Model Driven Data Management in Healthcare

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Keywords: ISO11179, Metadata, Model Driven Engineering, UML.

Abstract: Healthcare research depends on the availability of data that is of high quality, that is easy to query, consistent and current. Traditionally, healthcare data has relied on multiple diverse datasets being integrated by domain experts. These integration processes are executed with a high degree of human involvement, integrating datasets can be time-consuming and can result in the introduction of errors into the data. This paper describes work to build an integration toolset for healthcare datasets based on the ISO11179 Standard for metadata registries. It describes issues encountered whilst implementing the standard and shows how these short-comings were overcome by using techniques from the field of Model Driven Engineering (MDE).

1 INTRODUCTION

A fundamental problem in the UK and elsewhere is how to make clean data from heterogeneous healthcare datasets available to researchers easily and quickly. Much of this information is from research sources such as clinical trials and electronic patient records, however currently researchers can spend more time wrangling and cleaning the data than is spent in analysis, some reports put this at between 60-80% of time taken in analysis tasks (Press, 2016). Analysis generally involves python and R scripts which are unique to the researcher that wrote them, if new information is not in the form that the researcher anticipated then the text has to be re-written, and checked for accuracy. In an ideal world data would be input to a data warehouse in a form that allows the same query to be run repeatedly, because the data that comes into the repository is guaranteed to be in the same format, and if the format changes the dataset and query can be updated to take account of this without a lengthy script re-write.

One approach to tackling this problem is the use of the standardized dataset, the idea being is that a set of data items are defined in the standard, and all reporting of any such data-items is made to confirm to the standard. This will enforce some simple rules such as a patient identifier in the NHS needs to be an integer of a certain length, conforming to a certain set of rules, very often encoded with a regular expression. This approach helps enormously, but its application hasn’t been entirely successful to date for several reasons. Firstly, it is impossible currently to mandate that everyone uses the same standard, or set of standards. Secondly, some standards are strong in some areas and weak in others. Thirdly standards evolve.

There are currently a number of different data standards in healthcare, each one having emerged from a different specialist area, such as pathology or pharmaceutical research. If data from heterogeneous datasets are presented in one, and only one of these standards, but for instance, in different formats, some being in XML, some in CSV and some in RDF, then they can relatively easily be fused with data from other datasets conforming to the same standard. Where data standards have clinically endorsed mappings between them, then data from different datasets can easily be merged. Where data is available in datasets which do not comply with a common standard, then a set of mappings needs to be made to merge that data. This is normally carried out using standard Extract, Load and Transform (ETL) techniques.

Some dataset standards, such as the OMOP CDM (OHDSI, 2018) claim the title of Common Data Model (CDM), and aim to be the only dataset definition for the whole industry. In addition, there are many datasets that have been built up within research organisations or within particular clinical specialist areas that are in use throughout the U.K.’s National Health Service (NHS). Certain of these clinical datasets were used in this investigation, in particular COSD, FHIR, SNOMED CT and the NHS Data Dictionary.
This research started out to discover better ways of integrating datasets, in particular if automated techniques using metadata could be applied to managing, wrangling and cleaning data used in clinical data analytics. Some work had already gone into applying the ideas in ISO/IEC 11179, the ISO standard for metadata registries, and so initially this was adopted as a way forward in the research program. We built and tested the interchange of several datasets using an ISO11179 compliant metadata registry with mixed results, we examined problems which arose, and then built a revised metadata registry built on model driven engineering principles. Repeating the same experiments we found that the improved metadata registry performed more efficiently, and was easier for clinicians and healthcare business analysts to use. We achieved the main research goals of identifying automated techniques for using metadata to manage, wrangle and clean dataset far more efficiently than the techniques previously being used. We then looked at the problems we had with the ISO11179 standard, and examined what improvements can be made to the ISO11179 standard to make it more workable and effective in achieving the purposes stated in ISO/IEC11179-1:2015E.

This paper is split into several sections; the section on related work details previous research efforts to achieve interoperability between heterogeneous clinical datasets ISO11179 and Model Driven Engineering techniques. In the section called Background the nature of the problem is described in detail, including a short summary of some of the main dataset standards encountered in the course of this research. In the section titled ISO11179: ISO Standard for Metadata Registries the ISO11179 approach to interoperability is examined. The next section Evaluation evaluates the effectiveness in applying ISO11179 to clinical dataset management. A review of the results is given in the next section Results, firstly of the overall research effort, and secondly on the role of ISO11179. Lastly, there is a section outlining Conclusions and suggesting future work. The main contributions of this research are as follows: first, providing a set of techniques for automating the management of datasets using metadata, and more specifically using tools built around a metadata registry, second, providing a record of experiences in applying the ISO11179 to medical dataset management; third, identifying shortcomings in the ISO11179 metadata registries standard; fourth, identifying ways to overcome these shortcomings using model driven engineering principles, and last, the design of an improved metamodel for healthcare metadata registries.

