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Del 12.3
Second Report on core set of FAIR software tools and on extended set of unified FAIR data standards applied in EJP RD

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# Table of content

1. Scope ................................................................................................................................. 6  
2. Sources ................................................................................................................................. 6  
3. Development status ............................................................................................................... 8  
4. Format ................................................................................................................................ 8  
5. Used standards and tools ..................................................................................................... 9  
   5.1. Standards ......................................................................................................................... 9  
      5.1.1. Meta-data standards ................................................................................................... 9  
         5.1.1.1. DCAT - Data Catalog Vocabulary (DCAT) ................................................................. 9  
         5.1.1.2. Dublin Core Metadata Terms and Element Set (“Dublin Core terms’) ................. 9  
         5.1.1.3. EJP RD Metadata Model ...................................................................................... 9  
         5.1.1.4. FASTA, FASTQ .................................................................................................... 10  
         5.1.1.5. ISO 21838 ............................................................................................................. 10  
         5.1.1.6. ISO 23903:2021 Health informatics - Interoperability and integration reference architecture - Model and framework ................................................................. 10  
         5.1.1.7. Maelstrom Data harmonization guidelines ............................................................. 11  
         5.1.1.8. MIABIS 2.0 ............................................................................................................ 11  
         5.1.1.9. Phenopackets schema .......................................................................................... 11  
         5.1.1.10. RD3 (Rare Disease Data about Data) data model .............................................. 12  
   5.1.2. Standards on data file formats, markup, and annotation ............................................. 12  
      5.1.2.1. BED, GFF .............................................................................................................. 12  
      5.1.2.2. CSV ...................................................................................................................... 12  
      5.1.2.3. ISA-Tab+serialisations (ISA-JSOn etc) ................................................................. 12  
      5.1.2.4. JSON LD .............................................................................................................. 13  
      5.1.2.5. OML ..................................................................................................................... 13  
      5.1.2.6. OpenAPI Specification ......................................................................................... 13  
      5.1.2.7. Schema.org .......................................................................................................... 13  
      5.1.2.8. XML ..................................................................................................................... 13  
   5.1.3. Standard data element sets ........................................................................................... 14  
      5.1.3.1. CCE - Common Condition of use Elements ......................................................... 14  
      5.1.3.2. CDE - Common Data Elements .......................................................................... 14  
   5.1.4. Standards on data models ........................................................................................... 14  
      5.1.4.1. Basic Formal Ontology (BFO) and The Open Biological and Biomedical Ontology (OBO) Foundry ontologies .............................................................................. 14  
      5.1.4.2. Clinical Data Interchange Standards Consortium (CDISC) ................................. 15  
      5.1.4.3. bioCADDIE Data Tag Suite (DATS) ....................................................................... 15  
      5.1.4.4. DUC – Digital Use Conditions ........................................................................... 15  
      5.1.4.5. JRC Common Data Element Semantic Data Model ........................................... 15
5.1.5. Standards on data ontology, terminology and vocabulary ..................17
5.1.5.1. Anatomical Therapeutic Chemical (ATC) ......................................17
5.1.5.2. Data Catalogue Vocabulary (DCAT) .............................................17
5.1.5.3. Data Use Ontology (DUO) ..........................................................17
5.1.5.4. FAIRplus recipe on how to choose controlled vocabulary .................18
5.1.5.5. Gene Ontology (GO) .................................................................18
5.1.5.6. GENO ..................................................................................18
5.1.5.7. HGNC (HUGO) .......................................................................18
5.1.5.8. Sequence Variant Nomenclature (HGVS) .......................................18
5.1.5.9. HPO-ORDO ontological module (HOOM) .....................................19
5.1.5.10. Human Phenotype Ontology (HPO) ..............................................19
5.1.5.11. International Classification of Diseases (ICD) ..............................19
5.1.5.12. International Classification of Functioning, Disability and Health (ICF) 19
5.1.5.13. International Classification of Diseases for Oncology (ICD-O) .......20
5.1.5.14. Informed Consent Ontology (ICO) .................................................20
5.1.5.15. LOINC ..................................................................................20
5.1.5.16. MedDRA - Medical Dictionary for Regulatory Activities Terminology ..20
5.1.5.17. MeSH - Medical Subject Headings ...............................................20
5.1.5.18. NCIT - National Cancer Institute Thesaurus ................................21
5.1.5.19. Online Mendelian Inheritance in Man (OMIM) ............................21
