diagnostic Study, Performed
- Encounter, Performed
- Intervention, Performed
- Laboratory Test, Performed
- Medication, Active
- Medication, Administered
- Patient Characteristic, Payer
- Physical Exam, Performed
- Procedure, Performed
- Substance, Administered

3.0.2 Prevalence period

Prevalence period is specific to indicate those datatypes that indicate the difference between onset and abatement dateTimes for three datatypes:

- Diagnosis
- Allergy
- Symptom

---

11 Health Level 7 (HL7) Fast Health Interoperability Resources (FHIR) information is available at https://www.hl7.org/fhir/overview.html.
3.0.3 Participation period

*Participation period* is specific to the time an individual participates in a health plan, health management program or a clinical treatment program for one datatype:

- Participation

3.0.4 Location periods

*Location periods* are specific to indicate the difference between the arrival and departure times with respect to the locations where the patient is treated (as an attribute of Encounter, Performed). QDM includes location periods for one datatype:

- Encounter, Performed

Note that any given Encounter, Performed can include zero to many facility locations (i.e., the facility locations has a cardinality of 0..*). Each facility locations attribute listed can include a single location period (i.e., the physical *arrival* and physical *departure* times for that facility location).

3.1 Use of Timing Periods Vs Author Times

Note that some datatypes list periods and author dateTimes as attributes. The reason is to allow a measure developer to differentiate in their logic the time an action occurs versus the time it is documented. In most cases, when the start dateTime of a period is not known, the retrieve should default to the author dateTime without having to specify author dateTime in the measure. Section 4.1 lists the attribute definitions.
4. QDM Definitions

4.0 Categories and Datatypes

4.0.1 Adverse Event

*Adverse Event is used to define* any untoward medical occurrence associated with the clinical care delivery, whether or not considered drug related.

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse Event</td>
<td>Data elements that meet criteria using this datatype should document the Adverse Event and its corresponding value set. The Relevant Period references: <code>startTime</code> – the time the adverse event began; <code>stopTime</code> – the time the adverse event completed.</td>
<td>• Relevant Period&lt;br&gt;code&lt;br&gt;Type&lt;br&gt;Severity&lt;br&gt;FacilityLocations&lt;br&gt;Author dateTime&lt;br&gt;id</td>
</tr>
</tbody>
</table>

4.0.2 Allergy/Intolerance

*Allergy is used to* address immune-mediated reactions to a substance such as type 1 hypersensitivity reactions, other allergy-like reactions, including pseudo-allergy.

*Intolerance* is a record of a clinical assessment of a propensity, or a potential risk to an individual, to have a non-immune mediated adverse reaction on future exposure to the specified substance, or class of substance.

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allergy/Intolerance</td>
<td>Data elements that meet criteria using this datatype should document the Allergy or intolerance and its corresponding value set. Timing: The Prevalence Period references the time from the <em>onset date</em> to the <em>abatement date</em>.</td>
<td>• Prevalence Period&lt;br&gt;Code (for Substance)&lt;br&gt;Type&lt;br&gt;Severity&lt;br&gt;Author Date/Time&lt;br&gt;id</td>
</tr>
</tbody>
</table>

4.0.3 Assessment

*Assessment* is a resource used to define specific observations that clinicians use to guide treatment of the patient. An assessment can be a single question, or observable entity with an expected response, an organized collection of questions intended to solicit information from patients, providers or other individuals, or a single observable entity that is part of such a collection of questions.
### Table 3. Assessment Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assessment, Performed</td>
<td>Data elements that meet criteria using this datatype should document completion of the assessment indicated by the QDM category and its corresponding value set. Timing: The time the assessment is completed; Author dateTime.</td>
<td>• Author dateTime • Negation Rationale • Reason • Method • Result (allow dateTime as a result response type; add percentage as a result response type) • relatedTo • Code • id • Components (may appear 0 or many times) (Each component will have: – code – result)</td>
</tr>
<tr>
<td>Assessment, Order</td>
<td>Data elements that meet these criteria using this datatype should document an order by a clinician or appropriately licensed care provider to a patient or an appropriate provider or organization to perform an assessment indicated by the QDM category and its corresponding value set. Timing: The time the order is authored (i.e., provided to the patient). NOTE: Orders address the time that the order is authored, a single point in time. Some assessment orders will be addressed as components of a care plan which incorporates the period (or timing) when the order is to be carried out. Measure developers should address feasibility of clinical workflow to capture Assessment, Order when evaluating measures.</td>
<td>• Author dateTime • Negation Rationale • Reason • Code • id</td>
</tr>
<tr>
<td>Assessment, Recommended</td>
<td>Data elements that meet these criteria using this datatype should document a recommendation for a request by a clinician or appropriately licensed care provider to a patient or an appropriate provider or organization to perform an assessment indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures.</td>
<td>• Author dateTime • Negation Rationale • Reason • Code • id</td>
</tr>
</tbody>
</table>
4.0.4 Care Experience

*Care Experience* represents the experience a patient has when receiving care or a provider has when providing care. The individual *Care Experience* datatypes should be consulted for further details.

**Table 4. Care Experience Datatypes and Attributes**

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Care Experience</td>
<td>Data elements that meet this criterion indicate the patient's care experience, usually measured with a validated survey tool. The most common tool is the Consumer Assessment of Healthcare Providers and Systems. Timing: The time the care experience is recorded; Author dateTime.</td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
<tr>
<td>Provider Care Experience</td>
<td>Data elements that meet this criterion indicate the provider's experience with availability of resources (e.g., scheduling, equipment, space, and such consumables as medications). Provider care experience gauges provider satisfaction with key structures, processes, and outcomes in the healthcare delivery system. Timing: The time the care experience is recorded; Author dateTime.</td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
</tbody>
</table>

4.0.5 Care Goal

*Care Goal* represents a defined target or measure to be achieved in the process of patient care, that is, an expected outcome. A typical goal is expressed as a change in status expected at a defined future time. That change can be an observation represented by other QDM categories (diagnostic tests, laboratory tests, symptoms, etc.) scheduled for some time in the future and with a particular value. A goal can be found in the plan of care (care plan), the structure used by all stakeholders, including the patient, to define the management actions for the various conditions, problems, or issues identified for the target of the plan. This structure, through which the goals and care-planning actions and processes can be organized, planned, communicated, and checked for completion, is represented in the QDM categories as a Record Artifact. A time/date stamp is required. Specifically, a care plan is composed of the following elements:

- **Problem**, which is managed by other QDM standard categories (condition/diagnosis/problem) and their related data elements.
- **Procedure**, which is managed by other standard categories and their related data elements. Note that procedures are a continuum of interventions ranging from actions patients can do for themselves to those that can be performed by others (caregivers or clinical professionals), including detailed complex surgical procedures requiring highly trained physicians, nurses, and state-of-the-art facilities.
- **Goal**, which is what is expected to happen.
- **Outcome**, which is what happened. An outcome can be shown by other QDM standard categories and their related data elements.
Table 5. Care Goal Datatype and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Care Goal</td>
<td>Unlike other QDM datatypes, the Care Goal datatype does not indicate a specific context of use. Instead, to meet this criterion, there must be documentation of a care goal as defined by the Care Goal QDM category and its corresponding value set. Timing: The Relevant Period references the period between: • startTime – when the goal is recorded, and therefore should be considered effective, • stopTime – when the target outcome is expected to be met</td>
<td>• relatedTo (the problem, procedure (or other class of information) to which the care goal is related) • Relevant Period • Target Outcome (add percentage as a target outcome response type) • Code • id</td>
</tr>
</tbody>
</table>

4.0.6 Communication

*Communication* represents the transmission, receipt, or acknowledgement of information sent from a source to a recipient, such as from one clinician to another regarding findings, assessments, plans of care, consultative advice, instructions, educational resources, etc.

The HL7 FHIR 3.0 resource *Communication* defines communication as a conveyance of information from one entity, a sender, to another entity, a receiver. The senders and receivers may be patients, practitioners, related persons, organizations, or devices. Established FHIR resources include:

- A reminder or alert delivered to a responsible provider
- A recorded notification from the nurse that a patient's temperature exceeds a value
- A notification to a public health agency of a patient presenting with a communicable disease reportable to the public health agency
- Patient educational material sent by a provider to a patient
- Non-patient specific communication use cases may include:
  - A nurse call from a hall bathroom
  - Advisory for battery service from a pump

Generally, direct patient counseling, training or education represents an *intervention* (section 4.0.14), or *procedure* (section 4.0.19). A time and date stamp is required.

QDM through version 5.3 included 3 communication QDM datatypes:

- Communication, Provider to Patient
- Communication, Patient to Provider
- Communication, Provider to Provider
These three datatypes restricted the measure developer to communication between patients and providers and among providers. The structure of the datatypes also provided implicit guidance to implementers to define what type of provider based on the intent of the measure.

QDM 5.4 merges the three communication QDM datatypes into one, Communication, Performed to allow greater expressivity and more explicit definition of the sender and receiver, and to indicate the anticipated communication medium. This new QDM modeling is more consistently with other QDM datatypes, and it is also modeled more consistently with existing communication interoperability resources (e.g., HL7 FHIR).

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Communication: Performed</td>
<td>To meet criteria using this datatype, the communication indicated by the Communication QDM category is a conveyance of information from one entity (e.g., person, organization, or device) to another. Timing: Relevant Period: The time the communication is sent (start time) to the time the communication is received (end time) Author date/time: the time the communication is documented (specifically significant for negation rationale, i.e., Communication Not Performed)</td>
<td>• Category • Sender • Recipient • Medium • Negation Rationale • Relevant Period • Author date/time • relatedTo • Code • id</td>
</tr>
</tbody>
</table>

### 4.0.7 Condition/Diagnosis/Problem

*Condition/Diagnosis/Problem* represents a practitioner’s identification of a patient’s disease, illness, injury, or condition. This category contains a single datatype to represent all of these concepts: *Diagnosis*. A practitioner determines the *diagnosis* by means of examination, diagnostic test results, patient history, and/or family history. *Diagnoses* are usually considered unfavorable, but may also represent neutral or favorable conditions that affect a patient’s plan of care (e.g., pregnancy).

The QDM does not prescribe the source of *diagnosis* data in the EHR. *Diagnoses* may be found in a patient’s problem list, encounter diagnosis list, claims data, or other sources within the EHR. The preferred terminology for diagnoses is SNOMED-CT, but diagnoses may also be encoded using ICD-9/10.

The *Diagnosis* datatype should not be used for *differential diagnoses* or *rule-out diagnoses* (neither of which are currently supported by the QDM).
### Table 7. Condition/Diagnosis/Problem Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Diagnosis      | Data elements that meet criteria using this datatype should document the Condition/Diagnosis/Problem and its corresponding value set. The onset dateTime corresponds to the implicit start dateTime of the datatype and the abatement dateTime corresponds to the implicit stop dateTime of the datatype. If the abatement dateTime is not present, then the diagnosis is considered to still be active. When this datatype is used with timing relationships, the criterion is looking for an active diagnosis for the time frame indicated by the timing relationships. Timing: The Prevalence Period references the time from the onset dateTime to the abatement dateTime. | • Prevalence Period  
• Anatomical Location Site  
• Severity  
• Code  
• Author dateTime  
• id |

### 4.0.8 Device

*Device* represents an instrument, apparatus, implement, machine, contrivance, implant, in-vitro reagent, or other similar or related article, including a component part or accessory, intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease and not dependent on being metabolized to achieve any of its primary intended purposes.12

Documented evidence of a device may exist in a clinical record in various ways.