2 RELATED WORK

The work described in this paper has been informed by work carried out by colleagues at the University of Oxford on the CancerGrid project (Davies et al., 2014), where an ISO/IEC 11179-compliant metadata registry was developed as detailed in (Davies et al., 2015). Initially the test software for these studies was developed using the eXist XML database, but it was found to have problems scaling once the number of data elements increased over about 10,000, and so new work was carried out to build a more scalable metadata registry using java-based web frameworks.

One of the earliest efforts to apply the principles of ISO11179 in practice was the caBIG initiative by the National Cancer Institute in the USA, (Kunz et al., 2009); they built a software development kit which allows developers to build web service stubs around data elements, (Komatsoulis et al., 2008), however it doesn’t appear to have been widely adopted. Indeed there are very few examples of ISO11179 metadata registries in practice, one study has used semantic web technology to integrate metadata registries, Sinaci and Erturkmen (Sinaci and Erturkmen, 2013) describe a semantic metadata registry framework where Common Data Elements (CDEs) are exposed as Linked Open Data resources. CDEs are described in the Resource Description Framework (RDF), and can be queried and interlinked with CDEs in other registries using the W3C Simple Knowledge Organization System (SKOS). An ISO11179 ontology has been defined as part of the framework, and the Semantic MDR has been implemented using the Jena framework.

Metadata Registries, such as those conforming to the ISO11179 standard, can help to solve the problem of data incompatibility, provenance and compliance, as is indicated in studies such as those conducted by Ulrich et al. (Ulrich et al., 2016). In this study a hybrid architecture consisting of an ISO 11179-3 conformance MDR server application for interactively annotating and mediating data elements and the translation of these data elements into Fast Health Interoperability Resources (FHIR) (HL7-FHIR-Foundation, 2017) resources was used to manage data for the North German Tumor Bank of Colorectal Cancer.

Tao et al. (Tao et al., 2011) present case studies in representing HL7 Detailed Clinical Models (DCMs) and the ISO11179 model in the Web Ontology Language (OWL); a combination of UML diagrams and Excel spreadsheets were used to extract the metamodels for fourteen HL7 DCM constructs. A critical limitation of this approach is that the transformation from metamodels to their ontological representa-
tion in OWL is based on a manual encoding. Leroux et al. (Leroux et al., 2012) use existing ontologies to enrich OpenClinica forms.

Model driven engineering techniques have also been applied to healthcare problems, Schlieter et al. (Schlieter et al., 2015) record their experience gained from using model-driven engineering to implement an application for path-based stroke care; amongst the lessons learned they recommend using existing ontological models where possible, and being prepared to reconcile a heterogeneity of models from the various stakeholders under a common metamodel. Atanasovski et al. (Atanasovski et al., 2018) presents a formal meta-model used to specify healthcare process management in an electronic health record system using FHIR and OpenEHR. Marcos et al. (Marcos et al., 2013) describe the implementation of an OpenEHR system to enable interoperability in clinical trial data using OpenEHR archetypes. Archetypes are part of a domain specific language (DSL) which in turn is part of the OpenEHR standard, discussed in more detail in the next section, but are described in detail in (Costa et al., 2011) The problems with integrating data encoded using different datasets and terminologies are clearly identified by Jian (Jian et al., 2007), and solutions using OpenEHR technology are put forward in (Martínez-Costa et al., 2010).

In the Model Driven Health Tools (MDHT) (Open-Health-Tools, 2008) project, the HL7 Clinical Document Architecture (CDA) standard (Dolin et al., 2006) for managing patient records is implemented using Eclipse UML tools (Eclipse-Foundation, 2018). MDHT supports only the CDA standard, whereas the Model Catalogue can interoperate with any metadata standard. The CDA standards are large and complex: Scott and Worden (Scott and Worden, 2012) advocate a model-driven approach to simplify the HL7 CDA, supported by three case studies: the NHS England ‘Interoperability Toolkit’, a simplification of US CDA documents, and the Common Assessment Framework project for health and care providers in England.

3 BACKGROUND

As mentioned earlier one goal of this work was to discover more efficient ways of integrating healthcare data from heterogeneous datasets. During the course of this work a number of standardised datasets were encountered, some being local standards, some National (such as the Cancer and Outcomes Dataset, managed by NHS England) and some International (such as SNOMED CT and OpenEHR).