5.1.5.20. Orphanet nomenclature of rare diseases (ORPHAcodes) and Orphanet ontology of rare diseases (ORDO) ................................21
5.1.5.21. PROV Ontology ......................................................................21
5.1.5.22. Semanticscience Integrated Ontology (SIO) ..................................22
5.1.5.23. SNOMED CT ..........................................................................22
5.1.6. Standards on data discovery ...............................................................22
5.1.6.1. Automatable Data Discovery and Access Matrix (ADA-M) ..............22
5.1.6.2. Beacon-1 .................................................................................22
5.1.6.3. Beacon-2 API .........................................................................23
5.1.6.4. Bioschemas .............................................................................23
5.1.7. Standards on data exchange mechanisms ...........................................23
5.1.7.1. DOIP - Digital Object Interface Protocol ......................................23
5.1.7.2. FHIR - Fast Healthcare Interoperability Resources ..........................23
5.1.7.3. HL7 FHIR4FAIR – FHIR Implementation Guide ............................24
5.1.7.4. ISO /AWI TR 24305 Health informatics - Guidelines for implementation of HL7/FHIR based on ISO 13940 and ISO 13606 ..........................24
5.1.7.5. mzML/mzIdentML ..................................................................24
5.1.7.6. PhenoPackets ............................................................................................................. 24
5.1.7.7. REST ....................................................................................................................... 25
5.1.8. Standards on security, authentication, and authorisation ........................................ 25
5.1.8.1. GA4GH Passport .................................................................................................... 25
5.1.8.2. GA4GH Visa .......................................................................................................... 25
5.1.8.3. Open ID Connect .................................................................................................... 25
5.1.8.4. SAML 2.0 ............................................................................................................... 26
5.2. Tools ................................................................................................................................ 27
5.2.1.1. FAIR Genomes – ELSI Framework ..................................................................... 27
5.2.1.2. FAIR Genomes – Metadata scheme .................................................................... 27
5.2.1.3. FAIR Genomes – Metadata codebook ............................................................... 28
5.2.1.4. FAIR Genomes – VARDA .................................................................................... 28
5.2.1.5. FAIR Genomes – Mutalyzer ................................................................................ 28
5.2.2. Authentication and Authorization Infrastructure ..................................................... 29
5.2.2.1. LifeScience AAI .................................................................................................... 29
5.2.3. Pseudonymisation ..................................................................................................... 29
5.2.3.1. EUPID .................................................................................................................. 29
5.2.3.2. SPIDER ................................................................................................................ 29
5.2.4. Mapping / Alignment Services ................................................................................ 30
5.2.4.1. Data Model Alignment Service .......................................................................... 30
5.2.4.2. OxO ....................................................................................................................... 30
5.2.4.3. The FAIR Evaluator .............................................................................................. 30
5.2.5. Consent and use conditions ...................................................................................... 31
5.2.5.1. CCE / DUC Profile Creation Tool [ULEIC] ......................................................... 31
5.2.5.2. CCE / DUC Profile Creation Tool [UMCG] ........................................................ 31
5.2.6. Record linkage services ............................................................................................ 31
5.2.6.1. EUPID .................................................................................................................. 31
5.2.6.2. SPIDER ................................................................................................................ 31
5.2.7. Resource/Data/Sample discovery and access ............................................................. 31
5.2.7.1. Beacon in a box ...................................................................................................... 32
5.2.7.2. bio.tools ............................................................................................................... 32
5.2.7.3. Castor .................................................................................................................... 32
5.2.7.4. CDE in a box/FAIR in a box ................................................................................. 32
5.2.7.5. Combined RD-Nexus + CDE/FAIR-in-a-Box ....................................................... 33
5.2.7.6. FAIR Data Point (FDP) ......................................................................................... 33
5.2.7.7. FDP reference implementation ........................................................................... 34
5.2.7.8. GRLC ..................................................................................................................... 34
5.2.7.9. I2b2 / tranSMART ............................................................................................... 34
5.2.7.10. INFRAFRONTIER ........................................................................................................35
5.2.7.11. MOLGENIS ........................................................................................................35
5.2.7.12. REDCap ...............................................................................................................35
5.2.7.13. RD-Connect GPAP ..............................................................................................36
5.2.7.14. Rare Disease Networked Exploration of the UnSeen (RD-NEXUS) ..............36
1. Scope