- A provider may document placement of a device as an intervention or procedure (e.g., the QDM datatypes Intervention, Performed, or Procedure, Performed). Example: Procedure, Performed: Pacemaker insertion.
- The provider may document presence of the device as a finding, perhaps including the finding on the problem list. Example: Diagnosis: Pacemaker present, or Assessment, Performed: Pacemaker present.
- A provider may also document the device in a specific device use statement or assessment. Example: Device, Applied: Transtelephonic monitoring of pacemaker assessment. Note that the current use of the QDM datatype “Device, Applied” usually references the procedure to “apply” the device (i.e., to use for the patient, to use on the patient’s body, or to implant in the patient’s body). Each of these current uses should address the concept using the QDM datatypes, Intervention, Performed, or Procedure, Performed.

Given the variation in determining evidence of device usage, a measure developer may need to include multiple queries (or retrieves) to assure capture of all devices present in the measure population. The QDM datatype, Device, Applied, provides the best method to determine specific details about the device in use if a device usage assessment exists to provide such information. It also allows a measure developer to explicitly express which devices should be included in the query.

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### Table 8. Device Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| **Device, Applied**       | Data elements that meet criteria using this datatype should document that the device indicated by the QDM category and its corresponding value set is in use, or impacts or alters the treatment, care plan, or encounter (e.g., an antithrombotic device has been placed on the patient's legs to prevent thromboembolism, or a cardiac pacemaker is in place). Timing: The Relevant Period addresses: startTime – When the device is inserted or first used stopTime – when the device is removed or last used                                                                 |  • Anatomical Location Site  
  • Negation Rationale  
  • Reason  
  • Relevant Period  
  • Author dateTime  
  • Code  
  • id |
| **Device, Order**         | Data elements that meet criteria using this datatype should document an order for the device indicated by the QDM category and its corresponding value set.                                                                                                 |  • Negation Rationale  
  • Reason  
  • Author dateTime  
  • Code  
  • id |
| **Device, Recommended**   | Data elements that meet criteria using this datatype should document a recommendation to use the device indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures. |  • Negation Rationale  
  • Reason  
  • Author dateTime  
  • Code  
  • id |

### 4.0.9 Diagnostic Study

*Diagnostic Study* represents any kind of medical test performed as a specific test or series of steps to aid in diagnosing or detecting disease (e.g., to establish a diagnosis, measure the progress or recovery from disease, confirm that a person is free from disease). The QDM defines diagnostic studies as those that are not performed in organizations that perform testing on samples of human blood, tissue, or other substance from the body. Diagnostic studies may make use of digital images and textual reports. Such studies include but are not limited to imaging studies, cardiology studies (electrocardiogram, treadmill stress testing), pulmonary-function testing, vascular laboratory testing, and others.

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### Table 9. Diagnostic Study Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diagnostic Study, Order</td>
<td>Data elements that meet criteria using this datatype should document a request by a clinician or appropriately licensed care provider to an appropriate provider or organization to perform the diagnostic study indicated by the QDM category and its corresponding value set. The request may be in the form of a consultation or a direct order to the organization that performs the diagnostic study. Diagnostic studies are those that are not performed in the clinical laboratory. Such studies include but are not limited to imaging studies, cardiology studies (electrocardiogram, treadmill stress testing), pulmonary function testing, vascular laboratory testing, and others. Timing: The time the order is signed; Author dateTime.</td>
<td>• Negation Rationale&lt;br&gt;• Reason&lt;br&gt;• Author dateTime&lt;br&gt;• Code&lt;br&gt;• id</td>
</tr>
<tr>
<td>Diagnostic Study, Performed</td>
<td>Data elements that meet criteria using this datatype should document the completion of the diagnostic study indicated by the QDM category and its corresponding value set. Timing: The Relevant Period addresses: startTime – when the diagnostic study is initiated stopTime – when the diagnostic study is completed Examples: • Initiation of a treadmill stress test to the time the treadmill stress test has completed • Initiation of the ultrasound study until completion of the ultrasound study</td>
<td>• Facility Location&lt;br&gt;• Method&lt;br&gt;• Negation Rationale&lt;br&gt;• Reason&lt;br&gt;• Result&lt;br&gt;• Result dateTime&lt;br&gt;• Relevant Period&lt;br&gt;• Status&lt;br&gt;• Code&lt;br&gt;• Author dateTime&lt;br&gt;• id&lt;br&gt;• Components (may appear 0 or many times) (Each component will have: – code&lt;br&gt;– result)</td>
</tr>
<tr>
<td>Diagnostic Study, Recommended</td>
<td>Data elements that meet criteria using this datatype should document a recommendation for a request by a clinician or appropriately licensed care provider to an appropriate provider or organization to perform the diagnostic study indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures.</td>
<td>• Negation Rationale&lt;br&gt;• Author dateTime&lt;br&gt;• Code&lt;br&gt;• id</td>
</tr>
</tbody>
</table>
4.0.10 Encounter

*Encounter* represents an identifiable grouping of healthcare-related activities characterized by the entity relationship between the subject of care and a healthcare provider; such a grouping is determined by the healthcare provider. A *patient encounter* represents interaction between a healthcare provider and a patient with a face-to-face patient visit to a clinician’s office, or any electronically remote interaction with a clinician for any form of diagnostic treatment or therapeutic event. Encounters can be billable events but are not limited to billable interactions. Each encounter has an associated location or modality within which it occurred (such as an office, home, electronic methods, phone encounter, or telemedicine methods). The *encounter location* is the patient’s location at the time of measurement. Different levels of interaction can be specified in the value associated with the element while modes of interaction (e.g., telephone) may be modeled using the data flow attribute.

**Table 10. Encounter Datatypes and Attributes**

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Encounter, Order           | Data elements that meet criteria using this datatype should document that an order for the encounter indicated by the QDM category and its corresponding value set has been recommended. Timing: The time the order is signed; Author dateTime. | • Facility Location  
  • Negation Rationale  
  • Reason  
  • Author dateTime  
  • Code  
  • id |
| Encounter, Performed       | Data elements that meet criteria using this datatype should document that the encounter indicated by the QDM category and its corresponding value set is in progress or has been completed. Timing: The Relevant Period addresses:  
  startTime – The time the encounter began (admission time)  
  stopTime – The time the encounter ended (discharge time)  
  The Author dateTime addresses when an Encounter is documented. Documentation can occur at the beginning, during, at the end or subsequent to the end of an Encounter. The Author dateTime should be used only if the Relevant Period cannot be obtained.  
  The Location Period is an attribute of the attribute facility location addresses:  
  startTIme = the time the patient arrived at the location;  
  stopTime = the time the patient departed from the location | • Relevant Period  
  • Admission Source  
  • Diagnoses  
  • Discharge Disposition  
  • Length of Stay  
  • Negation Rationale  
  • Principal Diagnosis  
  • Author dateTime  
  • Code  
  • id  
  • Facility Locations (may appear 0 or many times) Each component will have:  
  – Code  
  – location period |

---

### Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Encounter, Recommended</strong></td>
<td>Data elements that meet criteria using this datatype should document that the encounter indicated by the QDM category and its corresponding value set has been recommended. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures.</td>
<td>• Facility Location • Negation Rationale • Reason • Author dateTime • Code • id</td>
</tr>
</tbody>
</table>

### 4.0.11 Family History

*Family History* represents a diagnosis or problem experienced by a family member of the patient. Typically, a family history will not contain very much detail, but the simple identification of a diagnosis or problem in the patient’s family history may be relevant to the care of the patient.

If a relationship is specified, codes from the HL7 Personal Relationship Role Type value set (2.16.840.1.113883.1.11.19563) should be used to ensure compatibility with QRDA reporting constraints.

#### Table 11. Family History Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Family History</strong></td>
<td>To meet criteria using this datatype, the diagnosis/problem indicated by the FamilyHistory QDM category and its corresponding value set should reflect a diagnosis/problem of a family member. Timing: The time the family history item is authored (i.e., entered into the record). NOTE – Measure developers suggested that onset age for family history represents one item in a risk assessment for individual patients. Thus, onset age (when the family member developed the condition indicated in the Family History) can be determined using the Assessment, Performed QDM datatype.</td>
<td>• Author dateTime • Relationships • Code • id</td>
</tr>
</tbody>
</table>

### 4.0.12 Immunization

*Immunization* represents vaccines administered to patients in healthcare settings but does not include non-vaccine agents.
Table 12. Immunization Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immunization, Administered</td>
<td>Data elements that meet criteria using this datatype should document that the vaccine indicated by the QDM category and its corresponding value set was actually administered to the patient. Timing: The time the immunization is administered, i.e., a single point in time, or Author dateTime.</td>
<td>• Dosage</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Negation Rationale</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Reason</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Route</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
<tr>
<td>Immunization, Order</td>
<td>Data elements that meet criteria using this datatype should document a request for the immunization indicated by the QDM category and its corresponding value set. Timing: The time the order is signed; Author dateTime.</td>
<td>• Active dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Dosage</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Negation Rationale</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Reason</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Route</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Supply</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
</tbody>
</table>

4.0.13 Individual Characteristic

Individual Characteristic represents specific factors about a patient, clinician, provider, or facility. Included are demographics, behavioral factors, social or cultural factors, available resources, and preferences. Behaviors reference responses or actions that affect (either positively or negatively) health or healthcare. Included in this category are mental health issues, adherence issues unrelated to other factors or resources, coping ability, grief issues, and substance use/abuse. Social/cultural factors are characteristics of an individual related to family/caregiver support, education, and literacy (including health literacy), primary language, cultural beliefs (including health beliefs), persistent life stressors, spiritual and religious beliefs, immigration status, and history of abuse or neglect. Resources are means available to a patient to meet health and healthcare needs, which might include caregiver support, insurance coverage, financial resources, and community resources to which the patient is already connected and from which the patient is receiving benefit. Preferences are choices made by patients and their caregivers relative to options for care or treatment (including scheduling, care experience, and meeting of personal health goals) and the sharing and disclosure of their health information.