The challenge here is to build a metadata registry that can easily store data specifications from all these varying datasets and standards, to store and make this information available for other applications which are able to transform the data from one dataset to another. This section is designed to give the reader a brief insight into the variability of these datasets.

These healthcare data standards have evolved over the last 30 years, mostly starting in particular communities focusing on specific healthcare sub-domains, such as pathology or laboratory testing. Some evolved to address local problems, such as defining standard formats to send laboratory results back to clinicians, and some started with the broader goals of unifying global healthcare records with one common standard. Each dataset has a slightly different way of looking at data, is used in a slightly different way reflecting the different sub-domains and cultures that the dataset evolved from. For instance LOINC is widely used in laboratories carrying out clinical testing. SNOMED CT aims at being a clinical terminology which has terms to be used in the whole of the healthcare domains, however in the sub-domain of laboratory testing LOINC has more granularity, and so there is a need for mapping between the two data standards.

The data standards that are used in the healthcare domain, having evolved from specific sub-domains, tend to use the language adopted by that sub-domain, and very often the rules of how meaningful sentences are constructed in the area of laboratory testing for instance, are not exactly the same rules as are used in general English. So while these areas look to provide terminologies, the use of these terminologies can differ from standard to standard.

In some standards, such as SNOMED CT terms are designated as pre and post co-ordinated. Pre-coordinated terms are terms that have compound terms combined in advance to arrive at a specific designation, with a specific identifier.

This is illustrated in Figure 1 using the normative example, taken from the ITSDSO website, showing the two different ways of expressing a fractured tibia. The pre-coordinated term, ID 31978002, expresses the idea of a "fractured tibia", whereas the post-coordinated terms of tibia, ID 12611008 and fracture, ID 125605004 are coordinated during the actual diagnosis, to form the resultant concept "fractured tibia" using the set of SNOMED-CT post-coordinated terms.

SNOMED-CT is at core maintained as a formal ontology using the Web Ontology Language (OWL), and thus the coordination can be carried out using description logic, as illustrated in (Stevens and Sattler, 2013), and (Rector and Iannone, 2012).
In most healthcare centres today, it is much more likely to be carried out by other patient record and clinical diagnosis tools as appropriate, and a mapping made from the tools’ own dataset to the SNOMED-CT standard. This throws light on another issue faced in the process of data collection from heterogeneous sources: that the data very often needs to be transformed, normally using standard ETL processes. Datasets that use post-coordinated terms lead to less flexibility in such transformations, since the terms used (in this case the post-coordinated Fracture of Tibia as opposed to the pre-coordinated Fracture and Bone Structure Tibia) may not all have direct analogues in the dataset definitions of the data sources. It is more likely, though not guaranteed, that a better partial mapping can be made from the pre-coordinated terms since there are going to be more of them than from the post-coordinated terms.

3.1 Healthcare Standard Datasets

The data standards being used in healthcare originate from different specialist areas, use different formats and have many overlaps, resulting in a number of different viewpoints over which standards are more or less useful. This section gives an overview and brief review of some of the main standards encountered, key considerations from a modelling point of view are hierarchy depth, granularity, restraints and constraints and data element relationships. Whilst the domain of Healthcare might appear to be a single domain to the outsider, it is split into a number of sub-domains, who in turn have evolved different process, different applications and even different uses of language to describe the work being carried out in the subdomain. These factors are reflected in the differences between different datasets, and all contribute to making interoperability a complex problem to solve.

3.1.1 SNOMED and Semantic Technologies

The systemized nomenclature of medicine (SNOMED) began life as the systemised nomenclature of pathology (SNOP) in 1965, originated by the College of Pathologists (CAP) in the US.

At core SNOMED CT is a set of terms, which are attached to formally defined concepts, and each is given a code. It enables different medical conditions to be given a unique reference, which due to the hierarchical nature of the core ontology allows clinicians to go into the appropriate amount of detail. SNOMED CT has a deep hierarchy of terms, and include pre and post-coordinated terms, as discussed.

3.1.2 HL7 and FHIR

Health Level 7 is a set of International standards pertaining to information usage and interoperability in healthcare, produced by Health Level 7 International and adopted by both the American National Standards Institute (ANSI) and the International Standards Organization (ISO). HL7 has been around for almost 30 years, and in essence it is a set of messaging standards, definitions and resources, including Version 2.x Messaging Standard, Version 3.x Messaging Standard, Clinical Document Architecture (CDA), Continuity of Care Document (CCD), Structured Product Labelling (SPL), Clinical Context Object Workshop (CCOW) and Fast Healthcare Interoperability Resources (FHIR).