This report intends to provide an overview on the status of software tools and standards relevant to the European Joint Programme on Rare Diseases (EJP RD). Specifically, it lists those elements which have been identified in preceding work as being relevant to the Virtual Platform (VP).

This collection of items will be designated as “the list” in the following. The relevance to the VP of items in the list stems either from the VP design and implementation work or from the interaction with potential users of VP components and their existing or upcoming systems, respectively.

For the purpose of this document, “tools” are defined as all elements of the VP as described in the Virtual Platform Specification (VIPS)\(^1\), except for standards and data sources.

This document is the second version which is based on Deliverable 12.2 “First report on core set of FAIR software tools & on extended set of unified FAIR data standards applied in EJP RD” and will be completed by the Deliverable 12.4, “Report on extended set of FAIR software tools, applied in EJP RD, including overview of FAIRification guidelines for RD data managers”, due in month 60.

Therefore, the list is expected to grow and the classification of items regarding their types and VP adoption status is expected to be updated/changed in subsequent releases.

2. Sources

This report builds on numerous preceding activities in general and the following documents and sources in particular.

**Del 12.1 “Report on core set of unified FAIR data standards”**
This deliverable was concerned with standards relevant to the VP. It identified: (a) existing standards that can be used/piloted directly; and (b) standards with a need to aggregate and/or map between and/or extend existing standards before they can be used in the VP context. Del 12.1 provided a very comprehensive list of standards, potentially relevant to the VP.

**The list of elements compiled by the FAIRification Work Focus**
This list has been compiled by the FAIRification team while interacting with numerous stakeholders in EJP RD, and in particular the European Reference Networks (ERNs).

**Deliverable 12.2 “First report on core set of FAIR software tools & on extended set of unified FAIR data standards applied in EJP RD”**
Del. 12.2 was based on Del. 12.1 and the list of elements compiled by the FAIRification Work Focus. Del 12.2 is the base for Del. 12.3. “

**AD36 Additional Deliverable Virtual Platform Specification (VIPS)**

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The Virtual Platform (VP) provides RD stakeholders with resources relevant to RD research. The VIPS depicts the overall structure of the VP (architecture, functional and non-functional requirements), the use cases supported by the VP, the process how to link resources to the VP and the agreed guidelines and standards. Its level of detail aims at the managerial level, not at the implementation level. The VIPS provides the architectural framework for the VP and, as such, it defines a classification scheme for VP components and their underlying software/service developments, which have been adopted for the present report as well. In 2021, a first version of the VIPS was developed, which served as a source for the present deliverable.

AD49 Additional Deliverable Second Version – Virtual Platform Specification (VIPS)
In parallel to the present deliverable, a second version of the VIPS has been developed, which reflected the outcomes of our experiences since finalizing the VIPS version 1.

Beyond 1 Million Genome / Deliverable 3.7 – Documented best practices in sharing and linking phenotypic and genetic data
The Beyond 1 Million Genomes (B1MG, https://b1mg-project.eu) project aims to make it easier to share human health data around Europe. It will support the European Union’s 1+ Million Genomes Initiative (1+MG), which aims to provide access to at least one million sequenced genomes in the EU by 2022. The B1MG project will support this initiative by creating the infrastructure, the legal guidance and the best practices to enable this access. It will make it possible for scientists and clinicians to study the genotypic and phenotypic data from over one million people. This data will be linked, so the genetic data from one individual can be matched with their phenotypic data (like their weight, blood group and medical history).