Table 13. Individual Characteristic Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient Characteristic</td>
<td>Data elements that meet criteria using this datatype should document a characteristic of the patient not represented by one of the more specific Individual Characteristic datatypes. Timing: The time the characteristic is authored.</td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
<tr>
<td>Datatype</td>
<td>Definition</td>
<td>Attributes</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>Patient Characteristic Birthdate</td>
<td>The <em>Patient Characteristic Birthdate</em> data element should document the patient’s date of birth.</td>
<td>• Birth dateTime</td>
</tr>
<tr>
<td></td>
<td>Timing: The Patient Characteristic, Birthdate is a single point in time representing the date and time of birth. It does not have a start and stop time.</td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td>Note: <em>Patient Characteristic Birthdate</em> is fixed to LOINC code 21112-8 (Birth date) and therefore cannot be further qualified with a value set.</td>
<td>• id</td>
</tr>
<tr>
<td>Patient Characteristic Clinical Trial Participant</td>
<td>Data elements that meet criteria using this datatype should document that the patient is a clinical trial participant for the clinical trial indicated by the QDM category and its corresponding value set.</td>
<td>• Reason</td>
</tr>
<tr>
<td></td>
<td>Timing: The Relevant Period addresses: start Time – The time the clinical trial began stop Time – The time the clinical trial ended</td>
<td>• Relevant Period</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
<tr>
<td>Patient Characteristic Ethnicity</td>
<td>Data elements that meet criteria using this datatype should document that the patient has one or more of the ethnicities indicated by the QDM category and its corresponding value set.</td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td>Timing: Ethnicity does not have a specific timing. Measures using Patient Characteristic, Ethnicity should address the most recent entry in the clinical record.</td>
<td>• id</td>
</tr>
<tr>
<td>Patient Characteristic Expired</td>
<td>The <em>Patient Characteristic Expired</em> data element should document that the patient is deceased.</td>
<td>• Cause</td>
</tr>
<tr>
<td></td>
<td>Timing: The Patient Characteristic, Expired is a single point in time representing the date and time of death. It does not have a start and stop time.</td>
<td>• Expiration dateTime</td>
</tr>
<tr>
<td></td>
<td>Note: <em>Patient Characteristic Expired</em> is fixed to SNOMED-CT® code 419099009 (Dead) and therefore cannot be further qualified with a value set.</td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
<tr>
<td>Patient Characteristic Payer</td>
<td>Data elements that meet criteria using this datatype should document that the patient has one or more of the payers indicated by the QDM category and its corresponding value set.</td>
<td>• Relevant Period</td>
</tr>
<tr>
<td></td>
<td>Timing: The Relevant Period addresses: start Time – The first day of insurance coverage with the referenced payer stop Time – The last day of insurance coverage with the referenced payer</td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
</tbody>
</table>
### Datatype Definitions

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| **Patient Characteristic Race** | Data elements that meet criteria using this datatype should document the patient’s race. Timing: Race does not have a specific timing. Measures using Patient Characteristic, Race should address the most recent entry in the clinical record. | • Code  
• id |
| **Patient Characteristic Sex** | Data elements that meet criteria using this datatype should document that the patient's sex matches the QDM category and its corresponding value set. Timing: Birth (administrative) sex does not have a specific timing. | • Code  
• id |
| **Provider Characteristic**   | Data elements that meet criteria using this datatype should document a characteristic of the provider. Timing: The time the characteristic is authored. | • Author dateTime  
• Code  
• id |

#### 4.0.14 Intervention

*Intervention* represents a course of action intended to achieve a result in the care of persons with health *problems* that does not involve direct physical contact with a patient. Examples include patient education and therapeutic communication.

The QDM User Group carefully considered merging the Intervention and Procedure QDM categories as each is modeled identically and interoperability standards does not differentiate between interventions and procedures. However, the measure developers prefer to retain the intervention and procedure categories as they are more clinically expressive for clinicians to understand the measure intent.

**Table 14. Intervention Datatypes and Attributes**

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Intervention, Order | Data elements that meet criteria using this datatype should document a request to perform the intervention indicated by the QDM category and its corresponding value set. Timing: The time the order is signed; Author dateTime | • Negation Rationale  
• Reason  
• Author dateTime  
• Code  
• id |
### Interventions, Performed

Data elements that meet criteria using this datatype should document the completion of the intervention indicated by the QDM category and its corresponding value set. Timing: The Relevant Period addresses:
- **startTime** – The time the intervention begins
- **stopTime** – The time the intervention ends

**NOTE** - timing refers to a single instance of an intervention. If a measure seeks to evaluate multiple interventions over a period of time, the measure developer should use CQL logic to represent the query request.

<table>
<thead>
<tr>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Negation Rationale</td>
</tr>
<tr>
<td>• Reason</td>
</tr>
<tr>
<td>• Result</td>
</tr>
<tr>
<td>• Relevant Period</td>
</tr>
<tr>
<td>• Author dateTime</td>
</tr>
<tr>
<td>• Status</td>
</tr>
<tr>
<td>• Code</td>
</tr>
<tr>
<td>• id</td>
</tr>
</tbody>
</table>

### Interventions, Recommended

Data elements that meet criteria using this datatype should document a recommendation for the intervention indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient).

**NOTE**: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures.

<table>
<thead>
<tr>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Negation Rationale</td>
</tr>
<tr>
<td>• Reason</td>
</tr>
<tr>
<td>• Author dateTime</td>
</tr>
<tr>
<td>• Code</td>
</tr>
<tr>
<td>• id</td>
</tr>
</tbody>
</table>

### Laboratory Test

*Laboratory Test* represents a medical procedure that involves testing a sample of blood, urine, or other substance from the body. Tests can help determine a diagnosis, plan treatment, check to see if treatment is working, or monitor the disease over time.\(^\text{15}\) This QDM data category for Laboratory Test is only used for information about the subject of record.

### Table 15. Laboratory Test Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory Test, Order</td>
<td>Data elements that meet criteria using this datatype should document a request for the laboratory test indicated by the QDM category and its corresponding value set. Timing: The time the order is signed; Author dateTime.</td>
<td>• Negation Rationale</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Reason</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Author dateTime</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Code</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• id</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Laboratory Test, Performed                   | Data elements that meet criteria using this datatype should document the laboratory test indicated by the QDM category and its corresponding value set was performed. Timing: The Relevant Period addresses: startTime – When the laboratory test is initiated (i.e., the time the specimen collection begins) stopTime – when the laboratory test is completed (i.e., the time the specimen collection ends) Examples:  
  • Initiation of a venipuncture for a fasting blood glucose to the time venipuncture for the fasting blood glucose is completed – basically a single point in time for many specimen collections  
  • Initiation of a 24-hour urine collection for measured creatinine clearance until completion of the 24-hour urine collection Note – the time that the result report is available is a separate attribute than the time of the study (specimen collection) | • Method  
• Negation Rationale  
• Reason  
• Reference Range High  
• Reference Range Low  
• Result  
• Result dateTime  
• Relevant Period  
• Status  
• Author dateTime  
• Code  
• id  
• Components (may appear 0 or many times) (Each component will have:  
  – code  
  – result  
  – reference range High (optional)  
  – reference range Low (optional)) |
| Laboratory Test, Recommended                | Data elements that meet criteria using this datatype should document a recommendation for the laboratory test indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures. | • Negation Rationale  
• Reason  
• Author dateTime  
• Code  
• id |

### 4.0.16 Medication

*Medication* represents clinical drugs or chemical substances intended for use in the medical diagnosis, cure, treatment, or prevention of disease. Medications are defined as direct referenced values or value sets containing values derived from code systems such as RxNorm.16

---

Table 16. Medication Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Medication, Active| Data elements that meet criteria using this datatype should document that the medication indicated by the QDM category and its corresponding value set is being taken by the patient. Keep in mind that when this datatype is used with timing relationships, the criterion is looking for a medication being taken for the time frame indicated by the timing relationships. Timing: The Relevant Period addresses: \(\text{StartTime} =\) when the medication is first known to be used (generally the time of entry on the medication list); \(\text{StopTime} =\) when the medication is discontinued (generally, the time discontinuation is recorded on the medication list) | • Dosage  
• Frequency  
• Route  
• Relevant Period  
• Code  
• id |
| Medication, Administered | Data elements that meet criteria using this datatype should document that the medication indicated by the QDM category and its corresponding value set was actually administered to the patient. Timing: The Relevant Period addresses: \(\text{startTime} =\) when a single medication administration event starts (e.g., the initiation of an intravenous infusion, or administering a pill or IM injection to a patient); \(\text{stopTime} =\) when a single medication administration event ends (e.g., the end time of the intravenous infusion, or the administration of a pill or IM injection is completed - for pills and IM injections, the start and stop times are the same) NOTE – Measure developers should address multiple administrations over a period of time using CQL logic. Refer to Special Cases in Appendix A (Section A.3) for scenarios to consider in calculating cumulative medication duration. | • Dosage  
• Frequency  
• Negation Rationale  
• Reason  
• Route  
• Relevant Period  
• Author dateTime  
• Code  
• id |
| Medication, Discharge | Data elements that meet criteria using this datatype should document that the medications indicated by the QDM category and its corresponding value set should be taken by or given to the patient after being discharged from an inpatient encounter. Timing: The time the discharge medication list on the discharge instruction form is authored. | • Dosage  
• Frequency  
• Supply  
• Negation Rationale  
• Refills  
• Route  
• Author dateTime  
• Code  
• id |
<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Medication, Dispensed | Data elements that meet criteria using this datatype should document that a prescription for the medication indicated by the QDM category and its corresponding value set has been dispensed and provided to the patient or patient proxy. In the ambulatory setting, medications are primarily taken directly by patients and not directly observed. Hence, dispensed, or fulfillment, information is the closest health provider documentation of medication compliance. In settings where patients attest to taking medications in electronic format (perhaps a Personal Health Record), patient attestation of medication taken may be available. The QDM datatype, Medication, Administered addresses medication taken; to address the source of the information, a measure addressing such patient attestation would require use of the dataflow attribute, source. Timing: The time the medication dispensing event occurs; the Author dateTime. Timing: The Relevant Period addresses: startTime = when the first administration of the medication is expected. If not specified, the startTime defaults to the Author dateTime (the time the medication dispensing event occurs). stopTime = when the medication supply provided by the dispensing event is expected to be completed. Note that when calculating cumulative medication duration, the medication dispensed stopTime may be present directly in the fulfillment record. If the stopTime is not available, the duration in days is the difference between the Relevant Period start and stop times. The record may indicate which of the available refills the fulfillment represents but each dispensing event is unique. Therefore, the measure developer should use CQL logic to address multiple dispensing events over a period of time. Refer to Special Cases in Appendix A (Section A.3) for scenarios to consider in calculating cumulative medication duration. | • Dosage  
• Supply  
• Frequency  
• Negation Rationale  
• Refills  
• Route  
• Relevant Period  
• Author dateTime  
• Code  
• id                                              |
<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medication, Order</td>
<td>Data elements that meet criteria using this datatype should document a request to a pharmacy to provide the medication indicated by the QDM category and its corresponding value set.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Timing: The Author dateTime is the time the order is signed.</td>
<td>• Dosage</td>
</tr>
<tr>
<td></td>
<td>Timing: The Relevant Period addresses:</td>
<td>• Supply</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[startTime] = when the first administration of the medication is expected. The first administration may be expected at the time of the order or at a specified future date (i.e., the (active) time for the order); such information should be identified in the medication order. If the startTime is not specified in the order, the startTime defaults to the Author dateTime (the time the order is signed).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[stopTime] = when the medication supply provided by the medication order is expected to be completed, including all fulfillments covered by the number of refills.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Note that when calculating cumulative medication duration, the stopTime may be present directly in the medication order. If the stopTime is not available, the duration in days is the difference between the Relevant Period start and stop times multiplied by ((1 + \text{the number of refills})).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Refer to Special Cases in Appendix A (Section A.3) for scenarios to consider in calculating cumulative medication duration.</td>
<td></td>
</tr>
</tbody>
</table>

### 4.0.17 Participation

*Participation* represents a patient’s coverage by a program such as an insurance or medical plan or a payment agreement. Such programs can include patient-centered medical home, disease-specific programs, etc.\(^{17}\)

---

\(^{17}\) Definitions modeled similar to HL7 FHIR STU 3.0 - [https://www.hl7.org/fhir/coverage.html](https://www.hl7.org/fhir/coverage.html).
Table 17. Participation Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Participation  | Data elements that meet criteria using this datatype should document the type of plan or program in which the patient is expected to be enrolled. The program is identified as the Issuer (e.g., Aetna, BCBSA, Cigna, etc.). The code attribute indicates the coverage type indicating the program in which the subject of record participates (e.g., health insurance plan policy, disease specific policy, health maintenance organization policy, etc.). Timing: Participation Period addresses:  
• enrollmentStartdate – The time the patient enrolled in the program  
• enrollmentEnddate – The time the patient’s enrollment in the program ends | • Code  
• Participation Period  
• id |

4.0.18 Physical Exam

Physical Exam represents the evaluation of the patient's body and/or mental status exam to determine its state of health. The techniques of examination can include palpation (feeling with the hands or fingers), percussion (tapping with the fingers), auscultation (listening), visual inspection or observation, inquisition and smell. Measurements may include vital signs (blood pressure, pulse, respiration) as well as other clinical measures (such as expiratory flow rate and size of lesion). Physical exam includes psychiatric examinations.