3.1.3 Read Codes

Read codes are the standard clinical terminology system developed by Dr James Read in the 1980’s for use in the United Kingdom. There are 2 forms of read codes, a 4-byte version (the original version), a 5-byte version 2, and version 3 also called Clinical Terms 3 or CTV3. This latter vocabulary was merged with the US origin SNOMED-RT to form SNOMED-CT. CTV3 was mandated for use across the NHS in 1999, it currently consists of a vocabulary of about 200,00 data terms, together with tables that capture various hierarchies of elements.

3.1.4 International Classification of Diseases (ICD)

The International Classification of Diseases (ICD) is managed by the World Health Organization (WHO),
the authority for health matters in the United Nations.[2] The ICD is a health care classification system, its primary purpose is to provide a set of diagnostic codes for diseases and disease classification. This includes nuanced classifications of a wide variety of items such as signs, symptoms, abnormal findings, complaints, social circumstances, and external causes of injury or disease.

3.1.5 OpenEHR - ISO EN 13606

OpenEHR is an open standard for electronic health records, which is defined by the international standard ISO13606 (ISOTC215, 2008). From the perspective of data management the standard provides a reference model and a number of archetypes, which describe datasets from a clinical perspective and templates, which provide instantiations of those archetypes for specific use cases. The standard covers not only an architecture definition, but also a domain specific language for storing, managing and querying healthcare records.

3.1.6 OMOP CDM

The Observational Medical Outcomes Partnership (OMOP) Common Data Model (CDM) came out of work started in 2008 by a public-private partnership in the USA, between the FDA and the pharmaceutical industry to research the utility of using existing health care databases to evaluate safety issues relating to currently approved and available drugs.

3.1.7 LOINC

The Logical Observation Identifiers Names and Codes (LOINC), was started in 1994 with goal of codifying laboratory specimens and tests in a clinical context. It is designed as a data standard for laboratory tests and results, anything that can be observed or tested in relation to a patient’s healthcare. Part of LOINC can be used for normal observations, and part of it for laboratory results. It is at core a large terminology, more detailed in many areas of pathology than SNOMED CT. It is very often used in conjunction with both SNOMED CT and HL7, so that messages sent requested laboratory tests may be codified in HL7 using LOINC, and responses may use SNOMED in addition to LOINC.

3.1.8 NHS Data Dictionary

The NHS Data Dictionary is maintained by NHS Digital as a UML model, however it is publicly available as either HTML files, viewable on a website, or as XML/XSD files which can be downloaded. The model is relatively straightforward, and is used for most of the reporting carried out by NHS Digital in the UK.

4 ISO11179: ISO STANDARD FOR METADATA REGISTRIES

Metadata can be recorded using a metadata registry; a metadata registry (MDR) is a toolkit which allows definitions of datasets to be stored, curated and managed. Metadata is usually defined as data about data; this is an unfortunate definition, in that it can be interpreted in a variety of ways. Our work relates to data management and for most data management purposes it is normally taken to mean data which defines the structure of data, although data about the governance, provenance and other aspects of that data can also be relevant, and thus included in a metadata registry. By storing the definitions of every data element and also the relationships between data elements in a metadata registry a map of all the data elements and data flows in an organization or domain can be created. Such a map can be used to manage, understand and curate the datasets being used. The ISO standard 11179 is a standard for metadata registries, although this designation is extended in the body of the standard. The ISO11179 (ISOJTC1, 2015) standard, was issued in 2015, is composed of 6 parts:

- ISO/IEC 11179-1:2015 Framework (referred to as ISO/IEC 11179-1)
- ISO/IEC 11179-2:2005 Classification
- ISO/IEC 11179-3:2013 Registry metamodel and basic attributes
- ISO/IEC 11179-4:2004 Formulation of data definitions
- ISO/IEC 11179-5:2015 Naming and identification principles
- ISO/IEC 11179-6:2015 Registration

The standard is comprehensive, however we are only considering some key aspects of Part 3 in this research. Part 7 called Datasets, is planned and is currently under review. This introduces a metamodel for datasets into the standard, however this was not readily available at the time that this research was carried out, and is not considered here. To quote from the standard itself:

ISO/IEC 11179 addresses the semantics of data, the representation of data and the registration of the descriptions of that data. It is
through these descriptions that an accurate understanding of the semantics and a useful depiction of the data are found.

