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**HL7 Virtual Medical Record for Clinical Decision Support (vMR-CDS) Templates, Release 1 - US Realm**

January 2014

**U.S. Realm Draft Standard for Trial Use (DSTU) Ballot**

**Sponsored by:**

**HL7** **Clinical Decision Support Work Group and HL7 Templates Work Group**

**in collaboration with the Health and Human Services Standards and**

**Interoperability Framework Health eDecisions Working Group**

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**Note Regarding Realm:**

Per guidance from the HL7 Technical Steering Committee, this specification is being balloted as a U.S. Realm specification. It is anticipated that future releases of the specification may be balloted in the Universal Realm.

**Acknowledgements, Copyrights**

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# Introduction

A Virtual Medical Record (vMR) is a data model for representing the data that are analyzed and/or produced by clinical decision support (CDS) engines. The purpose of the vMR effort is to define a standard vMR that (i) can be used across CDS implementations and (ii) is simple and intuitive for a typical CDS knowledge engineer to understand, use, and implement.

This specification defines vMR templates that constrain the base vMR model to facilitate semantic interoperability, similar to how Consolidated Clinical Documentation Architecture (C-CDA) templates constrain the base CDA model. The vMR templates are informed by the templates defined for the C-CDA and Quality Reporting Document Architecture (QRDA) standards as specified in the “HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation, Release 1.1 - US Realm” and the “HL7 Implementation Guide for CDA® R2: Quality Reporting Document Architecture - Category I (QRDA) DSTU Release 2 (US Realm)” standards.

## Purpose

The purpose of this specification is to define a set of templates for the vMR.

## Methodology

This specification leverages a template development methodology that is being defined by the HL7 Templates Work Group. This project is informed by, and contributing to, the specification of this template development methodology by the HL7 Templates Work Group.

## Intended Audience

The intended audience for this implementation guide is CDS implementers.

### Requisite Knowledge

Knowledge of the HL7 vMR Logical Model, Release 2, Version 3.0 is a prerequisite.

### Referenced Standards

The templates are defined in the context of the following specifications:

* HL7 vMR Logical Model Release 2, Version 3.0
* HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation, Release 1.1 - US Realm
* HL7 Implementation Guide for CDA® R2: Quality Reporting Document Architecture - Category I (QRDA) DSTU Release 2 (US Realm)

For documentation on the meaning of vMR model elements, as well as examples, please refer to the HL7 vMR Logical Model specification, as well as the HL7 vMR XML specification.

## Organization of this Specification

This specification defines a set of vMR templates and provides a narrative explanation of a sample template.

## Definitions and Acronyms

|  |  |
| --- | --- |
| **Term** | **Definition** |
| **C-CDA** | Consolidated Clinical Document Architecture |
| **CDS** | Clinical Decision Support |
| **OID** | Object Identifier |
| **QRDA** | Quality Reporting Document Architecture |
| **vMR** | Virtual Medical Record |

# vMR TEMPLATES

The vMR templates are defined in the accompanying Excel spreadsheets. Also, XML instance examples of select templates are available in the accompanying supplemental files.

Of note, we anticipate the development of vMR templates to be an ongoing, iterative process. In particular, the templates listed in Section 2.2 have been identified as potentially being useful and are under consideration for future addition to the vMR templates. Moreover, the potential leveraging of detailed clinical models within vMR templates is under investigation.