Table 18. Physical Exam Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Physical Exam, Order | Data elements that meet criteria using this datatype should document a request for the physical exam indicated by the QDM category and its corresponding value set. The datatype is expected to be used to identify orders such as "vital signs, frequency every x hours," or "pedal pulse check, frequency every 15 minutes for x hours." Timing: The time the order is signed; Author dateTime. | • Anatomical Location Site  
• Negation Rationale  
• Reason  
• Author dateTime  
• Code  
• id |
<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Physical Exam, Performed         | Data elements that meet criteria using this datatype should document the completion of the physical exam indicated by the QDM category and its corresponding value set. Timing: The Relevant Period addresses: startTime – The time the physical examination activity begins stopTime – The time the physical examination activity ends NOTE - timing refers to a single instance of a physical examination activity. If a measure seeks to evaluate multiple physical examination activities over a period of time, the measure developer should use CQL logic to represent the query request. | • Anatomical Location Site  
• Method  
• Negation Rationale  
• Reason  
• Result  
• Relevant Period  
• Code  
• Author dateTime  
• id  
• Components (may appear 0 or many times) (Each component will have:  
– code  
– result) |
| Physical Exam, Recommended       | Data elements that meet criteria using this datatype should document a recommendation for the physical exam indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures. | • Anatomical Location Site  
• Negation Rationale  
• Reason  
• Author dateTime  
• Code  
• id |

### 4.0.19 Procedure

*Procedure* is derived directly from HL7 and Canada Health Infoway: “An Act whose immediate and primary outcome (post-condition) is the alteration of the physical condition of the subject. … Procedure is but one among several types of clinical activities such as observation, substance-administrations, and communicative interactions … Procedure does not comprise all acts of *sic* whose intent is intervention or treatment.”

A *procedure* may be a surgery or other type of physical manipulation of a person’s body in whole or in part for purposes of making observations and diagnoses or providing treatment.

---


Table 19. Procedure Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Procedure, Order          | Data elements that meet criteria using this datatype should document a request for the procedure indicated by the QDM category and its corresponding value set. Timing: The time the order is signed; Author dateTime. | • Anatomical Location Site  
• Negation Rationale  
• Ordinality  
• Reason  
• Author dateTime  
• Code  
• id |
| Procedure, Performed      | Data elements that meet criteria using this datatype should document the completion of the procedure indicated by the QDM category and its corresponding value set. Timing: The Relevant Period addresses:  
\[ \text{StartTime = the time the procedure begins;} \]  
\[ \text{StopTime = the time the procedure is completed} \]  
NOTE:  
1) Timing refers to a single instance of a procedure. If a measure seeks to evaluate multiple procedures over a period of time, the measure developer should use CQL logic to represent the query request.  
2) The Incision dateTime is a single point in time available from the Operating Room and/or Anesthesia Record. | • Anatomical Location Site  
• Incision dateTime  
• Method  
• Negation Rationale  
• Ordinality  
• Reason  
• Result  
• Relevant Period  
• Author dateTime  
• Status  
• Code  
• id  
• Components (may appear 0 or many times) (Each component will have:  
  o Code  
  o result) |
| Procedure, Recommended    | Data elements that meet criteria using this datatype should document the recommendation for the procedure indicated by the QDM category and its corresponding value set. Timing: The time the recommendation is authored (i.e., provided to the patient). NOTE: Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures. | • Anatomical Location Site  
• Negation Rationale  
• Ordinality  
• Reason  
• Author dateTime  
• Code  
• id |

4.0.20 Substance

Substance represents a homogeneous material with definite composition that includes allergens, biological materials, chemicals, foods, drugs and materials.\(^\text{\textsuperscript{20}}\) QDM distinguishes between medications from non-medication substances by separately listing medication datatypes.

Substance may or may not have a code or be classified by a code system such as RxNorm. Examples of a substance may include environmental agents (e.g., pollen, dust) and food (e.g., vitamins).

Measure developers have considered the use of the QDM substance category in light of alignment with existing interoperability data standards. Two use cases require further investigation and effort in the HL7 FHIR realm:

**Nutrition [Considerations for Future Use]**
The nutrition mappings from QDM to FHIR resources require continued effort as of the printing of this QDM 5.4 version. The [FHIR NutritionOrder resource](https://www.hl7.org/fhir/structure.html#nutritionorder) is a record of the request for the supply of a diet, oral supplement or enteral formulas for a patient, and thus allow specification of a complete diet order, but not an order for an individual nutrient. The NutritionOrder resource is currently at a low maturity level (i.e., it has had limited testing).

Therefore, for the current QDM 5.4 reference, substance is modeled similarly to medications.

**Blood Products [Considerations for Future Use]**
The HL7 FHIR version 3.0 resource for blood products allow specification of the blood product as a biologically derived product but specification for how such products are administered remains in development. Therefore, for the current QDM 5.4 reference, blood product administration can be managed as a Procedure, Performed or a Substance, Administered based on measure developer findings during measure implementation testing.

### Table 20. Substance Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Substance, Administered | Data elements that meet criteria using this datatype should document that the substance indicated by the QDM category and its corresponding value set was actually given to the patient. Timing: The Relevant Period addresses: **StartTime** = when a single substance administration event starts (e.g., the initiation of an intravenous infusion, or administering the substance orally or topically to a patient); **StopTime** = when a single substance administration event ends (e.g., the end time of the intravenous infusion, or the administration of a substance orally or topically is completed - for oral or topical administration, the start and stop times are the same) | • Dosage  
• Frequency  
• Negation Rationale  
• Route  
• Relevant Period  
• Author dateTime  
• Code  
• id |
| Substance, Order | Data elements that meet criteria using this datatype should document a request for the substance indicated by the QDM category and its corresponding value set. Timing: The time the order is signed; Author dateTime. | • Dosage  
• Frequency  
• Negation Rationale  
• Reason  
• Supply  
• Refills  
• Route  
• Author dateTime |
### Substance, Recommended

Data elements that meet criteria using this datatype should document a recommendation for the substance indicated by the QDM category and its corresponding value set.

**Timing:** The time the recommendation is authored (i.e., provided to the patient).

**NOTE:** Recommendations address the time that the recommendation occurs, a single point in time. Vendors have expressed concerns that recommendations are not necessarily captured or managed in a standard manner as part of structured data capture in clinical workflow; many are documented as part of assessments in narrative text. Measure developers should address feasibility of clinical workflow to capture recommendations when evaluating measures.

<table>
<thead>
<tr>
<th>Attributes</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Code</td>
</tr>
<tr>
<td>• id</td>
</tr>
<tr>
<td>• Dosage</td>
</tr>
<tr>
<td>• Frequency</td>
</tr>
<tr>
<td>• Negation Rationale</td>
</tr>
<tr>
<td>• Reason</td>
</tr>
<tr>
<td>• Refills</td>
</tr>
<tr>
<td>• Route</td>
</tr>
<tr>
<td>• Author dateTime</td>
</tr>
<tr>
<td>• Code</td>
</tr>
<tr>
<td>• id</td>
</tr>
</tbody>
</table>

### 4.0.21 Symptom

*Symptom* represents an indication that a person has a condition or disease. Some examples are headache, fever, fatigue, nausea, vomiting, and pain.\(^{21}\) Also, symptoms are subjective manifestations of the disease perceived by the patient.\(^{22}\) As an example to differentiate *symptom* from *finding*, the patient’s subjective symptom of fever is distinguished from the temperature (a finding). For a finding, there is either a source of either a temperature-measuring device together with a recorder of the device (electronically) or an individual (healthcare provider, patient, etc.).

The QDM User Group reviewed the QDM datatype, Symptom in that many symptoms are entered as findings as part of assessments and, therefore, may not be included in a specific EHR “symptom” section. However, for clarity of expression in the human readable eCQM, the User Group agreed to retain the QDM datatype, Symptom, without change.

#### Table 21. Symptom Datatypes and Attributes

<table>
<thead>
<tr>
<th>Datatype</th>
<th>Definition</th>
<th>Attributes</th>
</tr>
</thead>
</table>
| Symptom  | Data elements that meet criteria using this datatype should document the symptom and its corresponding value set. **Timing:** The Prevalence Period references the time from the *onset date* to the *abatement date*. The *onset dateTime* corresponds to the implicit start *dateTime* of the datatype and the abatement *dateTime* corresponds to the implicit stop *dateTime* of the datatype. If the abatement *dateTime* is null, then the symptom is considered to still be active. When this datatype is used with timing relationships, the | • Prevalence Period  
• Severity  
• Code  
• id |

---


4.1 Attributes

QDM 5.3 includes a new cardinality column in the attribute definition table. Cardinality refers to the number of instances of the attribute that can be included in the measure description or the performance report (e.g., only once, or multiple times). The purpose of this new information is to be more explicit in guiding specification of QDM data elements. In previous versions of QDM, the cardinality was assumed. Therefore, implementation guides provided the cardinality definitions (QDM-based HQMF for measure specifications and QRDA Category I for individual performance reporting). Note that cardinality for most attributes is 0.. 1 (i.e., can occur up to 1 time). The only attributes that have cardinality of 0.. * (i.e., can occur multiple times) are:

- **diagnoses** used with Encounter, Performed
- **components** used with Assessment, Performed; Diagnostic Study, Performed; Laboratory Test, Performed; Physical Exam, Performed; and Procedure, Performed
- **facility locations** used with Encounter, Performed
- **relatedTo** used with Care Goal; Communication, Performed; Assessment, Performed

Note that, where cardinality is greater than 1 (i.e., 0.. *), the attributes use the plural form rather than singular (except relatedTo which can be plural or singular). The reason is that use of the plural more clearly informs the implementer to search among all instances of the element to retrieve the appropriate content and avoid errors in the retrieval. For example, Encounter, Performed: Inpatient Hospitalization (diagnoses: atrial fibrillation) informs the implementer to search all relevant diagnosis addressed during the encounter (i.e., all on the diagnosis list) to find atrial fibrillation. The singular ‘diagnosis’ term could cause an error if the implementer interprets it to look for atrial fibrillation as the first diagnosis listed. Refer to Table 22 for definitions of each of the attributes.