The purpose of ISO/IEC 11179 is to promote the following:

- standard description of data
- common understanding of data across organizational elements and between organizations
- re-use and standardization of data over time, space, and applications
- harmonization and standardization of data within an organization and across organizations
- management of the components of descriptions of data
- re-use of the components of descriptions of data

Part 3 of the standard provides a registry metamodel, specified using UML diagrams, and it was thus chosen as the starting point for building this implementation. There is a warning at the beginning of Part 3, stating that this part prescribes a conceptual model, not a physical implementation. This part of ISO/IEC 11179 also prescribes a list of basic attributes (see clause 12) for situations where a full conceptual model is not required or not appropriate. The other 5 parts were used to inform the core metadata registry metamodel as specified in ISO11179: Part 3.

4.0.1 Part 3 - Registry Metamodel and Basic Attributes

The objectives of the metadata registry metamodel are defined, in the standard, as:

- Providing a unified view of the concepts, terms, value domains and value meanings
- Promoting a common understanding of the data described
- Providing the specification at a conceptual level to facilitate the sharing and reuse of the contents of the implementations

The standard continues to split the metamodel up into 6 packages, Basic, Registration, Concepts, Binary relations, Data description and Identification, Designation and Definition.

In section 4 it is pointed out that clauses 7-9 are needed to implement a Concept Systems Registry, clause 10 will allow the implementation of an Extended Concept Systems Registry, and clause 11 specifies a metadata registry, whereas an extended metadata registry will implement all clauses 7-11. Our initial scope was to implement an extended metadata registry as described in clauses 7-11.

4.0.2 Data Description Package

This package specifies a metamodel for handling data, and although it references other packages which are mostly dealing with the more administrative aspects of registering metadata, it primarily puts forward a conceptual metamodel for handling data. Hence for the purposes of data interoperability it is the most relevant part of the standard.

The core model given for data description in ISO11179 is that reproduced in Figure 2, it shows the linkage between a Data Element, a Value Domain, a Conceptual Domain and a Data Element Concept. The area above the dotted red line is defined as the semantic or conceptual level, whereas the area below the red dotted line is defined as the representational level. The assumption is that the Data Element and Value Domain are objects which are being registered and classified, as per the processes defined in other parts of the standard.

This arrangement can be illustrated by the idea of a visit to the doctor, we can define a concept called reason for visit to healthcare centre and call this a data element concept, and from this we would implement a data element called reason for attendance and perhaps represent that with a set of enumerated codes, each representing a different reason. In ISO11179 structuring we would split the data element concept into an object class: Person, and a property: Reason for clinic attendance.


Part 3 of the standard, titled Registry Metamodel and basic attributes, and revised in 2013, was taken as the reference point for building the Metadata Registry implementation. In reviewing the standard several problems came to light, especially with regard to implementation, and especially with regard to any kind of conformity. These are listed as:

- ISO11179 introduces representational items, such as Conceptual Domain, Data Element, Data El-
element Concept, etc., indicating that they are part of the standard or ideal metadata registry metamodel with no indication of how the ISO11179-compliant models so defined are generated, used or related, nor how data can be transformed into this particular model or what the actual advantage is over any other metamodel/model.

- Partial UML models are specified at different levels of granularity, with no explicit connection to show how they relate.
- The text does not allow an overall model to be built with any degree of certainty of whether or not it conforms to the standard.
- Basic types used in the metamodel include types which in most computer science contexts would be viewed as derived types, this makes implementation needlessly difficult and confusing.
- The introduction asserts that metadata registry is specified in the form of a conceptual data model, however, despite references to other standards, and a brief explanation in Appendix E, no definitive explanation of what is meant by conceptual data model is provided.
- The examples provided are very concept specific, for instance, the example of country codes works for concepts which are used as a list, but many concepts used in healthcare are not used in such a straightforward fashion.
- Many concepts and terms are described, some are specified, and some specifications overlap with other definitions; for instance value domains are specified with the same definition that is used to describe data types.
- The UML models provide a great deal of detail for each sub-section of a metadata registry, however no system diagram or clear description is provided, it is therefore impossible to build a working system based on the UML diagrams alone, considerable interpretation is required, which detracts from the specification provided.

The standard is declared as being a standard for metadata registries, and although it contains a lot of disparate ideas on the subject of interoperability which can be applied to a metadata registry, no core set of definitions, core language or metamodel was found that could be used as a measure for conformance. Standards by definition should be conformed to, and whilst conformance is mentioned, it is lacking a clear set of definitions which can be used as a measurement for conformity. Therefore the goal of building an ISO11179 conformant metadata registry was abandoned early on, when it became apparent that clear objective conformance criteria were not present in the standard, despite the subject of conformance being discussed. That said, there is much in the standard document which can be usefully incorporated into the design of a metadata registry, and development continued with a view to include those aspects of the standards which could be shown to be beneficial to the construction of a metadata registry.