## Overview of Templates

As a matter of explanation, each template contains the following information:

* Various meta-data about the template, including template name, OID, description, effective date, status, version, and expiration date
* Data expected to be included in the vMR if no constraints are specified
  + For example, the SimpleLabResult template specifies that if no constraints are specified, all lab results available up to the current time should be provided
* Constraints that may be specified to restrict the data that are provided using the template
  + For example, the SimpleLabResult template specifies that restricting the search back period is allowed using the observationEventTime. This would allow, for example, a CDS service provider to specify that only lab results from the past 1 year are needed.
  + Similarly, the SimpleLabResult template specifies that restricting the data according to the observationFocus is allowed. This would allow, for example, a CDS service provider to specify that only Hemoglobin A1c and LDL cholesterol lab results are needed.
  + The approach to these constraints is specified in the vMR Logical Model within the CDS Input Specification component of the model.
* For each included vMR data element, the following constraints are specified. Note that many of these constraints are defined in Chapter 2 of the “HL7 Version 3 Standard: Refinement, Constraint, and Localization (RCL) to V3 Messages, R2” specification (<http://www.hl7.org/implement/standards/product_brief.cfm?product_id=76>). Those constraints defined in that chapter are specified with an asterisk below:
  + The data element’s path in the vMR data model
  + The data element’s cardinality\*
  + Whether the data element is mandatory (i.e., must be present in instances and be non-null)\*
    - Because there are no nullFlavors in the vMR, a data element is always mandatory if the cardinality is 1..X, and always non-mdantory if it is 0..X
  + The data element’s conformance\*
    - R means required\*. This conformance requirement is placed on a data element when at least one of the following conditions applies:
      * Cardinality is 1..X
      * Cardinality is 0..X, but it may be conditionally required based on other content within the template
      * Cardinality is 0..X, but is required when the data is available
  + Whether the data element has a fixed value in all instances\*. An example of a fixed value is where the template ID of an instance using a particular template must always be the same.
  + Whether the data element has an ad-hoc constraint, which means that it has a constraint that depends on other content within the template. Note that such conditional constraints are currently specified using narrative free text.
  + The data element’s data type\*
  + A narrative specification of further constraints, in particular value restrictions
  + Comments, including in particular any deviation from C-CDA or QRDA value restrictions. C-CDA and QRDA templates were explicitly reviewed in the development of vMR templates to maximize semantic interoperability.

Also of note:

* The subject of a clinical statement is identified in the EvaluatedPerson object which contains the clinical statement. Therefore, individual clinical statements do not specify a patient identifier. If one desired this attribute to be available with each clinical statement, this could be done using the clinical statement.evaluatedPersonId attribute.
* The provider associated with a clinical statement can be identified using a related entity relationship. However, because such information is typically not required for clinical decision support, it is not included in this specification.

## Additional Templates Under Review and/or Development

The following templates have been identified as being potentially useful for CDS and are currently under review and/or development. Those who are interested in contributing to the continued development of vMR templates and their associated value sets are encouraged to contact the CDS Work Group through its HL7 list-serv.

* Templates for images and imaging results, including as a part of an encounter
* Templates related to performance on quality measures
* Templates related to results returned by a CDS service
* Templates related to quality measurement and continuity of care
* Templates for clinical assessments (e.g., for nursing documentation of the skin, stool, wound, etc.)
* Templates to support handling of unvalidated data, where there is no authenticator
* Templates to support indicating that the data was authored from a device
* Templates to support orders, such as for medication orders, lab orders, and respiratory care orders
* Additional templates to support proposals
* Enhancements to problem templates to allow prioritization of active problems for a patient within a given context, such as by a CDS system. Also, enhancements to enable identification of the clinical domain and user expertise of those who identify each active problem.
* Enhancements to immunization-related templates, for example, to enable expressing the following:
  + The patient does not need a vaccination of a certain type because of reason X
  + The patient has completed a vaccination series, is immune against disease X, and no further doses are needed
* Enhancements to observation templates to allow the expression of result report times (as opposed to physiologic times) and associated report statuses such as ‘preliminary’, ‘final’.
* Additional templates to represent nutrition proposals, orders, and events.
* Templates for radiology and chemotherapy.
* Templates for adverse events which are not caused by exposure to some agent – e.g., falls, hospital acquired infections.
* Device-acquired vital sign measurements (e.g., resulting from work performed by Project 850)