### Table 22. Attribute Definitions

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active dateTime</td>
<td>When the order indicates the first immunization administration should occur. Active dateTime is most often used to specify immunizations for which administration is intended at a specific time in the future. If Active dateTime is not specified, it defaults to the Author (signed) dateTime.</td>
<td>Immunization, Order</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Admission Source</td>
<td>The location from which the patient was admitted (e.g., physician referral, facility from which the patient was transferred).</td>
<td>Encounter, Performed</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Anatomical Location Site</td>
<td>The anatomical site or structure</td>
<td>• Diagnosis (a)</td>
<td>0.. 1</td>
</tr>
<tr>
<td></td>
<td>• Where the diagnosis/problem manifests itself (a).</td>
<td>• Device, Applied (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• That is the focus of the action represented by the datatype (b).</td>
<td>• Physical Exam, Order (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Physical Exam, Performed (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Physical Exam, Recommended (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Procedure, Order (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Procedure, Performed (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Procedure, Recommended (b)</td>
<td></td>
</tr>
<tr>
<td>Author dateTime</td>
<td>The time the data element was entered into the clinical software.</td>
<td>• Assessment, Performed</td>
<td>0.. 1</td>
</tr>
<tr>
<td></td>
<td>Note, some datatypes include both Relevant Time and Author dateTime attributes.</td>
<td>• Care Goal</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Communication, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Device, Applied</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Device, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Device, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Diagnostic Study, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Diagnostic Study, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Encounter, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Encounter, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Encounter, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Immunization, Administered</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Immunization, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Administered</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Discharge</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Dispensed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Patient Care Experience</td>
<td></td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>-----------------</td>
<td>----------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Category</td>
<td>The type of message conveyed by a communication (e.g., notification, alert, reminder, instruction, etc.).</td>
<td>• Communication, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td>Cause</td>
<td>The recorded cause of death.</td>
<td>• Patient Characteristic Expired</td>
<td>0..1</td>
</tr>
<tr>
<td>Code</td>
<td>The single code or a member of the value set used to represent the quality data element. The code attribute explicitly specifies the value set (or direct referenced code) filter such that the query will retrieve only values defined by the QDM data element value or value set. Previous versions of QDM datatypes implicitly refer to attributes about a set of items that are included in a value set.</td>
<td>• All Datatypes&lt;br&gt;• Code can also be used with QDM attributes (see individual attribute descriptions)</td>
<td>0..1</td>
</tr>
</tbody>
</table>

Earlier versions did not explicitly state that the datatype attributes only refer to a specific item that is a member of the
<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>defined value set.</td>
<td>The code is applied in the CQL Model Info used to translate the CQL into the Expression Logical Model (ELM) to enhance computer readability. Measure developers do not need to specify any additional 'code.'</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Birth dateTime</td>
<td>The date and time that the patient was born.</td>
<td>● Patient Characteristic Birthdate</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Components</td>
<td>Elements included or documented as part of evaluations or test panels. Examples include: specific questions included in assessments, tests included in a laboratory test panel, observations included in a cardiac exam during a physical examination. Each assessment, diagnostic study, laboratory test, physical exam or procedure may have one or more components.</td>
<td>● Assessment, Performed</td>
<td>0.. *</td>
</tr>
<tr>
<td>Expiration dateTime</td>
<td>The date and time that the patient passed away.</td>
<td>● Patient Characteristic Expired</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Diagnoses</td>
<td>Coded diagnoses/problems addressed during the encounter. The diagnoses attribute is intended to capture ALL diagnoses, including principal diagnosis. Use of the Encounter, Performed: diagnoses attribute and the Diagnosis datatype is redundant for relating the diagnosis to the Encounter, Performed. The Encounter, Performed: diagnoses syntax is preferred. Referencing the same diagnosis using Encounter, Performed (diagnoses attribute) and Diagnosis (datatype) should only occur if the measure must define a specified length of a prevalence period, e.g., The measure must assure that the diagnoses have been present for at least some defined time period before the encounter, and were addressed during the Encounter.</td>
<td>● Encounter, Performed</td>
<td>0.. *</td>
</tr>
<tr>
<td>Discharge Disposition</td>
<td>The disposition, or location to which the patient is transferred at the time of hospital discharge.</td>
<td>● Encounter, Performed</td>
<td>0.. 1</td>
</tr>
</tbody>
</table>

23 CQL supports other data models besides QDM. CQL Model Info is used in implementations to map the particular data model to CQL expressions.
<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dosage</td>
<td>Details of how medication is taken or is to be taken, i.e., the quantity (mg, cc, tablets) to be taken at a single administration.</td>
<td>• Immunization, Administered&lt;br&gt; • Immunization, Order&lt;br&gt; • Medication, Active&lt;br&gt; • Medication, Administered&lt;br&gt; • Medication, Dispensed&lt;br&gt; • Medication, Order&lt;br&gt; • Medication, Discharge&lt;br&gt; • Substance, Administered&lt;br&gt; • Substance, Order&lt;br&gt; • Substance, Recommended</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>Facility Location Facility Locations</td>
<td>The particular locations in a facility in which the diagnostic study or encounter occurs or occurred. Examples include, but are not limited to, intensive care units (ICUs), non-ICUs, burn critical-care unit, neonatal ICU, and respiratory-care unit. Each Encounter, Performed may have 1 or more locations. For example, a patient treated in multiple locations during an individual encounter might be expressed as:&lt;br&gt; • Encounter, Performed: Inpatient Admission&lt;br&gt;   o ICU (location period)&lt;br&gt;   o Non-ICU Admission (location period)&lt;br&gt;   o Rehab (location period)</td>
<td>• Adverse Event&lt;br&gt; • Diagnostic Study, Performed&lt;br&gt; • Encounter, Order&lt;br&gt; • Encounter, Performed&lt;br&gt; • Encounter, Recommended</td>
<td>• 0.. 1 (singular) for all except Encounter, Performed&lt;br&gt; • 0.. * (plural) for Encounter Performed</td>
</tr>
<tr>
<td>Frequency</td>
<td>Indicates how frequently the medication or substance&lt;br&gt; • Is administered to a patient for an active medication (a).&lt;br&gt; • Was administered to the patient (b).&lt;br&gt; • Should be taken by the patient or administered to the patient (c).&lt;br&gt; • Is recommended to be given to the patient (d).</td>
<td>• Medication, Active (a)&lt;br&gt; • Medication, Administered (b)&lt;br&gt; • Substance, Administered (b)&lt;br&gt; • Medication, Discharge (c)&lt;br&gt; • Medication, Dispensed (c)&lt;br&gt; • Medication, Order (c)&lt;br&gt; • Substance, Order (c)&lt;br&gt; • Substance, Recommended (d)</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>id</td>
<td>The identifier of a specific instance of any QDM data element. CQL logic uses the .id reference to specify that a query expects a specific instance of an element to be retrieved. See Appendix A.4 for examples for using id to indicate specific instances of QDM data elements in CQL</td>
<td>• Adverse Event&lt;br&gt; • Allergy/Intolerance&lt;br&gt; • Assessment, Performed&lt;br&gt; • Assessment, Recommended&lt;br&gt; • Patient Experience&lt;br&gt; • Provider Care Experience</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>-----------</td>
<td>------------</td>
<td>-------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
|           | statements with the `relatedTo` attribute. | ● Care Goal  
● Communication, Performed  
● Diagnosis  
● Device, Applied  
● Device, Order  
● Device, Recommended  
● Diagnostic Study, Order  
● Diagnostic Study, Performed  
● Diagnostic Study, Recommended  
● Encounter, Order  
● Encounter, Performed  
● Encounter, Recommended  
● Family History  
● Immunization, Administered  
● Immunization, Order  
● Patient Characteristic  
● Patient Characteristic Birthdate  
● Patient Characteristic Clinical Trial Participant  
● Patient Characteristic Ethnicity  
● Patient Characteristic Expired  
● Patient Characteristic Payer  
● Patient Characteristic Race  
● Patient Characteristic Sex  
● Provider Characteristic  
● Intervention, Order  
● Intervention, Performed  
● Intervention, Recommended  
● Laboratory Test, Order  
● Laboratory Test, Performed  
● Laboratory Test, Recommended |
<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incision date_Time</td>
<td>The date and time of the first incision of the procedure. Incision date_Time is a single point in time available from the Operating Room and/or Anesthesia Record.</td>
<td>• Procedure, Performed</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Length of Stay</td>
<td>The difference of the admission date_time and the discharge date_time for the encounter. This attribute should not be used for outpatient encounters.</td>
<td>• Encounter, Performed</td>
<td>0.. 1</td>
</tr>
<tr>
<td>Location Period</td>
<td>The time the patient arrived at the location to the time the patient departed from the location. Each Encounter, Performed may have 1 or more locations. For example, a patient treated in multiple locations during an individual encounter might be expressed as:</td>
<td>• Encounter, Performed</td>
<td>0.. 1</td>
</tr>
</tbody>
</table>

- Encounters, Performed: Inpatient Admission
  - ICU (location period)
  - Non-ICU Admission (location period)
  - Rehab (location period)