5 IMPLEMENTATION

In the initial design work, the UML diagrams contained in Part 3 section 11 was taken to be the basic metamodel around which the core metadata registry would be built.

As detailed here, this very quickly became unworkable, mostly as a result of user’s being unable to translate data structures into the form dictated by the ISO11179 metamodel.

5.1 Implementation of ISO11179 UML Metamodel

Initially work began by implementing the UML models as specified in the standard (part 3, section 11), however when the initial prototype was run, many pieces of information were identified as being loaded more than once. Due to the model provided, there is an overlap of the conceptual structure of the metamodel, and the logical structure of the metamodel, although no such reference is available in the standard itself. Initially a basic domain model was built, using a Grails 2.4.3 toolkit, using the following basic representational items shown in the first column of Table 2.

The work was then shown to analysts and clinicians experienced in building healthcare datasets, with a view to having them enter suitable datasets and then take part in developing the data set curation functionality around the ISO11179 conformant metamodel.

5.2 Concerns over ISO11179 Conformance

The first major problem was in specifying exactly what metadata should be input into the prototype metadata registry, in particular how to translate or transform existing models or meta-models into the set of constructs defined and discussed in the standard. To illustrate this issue, consider taking a data item from an existing medical dataset, in this case COSD, as shown in Figure 3.
Using ISO11179 we take the date of clinical assessment as a data element, however it would be specified with the patient details, since conceptually one would need to describe the context of the data item, this results in a data element which has an object property of patient as an integral part of the construct. This is illustrated in Table 1

<table>
<thead>
<tr>
<th>ISO Artefact</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data Element</td>
<td>Patient and Date of clinical Assessment in form ccyy-mm-dd</td>
</tr>
<tr>
<td>Data Element Concept</td>
<td>Patient and Date of clinical Assessment</td>
</tr>
<tr>
<td>Value Domain</td>
<td>Date in form ccyy-mm-dd</td>
</tr>
<tr>
<td>Object Class</td>
<td>Patient</td>
</tr>
<tr>
<td>Property</td>
<td>Diagnosis Date</td>
</tr>
</tbody>
</table>

However if one is trying to enter the metadata for the Date of Clinical Assessment shown in the first row of Figure 3, there is an immediate disparity in that no object class is specified, available or immediately obvious from the spreadsheet. There is mention of the outpatient or patient in the description, however it is not in the view of the analyst preparing the dataset of any significance, and therefore was not included in a separate column. Therefore there is immediately a dilemma, do we enter outpatient, patient or simply leave the object property blank? The next issue is the idea of having the data element, if we drop the Patient from the name and then call the data element Date of Clinical Assessment, what then is the Data Element Concept? and how is it different from the description of the Data Element? Should the Data Element Concept include the patient? There is no obvious answer, and from a user perspective it appears that a simple set of dataset metadata, i.e. column headings on a spreadsheet, are being transformed into something more complex in order to manage them, but that management can only be carried out by experts versed in ISO/IEC11179.

5.2.1 User Difficulties
In the first few weeks of trying to enter standard existing healthcare datasets into the prototype metadata registry many objections were encountered from users experience with existing healthcare datasets, of the kind documented in the previous section. Metadata was seen to be entered twice or three times needlessly, the difference between the description of a Data Element and a Data Element Concept was not understood. Likewise the difference between a Value Domain and a Data Type whilst apparent in theory, was in practice not apparent, since the models being generated were not implementation specific. Therefore a representation of a set of numerical values would in nearly all cases be represented by the same data type, for instance the date of clinical assessment would have a value domain of date and a data type of date, which would then be implemented in a particular system as appropriate, e.g. text string, org.joda.time.format.DateTimeFormat as appropriate by the system concerned.

5.3 Update to Meta-model
After a few weeks it was decided to update the meta-model, at first, the number of ISO11179 elements was reduced, however this still didn’t gain any traction with users, who found the system confusing, counter-intuitive, time-consuming and needing a lot of extra work to understand the new language constructs in-
roduced by the standard. As a result the development went through a two iterations to arrive at the current model, the initial prototype we refer to as version 0.x, the second as 1.x and the third as 2.x. The third, has been fairly successful as is being used at over 7 hospital trusts and healthcare research centres in the UK currently. The changes in core constructs are shown in Table 2.