But the project will look ‘beyond’ the 1+MG Initiative and drive the development of a data sharing infrastructure that goes beyond the lifetime of 1+MG, and beyond 1 million genomes. Their deliverable 3.7 is the first version of documented best practices in sharing and linking phenotypic and genetic data. It identifies and describes best practices on sharing and linking phenotypic and genetic data in both the healthcare sector and in the research setting. The idea is to, as much as possible, avoid reinventing the wheel, learn from previous/current existing projects to improve performance and avoid mistakes made by other.

Results from B1MG Deliverable 3.7 have been analysed and some of their identified best practices have been included in the present deliverable.

The deliverable is available at https://zenodo.org/record/4819149#.Y4hiVRSZP-h
3. Development status

The Status of software tools/services/standards refers to its state of usage with respect to the VP development and has been aligned to table 6 of the VIPS.

<table>
<thead>
<tr>
<th>Status</th>
<th>Short Term</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Draft</td>
<td>The component being described has not been implemented yet or is under active development. However, it has not been shared or evaluated with stakeholders and end-users of the VP.</td>
</tr>
<tr>
<td>2</td>
<td>Trial use</td>
<td>The component being described is under active development and ready for trial use during the use case evaluation phase. However, no guarantees are made that future version remain compatible with older versions of the component.</td>
</tr>
<tr>
<td>3</td>
<td>Normative</td>
<td>The component described is in a stable state, can be used by clients and has guaranteed life cycle management process for future updates.</td>
</tr>
<tr>
<td>4</td>
<td>Deprecated</td>
<td>The component described is considered as being deprecated and might be removed from the document in the near future. For instance, a prototypic component developed in an iteration cycle might be dropped in favour for a more advanced component.</td>
</tr>
</tbody>
</table>

4. Format

FAIR standards and tools for data stewardship are presented for the RD community, particularly data stewards, in the following ways: (i) the ELIXIR RDMtoolkit presents basic guidance and a subset of tools via a web interface, (ii) the ELIXIR data stewardship wizard that smartly leads stewards to find appropriate standards and tools via a carefully crafted questionnaire, (iii) FAIR ‘cookbooks’ provide a third format to present FAIR tools and standards. Stewards thus have multiple sustainable ways to find tools and standards. A FAIR maturity indicator tool has been successfully tested on a number of RD resources in collaboration with ELIXIR and can be used as an additional aid in achieving FAIR compliance.

Providing a formal FAIR maturity status for standards and tools is outside of the scope of the EJP RD project. Here, the main intended use is providing stewards an aid in delivering resources that are compatible with a FAIR-based VP. The status in this document reflects the intended use in FAIRification as indicated by the submitter of the tool or standard.

Continuous updating and refinement of the list is foreseen to be a rolling task and the knowledge required to compile and maintain this list requires the full breadth and depth of the EJP RD community expertise.
5. Used standards and tools

5.1. Standards

5.1.1. Meta-data standards

5.1.1.1. DCAT - Data Catalog Vocabulary (DCAT)

DCAT is a W3C-recommended vocabulary, defined in RDF, which is designed to facilitate interoperability between data catalogues published on the Web. The version that is currently used in the EJP RD is DCAT version 2.

- Status of development: Normative
- Further Information: https://www.w3.org/TR/vocab-dcat-2/

5.1.1.2. Dublin Core Metadata Terms and Element Set (‘Dublin Core terms’)

The Dublin Core™ Metadata Terms² are widely used terms to denote metadata elements. They are maintained by the Dublin Core™ Metadata Initiative (DCMI)³. Included are the fifteen terms of the Dublin Core™ Metadata Element Set (also known as “the Dublin Core”⁴ plus several dozen properties, classes, datatypes, and vocabulary encoding schemes. The “Dublin Core” plus these extension vocabularies are collectively referred to as “DCMI metadata terms” (“Dublin Core terms” for short). They are expressed in RDF vocabularies for use in Linked Data. Creators of non-RDF metadata, such as in XML, JSON, UML, or relational databases, can use the terms by disregarding both the global identifier and the formal implications of term definitions in RDF, i.e., some of the machine-readable semantics will be lost. The terms are intended to be used in combination with metadata terms from other, compatible vocabularies in the context of application profiles. They are used as such in DCAT2 and thus the EJP RD Metadata model, as well as the ontologies that describe data elements, data access information, and provenance.