Location Period is used with a Code for the location. The pair of Code and Location Period is associated with each item in the Facility Locations list attribute.
<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medium</td>
<td>A channel used for a communication (e.g., fax, email, etc.)</td>
<td>• Communication, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td>Method</td>
<td>Indicates the procedure or technique used in its performance.</td>
<td>• Assessment, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Diagnostic Study, Performed (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Performed (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Physical Exam, Performed (b)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Procedure, Performed (b)</td>
<td></td>
</tr>
<tr>
<td>Negation Rationale</td>
<td>Indicates the reason that an action was not performed. Only QDM datatypes</td>
<td>• Assessment, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td></td>
<td>that represent actions (e.g., performed, recommended, communication, order,</td>
<td>• Assessment, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td>dispensed) allow the “negation rationale” attribute. The intent is to</td>
<td>• Communication, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>indicate a justification that such action did not happen as expected.</td>
<td>• Device, Applied</td>
<td></td>
</tr>
<tr>
<td></td>
<td>This attribute specifically does not address the presence or absence of</td>
<td>• Device, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td>information in a clinical record (e.g., documented absence of allergies Vs</td>
<td>• Device, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td>documentation about allergies). The syntax in the human readable HQMF is</td>
<td>• Diagnostic Study, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td>address in CQL examples and in the MAT User Guide. Prior versions of QDM</td>
<td>• Diagnostic Study, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>used the syntax, “Procedure, Performed not done.” QDM 5.0 DRAFT uses the</td>
<td>• Diagnostic Study, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td>syntax, “Procedure, not Performed.”</td>
<td>• Encounter, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Note: Some datatypes include both Relevant Time and Author dateTime</td>
<td>• Encounter, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td>attributes. The purpose is to accommodate Author dateTime if the actual</td>
<td>• Encounter Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td>start and stop times are not available when evaluating for feasibility,</td>
<td>• Immunization, Administered</td>
<td></td>
</tr>
<tr>
<td></td>
<td>and also to allow specification of a time for Negation Rationale.</td>
<td>• Immunization, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Performed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Laboratory Test, Recommended</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Administered</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Discharge</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Dispensed</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Medication, Order</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Physical Exam, Order</td>
<td></td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| Ordinality             | The scale in which different procedures are ordered in terms of the qualitative value, as opposed to a ranking performed strictly numerically or quantitatively. | • Procedure, Intolerance  
• Procedure, Order  
• Procedure, Performed  
• Procedure, Recommended | NA          |
| Participation Period   | The time from:  
• enrollmentStartdate – The time the patient enrolled in the program  
• enrollmentEnddate – The time the patient’s enrollement in the program ends | • Participation                                                                                                                             | 0.. 1       |
| Prevalence Period      | Prevalence Period is the time from onset dateTime to abatement dateTime.   | • Allergy/Intolerance  
• Diagnosis  
• Symptom                                                                                                           | 0.. 1       |
| Principal Diagnosis    | The coded diagnosis/problem established after study to be chiefly responsible for occasioning the admission of the patient to the hospital for care. | • Encounter, Performed                                                                                                                       | 0.. 1       |
| Reason                 | The thought process or justification for the datatype. In some measures, specific treatments are acceptable inclusion criteria only if a justified reason is present. Each of these measures uses a value set (often, but not exclusively, using SNOMED-CT®) to express acceptable justification reasons. Other measures specify reasons as justification for exclusions.  
• Examples include patient, system, or medical-related reasons for declining to perform specific actions. Each of these measures also uses a value set to express acceptable justification reasons for | • Assessment, Performed  
• Assessment, Recommended  
• Device, Applied  
• Device, Order  
• Device, Recommended  
• Diagnostic Study, Order  
• Diagnostic Study, Performed  
• Encounter, Order  
• Encounter, Recommended  
• Immunization, | 0.. 1       |
<table>
<thead>
<tr>
<th>Attribute</th>
<th>Definition</th>
<th>Datatype(s)</th>
<th>Cardinality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recipient</td>
<td>The entity (e.g., person, organization, clinical information system, or device) which is the target of a communication</td>
<td>• Communication, Performed</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>Reference Range High</td>
<td>The high bound (inclusive) of values that are considered normal.</td>
<td>• Laboratory Test, Performed</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>Reference Range Low</td>
<td>The low bound (inclusive) of values that are considered normal.</td>
<td>• Laboratory Test, Performed</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>Refills</td>
<td>The number of refills allowed by the prescription.</td>
<td>• Medication, Discharge</td>
<td>• 0.. 1</td>
</tr>
<tr>
<td>relatedTo</td>
<td>An attribute that indicates one QDM data element fulfills the expectations of another QDM data element.</td>
<td>• Care Goal</td>
<td>• 0.. *</td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Relationships</td>
<td>The relationship of the family member to the patient. To ensure compatibility with QRDA reporting constraints, relationship codes should come from the HL7 Personal Relationship Role Type value set (2.16.840.1.113883.1.11.19563).</td>
<td>● Family History</td>
<td>● 0.. 1</td>
</tr>
<tr>
<td>Relevant Period</td>
<td>Relevant Period addresses the time between the start of an action to the end of an action. Each datatype using relevant period defines specific definitions for the start and stop time for the action listed.</td>
<td>● Adverse Event ● Care Goal ● Communication, Performed ● Device, Applied ● Diagnostic Study, Performed ● Encounter, Performed ● Intervention, Performed ● Laboratory Test, Performed ● Medication, Active ● Medication, Administered ● Medication, Order ● Medication, Dispensed ● Patient Characteristic, Payer ● Physical Exam, Performed ● Procedure, Performed ● Substance, Administered</td>
<td>● 0.. 1</td>
</tr>
<tr>
<td>Result</td>
<td>The final consequences or data collected from the datatype. Results can be used in five ways, to express: ● That a result is present in the electronic record but any entry is acceptable ● A numerical result is reported directly as a value. Values may be integers or decimal numbers without units, or as a quantity with a value and units – examples: - 100 mg/dL for a lab test - 140 mmHg for blood pressure - as a percentage (actually a quantity with % as units) - as a ratio (e.g., 1:4, 1:80) Beginning with QDM 5.3 measures use CQL logic to express the mathematical logic.</td>
<td>● Assessment, Performed ● Diagnostic Study, Performed ● Intervention, Performed ● Laboratory Test, Performed ● Physical Exam, Performed ● Procedure, Performed</td>
<td>● 0.. 1</td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>operators to constrain desired results to those above or below a certain threshold</td>
<td>• A result that matches one of a specific set of coded concepts in a value set or a code that matches a direct referenced code</td>
<td>• A result as a dateTime (Assessment, Performed and components)</td>
<td></td>
</tr>
<tr>
<td>Result date time</td>
<td>The time the result report is generated and saved in the database.</td>
<td>• Diagnostic Study, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td>Route</td>
<td>Refers to the path by which the medication or substance should be taken into the body systems, such as intradermally, intrathecally, intramuscularly, intranasally, intravenously, orally, rectally, subcutaneously, sublingually, topically, or vaginally.</td>
<td>• Immunization, Administered</td>
<td>0..1</td>
</tr>
<tr>
<td>Sender</td>
<td>The entity (e.g., person, organization, clinical information system, or device) that is the source of a communication</td>
<td>• Communication, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td>Setting</td>
<td>Where the medication is expected to be consumed or administered. Examples:</td>
<td>• Medication, Order</td>
<td>0..1</td>
</tr>
<tr>
<td></td>
<td>• Inpatient – in the hospital during the current inpatient stay</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Outpatient – in an outpatient facility such as an ambulatory surgical center</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Community – in the ambulatory setting (e.g., at home)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severity</td>
<td>Indicates the intensity of the specified datatype (e.g., persistent, moderate, or severe).</td>
<td>• Allergy/Intolerance</td>
<td>0..1</td>
</tr>
<tr>
<td>Status</td>
<td>Indicates the particular stage of the action represented by the datatype.</td>
<td>• Diagnostic Study, Performed</td>
<td>0..1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Intervention, Performed</td>
<td></td>
</tr>
<tr>
<td>Attribute</td>
<td>Definition</td>
<td>Datatype(s)</td>
<td>Cardinality</td>
</tr>
<tr>
<td>--------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
</tbody>
</table>
| Supply             | The quantity (amount) of therapeutic agent that was provided to a patient (i.e., number of doses, number of tablets or pills, volume of medication) |  - Laboratory Test, Ordered  
  - Laboratory Test, Performed  
  - Laboratory Test, Recommended  
  - Procedure, Performed | 0.. 1          |
|                    | • Indicated to be given during a procedure, diagnostic test, or medication or substance administration |                                                                             |             |
|                    | • Note: Prior versions of the QDM (4.3 and earlier) addressed “dose” with two potential interpretations – (1) the quantity to be taken or administered with each administration and (2) the quantity of medication supplied (i.e., number of doses). QDM 5.0 and subsequent versions clarify the difference by defining “dosage” and “supply,” respectively. |                                                                             |             |
|                    | • Note: Medication, Discharge includes medications the provider has indicated the patient should take after discharge from the hospital. This medication list is part of the discharge instructions provided to a patient. The list may include medication supply if it incorporates medication orders written at discharge even though the supply will not be present for medications the patient already has at home or purchases over-the-counter (without a prescription). |                                                                             |             |
| Target Outcome     | The expected outcome that will indicate the care goal is achieved, or met. The outcome can be expressed using a code, integer, decimal, quantity (value and units), or ratio. |  - Care Goal                                                             | 0.. 1        |
| Type               | The characterization of the reaction (e.g., hypersensitivity, rash, gastroenteric symptoms, etc.) |  - Adverse Event  
  - Allergy/Intolerance                                                   | 0.. 1        |
Appendix A. Special Cases for Consideration

A.1 Use of Components Attributes

A.1.1 Laboratory Test, Performed Example Showing Components

Figure 5. in Section 2.5 uses the components attributes for Laboratory Test, Performed as follows:

Laboratory Test, Performed: Antinuclear Antibody Test

- Components: ANA Homogeneous Pattern (result >= 1:80)
- Components: ANA Speckled Pattern (result >= 1:80)

A.1.2 Encounter, Performed Example Showing Facility Locations

QDM versions for use with CQL model the facility locations attribute in the same way as components. Figure 7 in Section 2.5 shows the use of facility locations with Encounter, Performed as follows:

Encounter, Performed: Inpatient Admission (relevant period)

- Facility Locations: ED (location period)
- Facility Locations: MedSurg (location period)
- Facility Locations: ICU (location period)

A.1.3 Describing Results Associated with Components

The components attribute can be used with:

- Assessment, Performed
- Diagnostic Study, Performed
- Laboratory Test, Performed
- Physical Exam, Performed
- Procedure, Performed

When specifying a measure and determining components, the measure developer should consider the method for specifying results. As noted in Section 4.1, Table 22, results may be reported in several ways:

- That a result is present in the electronic record but any entry is acceptable
- A numerical result reported directly as a value (e.g., 100 mg/dL for a lab test, 140 mmHg for blood pressure), or as a ratio (e.g., 1:4, 1:80). A value is not limited to integers; it can reference fractions (decimals). Beginning with QDM 5.3 measures use CQL logic to express the mathematical operators to constrain desired results to those above or below a certain threshold
- A result that matches one of a specific set of coded concepts in a value set
- A result as a dateTime (Assessment, Performed)
- A result as a percentage (Assessment, Performed)
A.2 Specifying Results with Value Sets

When combining observable entities (assessments, diagnostic studies, laboratory tests, physical exams) or procedures into value sets, the method for specifying results may dictate whether similar sounding entities can be included in the same value set. Specifically, to constrain data elements for a measure to numerical results in the “normal” range, each test within the value set should define “normal” the same way. Since only observable entities with similar definitions of normal or abnormal can be grouped together, the measure may require different data elements to define tests, each with its respective result constraints.

Example:

All of the following are Lyme disease antibody tests identified by LOINC codes with three methods for reporting results. Hence, if the measure needs to constrain a result to a positive or negative response using the numerical value, the three types of tests cannot be included in the same value set. Three different data elements would be required in such an instance.

1. Laboratory test, Performed: Lyme Disease Presence
   - Value set:
     - 11006-4 Borrelia burgdorferi Ab [Presence] in Serum
     - 26006-7 Borrelia burgdorferi Ab [Presence] in Serum by Hemagglutination
   - Result can be reported as present or positive Vs absent or negative

2. Laboratory test, Performed: Lyme Disease IgG/IgM ratio
   - Value set (example):
     - 41279-1 Borrelia burgdorferi IgG/IgM Ab [Ratio] in Serum
   - Result can be reported as a ratio

3. Laboratory test, Performed: Lyme Disease Ab
   - Value set (example):
     - 22121-8 Borrelia burgdorferi Ab [Units/volume] in serum
   - Result can be reported in units/volume

Another option is to represent the results of a set of observable entities using SNOMED-CT concepts for positive and negative and require the local implementation site to interpret results that match the measure criteria. This option may be necessary in situations where the normal ranges vary by site.

Note, these considerations apply only when using value sets for observable entities with QDM datatypes or attributes. QDM datatypes that use direct referenced codes have only one code and, by definition only one result or response method.

When unsure how to proceed with respect to using value sets, consult with a terminologist.

A.3 Evaluating Cumulative Medication Duration

Cumulative medication duration (CMD) is the duration of medication therapy in days for a specific course of treatment. This concept is distinctly different than the lifetime cumulative dose that evaluates the total of all doses of a medication administered over the course of a patient’s lifetime. Lifetime cumulative dose is significant especially with respect to agents that cause significant toxicity above known thresholds. For example, the risk of developing congestive...
heart failure with cumulative doses of doxorubicin exceeding 400 mg/m². Such calculation requires information about all doses ever provided to the patient; it is not the subject of this section of the QDM guide. This section will focus only on the cumulative medication duration that represents a specific course of treatment.

Two scenarios have presented for existing measures:

- A physician should not order more than 90 days’ supply for some medications for patients ≥ 65 years.
- A patient discharged from the hospital with a principal diagnosis of major depressive disorder should receive medication to treat depression for at least 180 days after the discharge.

A.3.1 Options for determining CMD include:

- Use Medication, Order – This QDM datatype addresses medication order or prescriptions as written by the provider; i.e., the amount of medication supply available to the patient and the length of time addressed by that supply based on the frequency of planned administration and the number of available refills. The order is generally the most directly available information from an EHR even though the patient may not fill the prescription or the refills, and actual administration (or self-administration) may differ from the anticipated frequency and duration.