<table>
<thead>
<tr>
<th>ISO Artefact (v.0.x)</th>
<th>Iteration 2 (v.2.x)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Described Conceptual Domain</td>
<td>-</td>
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<tr>
<td>Object Class</td>
<td>DataClass</td>
</tr>
<tr>
<td>Property</td>
<td>-</td>
</tr>
<tr>
<td>Data Element Concept</td>
<td>-</td>
</tr>
<tr>
<td>Data Element</td>
<td>DataElement</td>
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<tr>
<td>Data Type</td>
<td>DataType</td>
</tr>
<tr>
<td>Value Domain</td>
<td>-</td>
</tr>
<tr>
<td>Described Value Domain</td>
<td>-</td>
</tr>
<tr>
<td>Enumerated Value Domain</td>
<td>EnumeratedType</td>
</tr>
<tr>
<td>Permissible Value</td>
<td>-</td>
</tr>
<tr>
<td>Enumerated Conceptual Domain</td>
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<tr>
<td>Relations</td>
<td>Relationship</td>
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<td>RelationshipMetadata</td>
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<tr>
<td>-</td>
<td>RelationshipType</td>
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<tr>
<td>Classification</td>
<td>-</td>
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<td>DataModelPolicy</td>
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<td>PrimitiveType</td>
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<td>-</td>
<td>ReferenceType</td>
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<td>Tag</td>
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<td>ValidationRule</td>
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</tbody>
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The first iteration resulted in version 1.x of the metadata registry, which went into service in Genomics England in 2015, however it met with many criticisms from users, and a complete overhaul was undertaken. This time a different approach was used, and the basic metamodel redeveloped, informed more by feedback from data analysts than reliance on the ISO/IEC11179 standard. The domain metamodel was developed using XText, which allowed for the fast iterative development of the metamodel, using the Eclipse toolkit. Once this was established the domain model was implemented using the Grails framework (v2.5.6), which allowed much of the existing codebase to be re-used.

5.3.1 MDML: A DSL for Metadata Management

Revising the core metamodel required examining a wide range of existing healthcare datasets, such as the ones mentioned in the introduction, and reviewing the conceptual background, language, context and purposes to which they were being used. Various modelling methodologies were examined, however the ECore/XText framework provided a relatively straightforward way of testing ideas quickly. A grammar was developed using XText which was called the Metadata Management Language (MDML), and from this the most recent metadata registry was developed.

When building this there were a number of features or requirements, that the Healthcare professions we dealt with required. These were absent in the ISO/IEC11179 standard, and are listed below:

- The ability to uniquely identify particular data elements, groupings, and identify a publishing state, i.e. is the dataset a draft, a current dataset, a superseded (but still used) dataset.

Some of these issues are touched upon in the standard, for instance data elements can be grouped using the construct of object property, however there is no notion of layers of grouping. With SNOMED CT there are 13 layers of hierarchy, with links between layers, and this kind of structure needs to be modelled and managed with the metadata registry.

The XText domain model included all the constructs from column 3 of Table 2, and provided a grammar which was then used to generate a set of domain objects, which in turn were able to represent the various healthcare datasets being used. The core enabled the automatic generation of grails domain classes using the code generation capabilities of XText. For illustrative purposes we show the key metadata registry structure in Figure 4.
The structure illustrated here allows the metadata registry to capture data elements, which is at the core of the standard. It also allows each element to have a status, which allows it to conform to the publishing aspects of the standard. By making each item in the registry a class inheriting from an abstract Catalogue Element each data element is captured by a unique identifier and a status relating to its publishing status. In the ISO standard metamodel the data element is the primary artefact under consideration, however there is no obvious direction on how groups of data elements should be handled.

In practice data elements appear in groups, whether spreadsheets, database tables or CSV files. In the data analysis business, groups of data need to be brought together and to do this datasets need to be mapped and transformed. In our first and second models it was easy to generated lists of data elements, however they were nearly always grouped using several layers of hierarchy into different entities, which made mapping and transformation difficult.

Further a DataModel is able to add in any number of extensions, which can be used to augment particular data elements for instance to mark them as user interface items so that user interfaces can be automatically generated. In addition imports can be added from other DataModels, so that new datasets can be created using data elements from several different datasets. This metamodel evolved over a period time in order to solve the problems of integrating data from a wide variety of heterogeneous sources, it is closely based around UML/Ecore, but is in essence a domain specific language or metamodel for data integration.

### 5.3.2 Core Elements of MDML

The main constructs of DataClasses, DataElements and DataTypes are common to most technical spaces, and can be used to group and manage data elements in a way that is easy for data analysts and developers to grasp. Metadata can then be attached to any of these core elements, so that security classifications, and governance directives, found in other parts of the ISO11179 standard can easily be added to particular DataElements or DataClasses.

At core MDML provides a metamodel, in essence a language for data structures, which allows additional metadata to be added on through a simple mechanism of extension values and tags. It allows datasets to be grouped and classified in a flexible and manageable way, so that one registry could hold several common healthcare metamodels, say FHIR, OMOP and LOINC, and build local models based on all three. There is a clear identification system built into the language which means that each version is separately identified, so that any mappings between FHIR and OMOP for instance will be detailed to particular versions of the datasets.