- Status of development: Normative
- Further Information and official definitions: https://dublincore.org/

5.1.1.3. EJP RD Metadata Model

The EJP RD Metadata Model is a machine-readable model for declaring information about a resource (e.g., a registry) for contributing to the functionality of the VP. The model extends DCAT2 (see 5.1.1.1) with resource types that are relevant for the VP. The model is defined in RDF by which it integrates with other models defined in RDF (a ‘Linked Data’ principle). Serialisation in non-RDF formats is possible (XML, JSON, etc.) at the expense of machine-readable semantics, similar to serialisation of the RDF definition of Dublin Core terms (see 5.1.1.2). Metadata about a resource that is standardised by the EJP RD Metadata Model (and thus also by DCAT2), and made accessible for machines with an interface that conforms to the FAIR Data Point (FDP)

² https://www.dublincore.org/specifications/dublin-core/dcmi-terms/
³ https://dublincore.org/
⁴ https://www.dublincore.org/specifications/dublin-core/dces/ The fifteen-element Dublin Core has been formally standardized as ISO 15836, ANSI/NISO Z39.85, and IETF RFC 5013.
specifications (see 5.2.4.1) increases the Findability and Accessibility of the resource globally and in the VP, and enables the VP to dynamically adapt its functionality to the capabilities of its resources.

- Status of development: Trial use
- Further Information: https://github.com/S2Olga/EJP-RD-metadata-model

5.1.1.4. FASTA, FASTQ

FASTA and FASTQ files are flat file formats used to exchange nucleotide or amino acid sequence information. FASTQ files include a sequencing quality indicator.

- Status of development: Normative
- Further Information:
  - https://www.bioinformatics.nl/tools/crab_fasta.html;
  - https://support.illumina.com/bulletins/2016/04/fastq-files-explained.html

5.1.1.5. ISO 21838

This standard specifies required characteristics of a domain-neutral top-level ontology (TLO) that can be used in tandem with domain ontologies at lower levels to support data exchange, retrieval, discovery, integration and analysis.

If an ontology is to provide the overarching ontology content that will promote interoperability of domain ontologies and thereby support the design and use of purpose-built ontology suites, then it needs to satisfy certain requirements. This document specifies these requirements. It also supports a variety of other goals related to the achievement of semantic interoperability, for example, as concerns legacy ontologies developed using heterogeneous upper-level categories, where a coherently designed TLO can provide a target for coordinated re-engineering.

- Status of development: Normative
- Further Information: https://www.iso.org/standard/71954.html

5.1.1.6. ISO 23903:2021 Health informatics - Interoperability and integration reference architecture - Model and framework

This standard enables the advancement of interoperability from the data/information exchange paradigm to knowledge sharing at decreasing level of abstraction, starting at IT concept level (semantic coordination) through business domain concept level (agreed service function level cooperation), domain level (cross-domain cooperation) up to individual context (skills-based end-user collaboration). The standard defines a model and framework for a harmonized representation of existing or intended systems with a specific focus on ICT-supported business systems. The Interoperability and Integration Reference Architecture supports ontology harmonization or knowledge harmonization to enable interoperability between, and integration of, systems, standards and solutions at any level of complexity without the demand for continuously adapting/revising those specifications. The approach can be used for analysing, designing, integrating, and running any type of systems. For realizing advanced interoperability, flexible, scalable, business-controlled, adaptive,
knowledge-based, intelligent health and social ecosystems need to follow a systems-oriented, architecture-centric, ontology-based and policy-driven approach.

- Status of development: Normative
- Further Information and official definitions: https://www.iso.org/standard/77337.html

5.1.1.7. Maelstrom Data harmonization guidelines

Maelstrom guidelines for retrospective data harmonization were developed by the Maelstrom Research team to ensure quality, reproducibility, and transparency of the data harmonization process. Based on these guidelines, retrospective harmonization is an iterative process involving a series of closely related, interdependent, and often integrated steps.