- Use Medication, Dispensed – This QDM datatype addresses the amount of the medication order that has been fulfilled; i.e., the quantity the patient received. While closer to administration, dispensed medication does not necessarily mean the patient’s self-administration will mirror the anticipated frequency or duration.

- Use Medication, Administered – This QDM datatype addresses the medication used to treat the condition identified. The information is generally available for medication administered by a clinician or direct observed therapy during which a clinician watches a patient take the drug. Alternatively, a patient’s documentation of self-administration might be used if such were available.

A.3.2 CMD Data Validity and Reliability Considerations

All of these calculations are confounded by changes to the medication regimen that occur based on telephone discussions between the patient and physician and unstructured clinical notes to represent the changes. Measure developers should address such issues when determining the feasibility, validity and reliability of their measures.

Data quality may impact measure reliability in determining a clinician’s compliance with expected prescribing guidelines. Measure developers using cumulative medication duration that relies on medication order frequency data should evaluate the impact of conflicting or missing data in the prescription. Surescripts’ analysis of 26,341 prescription (0.87%) found 66.1% with inappropriate content including 19% with patient directions (i.e., free text instructions) that conflicted with directions included in the designated standard structured field intended for that

---

Each measure developer should evaluate validity and reliability of the data during measure testing; the QDM data model and CQL expressions cannot evaluate such data quality issues. Note, the measure developer can use a CQL expression to request information about data that are missing or inconsistent with accepted standards-based content.

### A.3.3 Calculating CMD

#### A.3.3.1 CMD for Medication, Order

1. **Preferred Method** - Calculate based on start and stop times (Relevant Period) for Medication, Order

   - **Relevant Period for Medication, Order** is defined as:
     - `startTime` = when the first administration of the medication is expected. The first administration may be expected at the time of the order or at a specified future date (i.e., the active time for the order); such information should be identified in the medication order. If the startTime is not specified in the order, the `startTime` defaults to the Author dateTime (the time the order is signed).
     - `stopTime` = when the medication supply provided by the medication order is expected to be completed, including all fulfillments covered by the number of refills.

     Note that when calculating cumulative medication duration, the `stopTime` may be present directly in the medication order. If the `stopTime` is not available, the duration in days is the difference between the Relevant Period start and stop times multiplied by \((1 + \text{the number of refills})\).

   - **CMD** = \([\text{end of Relevant Period} - \text{start of Relevant Period}]\)

     Result provided in days

2. **Alternate, Non-preferred Method** - Calculate based on frequency and supply

   - **CMD** = \([\# \text{doses prescribed (supply)} / \text{daily frequency}] \times [1 + \# \text{refills}]\)

     This option uses existing QDM attributes but it is complex since frequency is listed as an abbreviation or phrase (e.g., every 8 hours, q8 hours, TID). The representation may vary by site and must be translated into the number of doses per day.

     Uses existing attributes but too complex and inconsistent data availability

#### A.3.3.2 CMD for Medication, Dispensed

- **Relevant Period for Medication, Dispensed** addresses:
  - `startTime` = when the first administration of the medication is expected. If not specified, the `startTime` defaults to the Author dateTime (the time the dispensing event is documented).

---

o `stopTime` = when the medication supply provided by the dispensing event is expected to be completed.

Note that when calculating cumulative medication duration, the medication dispensed stopTime may be present directly in the fulfillment record. If the stopTime is not available, the duration in days is the difference between the Relevant Period start and stop times. The record may indicate which of the available refills the fulfillment represents but each dispensing event is unique. Therefore, the measure developer should use CQL logic to address multiple dispensing events over a period of time.

- CMD = the sum of all dispensing events with each event providing [end of Relevant Period – beginning of Relevant Period] in days.

### A.3.3.3 CMD for Medication, Administered

- The Relevant Period addresses:
  o `startTime` = when a single medication administration event starts (e.g., the initiation of an intravenous infusion, or administering a pill or IM injection to a patient);
  o `stopTime` = when a single medication administration event ends (e.g., the end time of the intravenous infusion, or the administration of a pill or IM injection is completed - for pills and IM injections, the start and stop times are the same)

Measure developers should address multiple administrations over a period of time using CQL logic. CMD = [date of last Medication, Administered – date of first Medication, Administered]

### A.3.4 Cumulative Medication Duration Examples

- **Medication Dispensed**: Ibuprofen 500 mg every 8 hours to begin after dental procedure in one week from the order date and continue for 20 days with 2 refills, dispense 60.
  - Relevant Period start --- effectiveTime low = date dispensed
  - Relevant Period end --- effectiveTime high = date dispensed plus 20 days

  Cumulative medication duration = 20 days (effectiveTime high date - effectiveTime low date) --- Must sum each dispensing event to = 60 days

- **Medication Order**: Ibuprofen 800 mg every 8 hours (no start date noted so assume start = now) for 10 days with 1 refill, dispense 30
  - Relevant Period start --- effectiveTime low = today’s date
  - Relevant Period end --- effectiveTime high = today’s date + 10 days + 1 refill of 10 days

  Cumulative medication duration = 20 days

- **Medication Order**: Ibuprofen 800 mg every 12 hours (no start date noted so assume start – now) for 7 days (no refill), dispense 14
  - Relevant Period start --- effectiveTime low = today’s date
  - Relevant Period end --- effectiveTime high = today’s date + 7 days + no refills
Cumulative medication duration = 14 days

Note, the calculation subtracts the end date from the start date and provides a result in number of calendar days. Example:
Ibuprofen 800 mg every 12 hours, dispense 14 written at 17:00 on August 7 would include start dateTime as 201708071700 and the end dateTime as 20170814. For this example, the calculated result is August 14 minus August 7 = 7 days.

A.4 Use of relatedTo Attribute

The relatedTo attribute allows an expression to indicate that one QDM data element fulfills the expectations of another QDM data element. The attribute is allowed for datatypes:

- Assessment, Performed
- Communication, Performed
- Care Goal

The entity to which the QDM data element is relatedTo must be expressed by any fully defined QDM data element including its datatype and associated value set or direct referenced code and any required attributes. A QDM data element cannot be directly relatedTo an attribute. This entity (i.e., relatedTo QDM datatype:code) is identified in the CQL as an instance identifier (i.e., a specific instance of the activity defined by the QDM data element). The instance identifier is identified as .id as shown in the following two examples.

Two examples using CQL expressions:

1. Referral Management: Assess whether a specialist consultation has been performed as ordered for a patient with newly diagnosed hepatitis C. Based on workflow practices, measure developers have evaluated the use of Encounter, Order; Intervention, Order; and Communication, Performed, to express the request for a consult and receipt of the resulting consult report. Since adding the QDM datatype Assessment, Performed, the resulting consult report is more appropriately referenced as an assessment. The following three examples show the use of Intervention, Order; Encounter, Order; and Communication, Performed, to express referral management. Note the use of the reason attribute to indicate the reason, or topic of the consult. Intervention, Order or Encounter, Order may be more consistent with requests for consults as managed in clinical software. Measure developers should assess feasibility for retrieving the level of detail desired.

   a. Referral Management using Intervention, Order

   define “Referral”:
   [“Intervention, Order”: “Patient referral to specialist”] R
   define “Assessment”:
   [“Assessment, Performed”: “Consultation (reason: chronic hepatitis C disorder)”] A
   define “Fulfills With Assessment”:
   “Assessment” A
   where exists (“Referral” R where R.id in A.relatedTo) a – SNOMED-CT 103696004 (2017-03)
   b – SNOMED-CT 11429006 (2017-03)
   c – SNOMED-CT 128302006 (2017-03)
b. Referral Management using Encounter, Order

define “Referral”:
[“Encounter, Order”: “Patient referral to specialist”] R

define “Assessment”:
[“Assessment, Performed”: “Consultation” (reason: chronic hepatitis C disorder)”] A

define “Fulfills With Assessment”:
“Assessment” A
where exists (“Referral” R where R.id in A.relatedTo)
a – SNOMED-CT 103696004 (2017-03)
b – SNOMED-CT 11429006 (2017-03)
c – SNOMED-CT 128302006 (2017-03)

c. Referral Management using Communication, Provider to Provider

define “Referral”:
[“Communication, Performed”: “Patient referral to specialist” (sender=”primary care provider”, receiver=”specialist”)] R

define “Assessment”:
[“Assessment, Performed”: “Consultation” (reason: chronic hepatitis C disorder)”] A

define “Fulfills With Assessment”:
“Assessment” A
where exists (“Referral” R where R.id in A.relatedTo)
a – SNOMED-CT 103696004 (2017-03)
b – SNOMED-CT 11429006 (2017-03)
c – SNOMED-CT 128302006 (2017-03)

2. Assess if a Care Goal of weight loss has been established for a patient with a diagnosis of essential hypertension

define “Essential Hypertension”:
[“Diagnosis”: “Essential Hypertension”] E

define “Care Goal”:
[“Care Goal”: “Weight loss”] C

define “Fulfills With Care Goal”:
“Care Goal” C
where exists (“Diagnosis” E where E.id in C.relatedTo)
da – grouped value set 2.16.840.1.113883.3.464.1003.104.12.1011 (2017504)
e – SNOMED-CT 363806002 (201703)
Appendix B. Change Log

B.1 Changes in QDM 5.4

The Quality Data Model, Version 5.4 specification contains the following changes from the Quality Data Model, Version 5.3, Annotated:

- Added QDM datatype, Assessment, Order to be consistent with modeling of other QDM datatypes and allow for an order, not merely a recommendation for a future assessment.
- Added “setting” attribute to QDM datatype Medication, Order to allow a measure developer to specify the setting in which the medication order is expected to be consumed by the patient.
- Removed the “method” attribute from all datatypes with a recommended or order context (Assessment, Recommended; Assessment, Order; Diagnostic Study, Order; Diagnostic Study, Recommended; Laboratory Test, Order; Laboratory Test, Recommended; Procedure, Order; Procedure, Recommended; Substance, Recommended; Physical Exam, Order; Physical Exam, Recommended). QDM 5.4 retains the “method” attribute for Assessment, Performed; Diagnostic Study, Performed; Laboratory Test, Performed; Physical Exam, Performed; Procedure, Performed
- Provided additional guidance to the use of the QDM datatype Device, Applied.
- Provided additional guidance for the use of the QDM datatype, Symptom.
- Removed the “supply” attribute from QDM datatypes for which clinical data does not provide supply information – Medication, Active; Medication, Administered; Substance, Administered; Substance Recommended; and Immunization, Administered. QDM 5.4 retains the “supply” attribute for Medication, Dispensed; Medication, Order; Substance, Order; and Medication, Discharge.
- Removed the “anatomical approach site” attribute from QDM datatypes Device, Applied; Procedure, Performed; Procedure, Order; Procedure, Recommended. The pre-coordinated terminology for the procedure provides approach site information. Moreover, specific location terminology and device terminology can meet the current use cases without this attribute which has never been used in any CMS program measures.
- Retained the distinction between QDM categories Intervention and Procedure even though interoperability standards do not provide any differentiate between the two concepts. Measure developers find the QDM definitions useful and believe that clinicians reading the human readable eCQMs will find the distinction useful.
- Merged three Communication QDM datatypes – Communication, Provider to Patient; Communication, Patient to Provider; Communication, Provider to Provider into a single QDM datatype, Communication, Performed. This new datatype allows greater and more explicit expressivity adding attributes category (type of message – alert, notification, reminder, instruction, etc.), medium (channel of communication – e.g., email, fax, etc.), sender (entity that is the source of the communication), and recipient (entity that is the receiver of the communication) and assigned a relevant period to Communication, Performed.
- Provided some guidance on the use of the QDM substance category with existing use cases (blood product administration and exclusive breast milk feeding for newborn infants in the hospital).