There is a clear publishing cycle built into the metamodel, which allows change for draft models, but not for finalized models, therefore data elements in finalized models can be imported into other (draft) datamodels, but no new elements can be imported in currently finalized models. This is an important management feature; derived from the ISO standard which has been built into the language.

### 6 RESULTS

By using the metadata registry Genomics England was able to specify exactly what data they required from UK Hospital Trusts by building their own data models centrally online. The other organizations are then able to use these models to verify that outbound data is valid, and thus ensuring that data won’t be returned.

The main technique used is to import a new dataset into the metadata registry, normally in csv or excel format, to configure a mapping using XML, and then import the dataset definition into the metadata registry directly. From this new model a number of artefacts such as spreadsheets and XML Schema files could be generated from one source. This eliminated any doubt arising from spreadsheets which were disseminated after meetings. Instead the meetings could be held online, and the changes were immediately visible to all parties. Spreadsheets and XML files could
then be generated for local form or code generation, and rules could be attached to data elements and data types which could be automatically downloaded via a REST interface, rather than by disseminating spreadsheets and documents.

The MDML based metadata registry has been in use for the past 2 years at Genomics England, as well as at 7 other NHS Hospital Trusts in London. It’s core use is to manage datasets which are used by Genomics England to specify dataset requirements, a dataset or model is defined in the metadata registry, this is then used to automatically generate output specifications as excel, XML or as Case Report Forms. The latter is a specification for generating forms in the Open Clinica system, but which can automatically be loaded into Open Clinica to generate both the forms and data repository to store the data. Rules, embedded into the DataTypes as regular expressions, or as groovy code, are then used to validate datasets.

Data curation, which previously was carried out using excel spreadsheets, and involved regular physical meetings between data managers and clinical leads from all over the UK now takes place online. The models which are generated are given a unique identifier, and once all parties agree to issue a new finalized model it is finalized thus preventing further change to that version of the model. At any point in time the model, and it’s history can be referred to online, removing doubt and ambiguity over what the constituents are, further the model can be accessed using a REST interface, making it available for code generation utilities remotely. The results of using the metadata registry and toolkit are listed below:

- Reduction in time and effort in creating and curating datasets over spreadsheets and document-based techniques.
- Ability to clearly access the current version of a dataset and verify that it is the correct version, using unique identifiers.
- Ability to generate artefacts based on the data model, such as forms, XML and XML Schemas automatically.
- Ability to validate data against the model automatically using rules stored against individual data elements.
- Ability to map between different datasets in a detailed and precise fashion, again with the use of unique identifiers.

7 DISCUSSION

By applying MDE principles a metadata registry was built that satisfied the requirements of this research organisation for managing heterogeneous dataset. At the beginning of the work, much time and effort was put into implementing the international standard for metadata registries which is either lacking in clarity, or un-informed by the healthcare use cases that were encountered in this research effort. Whilst much in the standard was informative and relevant to the more general issue of data integration, the clear lack of a consistent metamodel or language around which to build a metadata registry was a great disappointment. Metadata registries are becoming commonplace in enterprise architecture today, especially since automatic management of large datasets, big data, is extremely difficult without them. So it was very disappointing that applying the ISO/IEC11179 standard proved so difficult in this instance. It must be pointed out that the shortcomings found and described in this paper only relate to 2 or 3 clauses in a standard which is several hundred pages in length, however they did cause significant problems in our development effort and it is hoped that clarifications and improvements to the standard will be made in due course.

8 CONCLUSION

There were a number of problems which arose from attempting to build a metadata registry which complies with the ISO/IEC11179 standard in the early stages, these have been documented, and steps taken to correct the problems encountered. In the course of correcting these problems a model driven engineering approach was taken, and an effective set of tools were built based around a revised metamodel for metadata management. The open source toolkit (MetadataWorks, 2018) developed from this research, has proved very effective, being used to simplify and speed up the work of data curation, data-wrangling and data-cleaning. In addition an MDE-based toolkit was built which provided a REST interface and allowed third parties to carry out automated data validation. Future work on extending the model is anticipated, in particular to produce automated tool chains which can generate both the input forms, repositories and mapping capabilities for heterogeneous datasets.
ACKNOWLEDGEMENTS

I would like to acknowledge the help of Adam Milward, Kathy Farndon, Amanda O’Neill and Samuel Hubble at Genomics England, and Jim Davies, Charles Crichton, Steve Harris and James Welch at the University of Oxford.

REFERENCES