B.2 Changes in QDM 5.3 Annotated

The Quality Data Model, Version 5.3 Annotated specification contains the following changes from the Quality Data Model, Version 5.3:

- Added Relevant Period to Medication, Order and Medication, Dispensed
- Removed Active date/Time from Medication, Order
• Changed QDM attributes with cardinality greater than 1 to plural (changes attributes diagnosis to diagnoses, component to components, and facility location to facility locations for Encounter, Performed)

• Added description of how QDM 5.3 data elements are expressed in tooling and in the HQMF and the CQL logic

• Added explanation for new timing attributes – periods

• Added explanation for new cardinality specification for QDM attributes

• Added explanations for special cases – use of components, specifying results with value sets, and considerations for cumulative medication duration

• Added clarifications for attributes relatedTo, Target Outcome, Result, Facility Locations, Location Period, and Code

• Added id attribute to all datatypes to allow expression of specific instances for any QDM data element in the CQL logic

B.3 Changes in QDM 5.3

The Quality Data Model, Version 5.3 specification contains the following changes from the Quality Data Model, Version 5.02 Proposed DRAFT specification:

• Removed Encounter, Active as this datatype can be managed by using Encounter, Performed with no end dateTime.

• Added Author dateTime to three QDM datatypes: Diagnosis, Allergy/Intolerance, and Adverse Event to account for situations in which the onset and abatement times are unknown.

• Added a QDM datatype for Participation and new attribute to support: Participation Period.

• Removed Radiation Dose and Radiation Duration attributes as they have not been used and the use case is not defined. The concepts can be expressed using Assessment, Performed or Diagnostic Test, Performed.

• Added guidance to Diagnosis attribute for Encounter, Performed to make it explicit that is intended to capture ALL diagnoses, including principal diagnosis.

• Modeled the Location attribute for Encounter, Performed to allow for describing the patient’s presence in more than one location during an individual Encounter.

• Removed Reason attribute from Encounter, Performed since it has never been used and the Encounter, Performed (diagnosis) attribute allows description of reason.

• Clarified that the QDM category Laboratory Test refers to tests performed for the subject of record.

• Added guidance to QDM data flow attributes to consider feasibility before using the attributes.

• Assigned cardinality to all QDM attributes – All datatypes use cardinality of 0..1 except:
  o Diagnosis (0.. *)
  o Facility location (only for the Encounter, Performed datatype – 0.. *)
  o Location Period (0.. *)
  o relatedTo (0.. *)

B.4 Changes in QDM 5.02 Proposed DRAFT

The Quality Data Model, Version 5.02 Proposed DRAFT specification contains the following changes from the Quality Data Model, Version 5.01 specification:
• Added authorTime to nine QDM datatypes that include the negation rationale attribute: Device, Applied; Diagnostic Study, Performed; Intervention, Performed; Encounter, Performed; Laboratory Test, Performed; Medication, Administered; Physical Exam, Performed; Procedure, Performed; Substance, Administered. The purpose is to more easily allow specification of a time for Negation Rationale.

• QDM 5.02 AuthorTimeErrata – changed all Author Times to Author dateTime to be consistent with expectations for data capture. Also, corrected Medication, discharge attribute from dose to dosage.

B.5 Changes in QDM 5.01 Proposed DRAFT

The Quality Data Model, Version 5.01 Proposed DRAFT specification contains the following changes from the Quality Data Model, Version 5.0 specification:

• Added Components attribute to Assessment, Performed; Diagnostic, Performed; Laboratory Test, Performed. The Components attribute includes a code and result and, for laboratory test, performed includes optional reference range high and reference range low.

B.6 Changes in QDM 5.0 DRAFT

The Quality Data Model, Version 5.0 DRAFT specification contains the following changes from the Quality Data Model, Version 4.3 specification:

• All logic content is removed. Logic is managed using CQL. The reader should refer to the CQL Formatting Guide to address logic.

• Added Allergy/Intolerance datatype

• Added Adverse Event datatype

• Removed all previous datatypes referencing Allergy, Intolerance and Adverse Reaction

• Added “code” attribute to all datatypes

• Added result response = dateTime or percentage to Assessment, Performed (result)

• Added Care Goal target outcome response = percentage

• Modified datatype timings to address:
  – Prevalence period for all datatypes requiring onset dateTime and abatement dateTime
  – Relevant period for action datatypes with specific timing descriptions of start and stop elements of relevant period
  – Author dateTime for datatypes addressing orders, recommendations and other points in time

• Removed Transfer to datatype

• Removed Transfer from datatype

• Added attributes for Encounter, Performed and Encounter, Active – Admission Source and Discharge Disposition (replaces Discharge status)

• Cumulative Medication Duration has been removed as the concept can be expressed directly using existing QDM datatypes and attributes with CQL logic expressions.
B.7 Changes in QDM 4.3

The Quality Data Model, Version 4.3 specification contains the following changes from the Quality Data Model, Version 4.2 specification:

- Added Assessment, Performed datatype
- Added Assessment, Recommended datatype
- Removed Risk Category Assessment datatype
- Removed Functional Status, Performed datatype
- Removed Functional Status, Recommended datatype
- Removed Functional Status, Ordered datatype
- Added clarification of timing for datatypes with Order actions
- Added clarification of feasibility requirements for all datatypes with Recommended actions

B.8 Changes in QDM 4.2

The Quality Data Model, Version 4.2 specification contains the following changes from the Quality Data Model, Version 4.1.2 specification:

- Enhanced support for encounter diagnoses and principal diagnoses
- Re-specified the Diagnosis datatypes
- Re-specified the Diagnosis, Family History datatype (now Family History)
- Re-specified the Symptom datatypes
- Added the Immunization category and datatypes
- Added reference range attributes to Laboratory Test, Performed
- Removed patient preference and provider preference attributes from all datatypes
- Removed negation rationale from inappropriate datatypes
- Fixed transposed descriptions of Transfer From and Transfer To datatypes

B.9 Changes in QDM 4.1.2

The Quality Data Model, Version 4.1.2 specification contains the following changes from the Quality Data Model, Version 4.1.1 specification:

- Removed anatomical approach site attribute from all Physical Exam datatypes
- Removed method attribute from Procedure, Intolerance datatype
- Clarified the meaning of start dateTime and stop dateTime in all Order datatypes
- Updated Appendix B to indicate where quantity-based temporal comparisons can and cannot be used

B.10 Changes in QDM 4.1.1

The Quality Data Model, Version 4.1.1 specification contains the following changes from the Quality Data Model, Version 4.1 specification:

- Replaced DateDiff and TimeDiff functions with DateTimeDiff
• Specified DateTimeDiff function to be used in measure observations only
• Added new section describing the use of attribute filters
• Added support for filtering date/time attributes by date
• Re-specified Overlaps to interpret missing end dates/times as ongoing

B.11 Changes in QDM 4.1
The Quality Data Model, Version 4.1 specification contains the following changes from the Quality Data Model, Version 4.0 specification:

• New QDM functions and operators
  – Union
  – Intersection
  – Fulfills
• New QDM temporal operators
  – Starts Concurrent With End Of
  – Ends Concurrent With Start Of
• Clarified names of existing QDM temporal operators
  – Starts Before or Concurrent With \( \rightarrow \) Starts Before or Concurrent With Start Of
  – Starts After or Concurrent With \( \rightarrow \) Starts After or Concurrent With Start Of
  – Starts Before or During \( \rightarrow \) Starts Before End Of
  – Ends Before or Concurrent With \( \rightarrow \) Ends Before or Concurrent With End Of
  – Ends After or Concurrent With \( \rightarrow \) Ends After or Concurrent With End Of
  – Ends Before or During \( \rightarrow \) Ends Before End Of
• Modified existing datatypes
  – Added signed datetime and active datetime to Medication, Order
  – Added target outcome to Care Goal
  – Renamed reason to cause in Patient Characteristic Expired
  – Clarified or removed ambiguous attributes (see QDM-46 for details)
• Removed System Characteristic Datatype
• Removed Measurement Start Date and Measurement End Date in favor of MeasurementPeriod
• Restricted QDM elements to use only one attribute at a time
• Added an appendix covering the representation of Cumulative Medication Duration

B.12 Changes in QDM 4.0
The Quality Data Model, Version 4.0 specification contains the following changes from the Quality Data Model, December 2013 specification:

• New document template and layout to improve readability
• Updated the wording in some sections to improve clarity
• Added definitions for all QDM functions, operators, and timing relationships
• New high-level QDM language constructs
  – Variable assignment
  – Inline comments
• New QDM functions and operators
  – Age At
  – Satisfies Any
  – Satisfies All
• New QDM temporal operators
  – Starts Before or Concurrent With
  – Starts Before of Concurrent With End Of
  – Starts After or Concurrent With
  – Starts After or Concurrent With End Of
  – Ends Before or Concurrent With
  – Ends Before or Concurrent With Start Of
  – Ends After or Concurrent With
  – Ends After or Concurrent With Start
• Incorporated guidance from eCQM Measure Logic and Implementation Guidance and Technical Release Notes, Version 1.7\textsuperscript{26}
  – Specific occurrences
  – Calculation of time intervals

## Acronyms

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Description</th>
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<tbody>
<tr>
<td>AHRQ</td>
<td>Agency for Healthcare Research and Quality</td>
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<tr>
<td>CDS</td>
<td>Clinical Decision Support</td>
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<tr>
<td>CMD</td>
<td>Cumulative Medication Duration</td>
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<tr>
<td>CMS</td>
<td>Centers for Medicare &amp; Medicaid Services</td>
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<tr>
<td>CPT®</td>
<td>Current Procedural Terminology</td>
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<tr>
<td>CQL</td>
<td>Clinical Quality Language</td>
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<td>eCQI</td>
<td>Electronic Clinical Quality Improvement</td>
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<td>Electronic Clinical Quality Measure</td>
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<td>EHR</td>
<td>Electronic Health Record</td>
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<td>FHIR®</td>
<td>Fast Health Interoperability Resources</td>
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<td>Health Quality Measure Format</td>
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<td>International Classification of Diseases, Ninth Revision</td>
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<tr>
<td>ICD-10</td>
<td>International Classification of Diseases, Tenth Revision</td>
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<tr>
<td>ISO</td>
<td>International Organization for Standardization</td>
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<td>MAT</td>
<td>Measure Authoring Tool</td>
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<td>ONC</td>
<td>Office of the National Coordinator for Health Information Technology</td>
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<tr>
<td>PHR</td>
<td>Personal Health Record</td>
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<td>QDM</td>
<td>Quality Data Model</td>
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<tr>
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<td>Quality Reporting Document Architecture</td>
</tr>
<tr>
<td>SI</td>
<td>International System of Units</td>
</tr>
<tr>
<td>SNOMED-CT®</td>
<td>Systematized Nomenclature of Medicine – Clinical Terms</td>
</tr>
<tr>
<td>UCUM</td>
<td>Unified Code for Units of Measure</td>
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