- Status of development: Normative
- Further Information: https://www.maelstrom-research.org/page/maelstrom-guidelines

5.1.1.8. MIABIS 2.0

The Minimum Information About Biobank data Sharing (MIABIS) aims to standardize data elements used to describe biobanks, research on samples and associated data. The MIABIS Community Standards work on several granularity levels, with the aim to support interoperability between biobanks sharing their data. General attributes to describe biobanks, sample collections and studies at an aggregated/metadata level are defined in MIABIS Core 2.0 (Merino-Martinez et al., 2016). New MIABIS modules describing samples and sample donors at individual level have been approved by BBMRI-ERIC and are described here (Eklund et al., 2020).

- Status of development: Normative
- Further Information: https://github.com/MIABIS/miabis/wiki

5.1.1.9. Phenopackets schema

The goal of the phenopacket-schema is to define the phenotypic description of a patient/sample in the context of rare disease, common/complex disease, or cancer. The schema as well as source code in Java, C++, and Python is available from the phenopacket-schema GitHub repository.

Version 1 of phenopackets was approved by GA4GH in October 2019. Based on initial experiences and feedback from multiple sources, and discussions in the GA4GH Clin/Pheno Workstream and Phenopackets Subgroups, version 1 has been extended to include better representation of the time course of disease, treatment, and cancer-related data. The current document refers to the version 2 of the Phenopackets schema. Version 2 is currently being finalized by the Global Alliance for Genomics and Health (GA4GH) Clinical & Phenotypic Data Capture workstream.

- Status of Development:
5.1.1.10. **RD3 (Rare Disease Data about Data) data model**

Metadata database to track and find samples processed in the Sandbox and submitted to the EGA (see 0), including details on patient phenotypes, sample preparation, sequencing and information about files collected. RD3 is now being generalized with Dutch FAIR genomes initiative, [http://fairgenomes.github.io](http://fairgenomes.github.io) in consideration of EJP RD metadata model. A reference implementation exists using MOLGENIS, see below.

- Status of development: Normative
- Further Information: [https://github.com/molgenis/RD3_database](https://github.com/molgenis/RD3_database)

5.1.2. **Standards on data file formats, markup, and annotation**

5.1.2.1. **BED, GFF**

The browser extensible data (BED) format is a concise and flexible way to represent genomic features and annotations

- Status of development: Normative
- Further Information:
  - [https://bedtools.readthedocs.io/en/latest/content/general-usage.html#bed-format](https://bedtools.readthedocs.io/en/latest/content/general-usage.html#bed-format)
  - [https://bedtools.readthedocs.io/en/latest/content/general-usage.html#gff-format](https://bedtools.readthedocs.io/en/latest/content/general-usage.html#gff-format)

5.1.2.2. **CSV**

Comma Separated Values is a common file format for tabular data, an old and simple flat-file format for representing data (text and values) in a rectangular matrix. It is not a formal “standard” as such but is very commonly used. A common alternative is the tab-delimited file format. [RFC4180](https://tools.ietf.org/html/rfc4180) describes the CSV format and mime-type for the internet community.

- Status of development: Normative
- Further Information: [https://datahub.io/docs/data-packages/csv](https://datahub.io/docs/data-packages/csv)

5.1.2.3. **ISA-Tab+serialisations (ISA-JSON etc)**

The ISA Abstract Model, originally developed as a tabular format (ISA-Tab) since 2007, has been developed with several international collaborators and in synergy with related, domain-specific effort. ISA is supported as a tabular format (ISA-Tab) and a JSON format (ISA-JSON), with additional machine readable semantics as Linked Data ([linkedISA](https://linkedISA)), and by a programmable Python API ([ISA_API](https://isa-api)).
5.1.2.4. **JSON LD**

JavaScript Object Notation for Linked Data (RDF) that allows for semantic capture in the JSON syntax.

- Status of development: Normative
- Further Information: [https://json-ld.org/](https://json-ld.org/)

5.1.2.5. **OML**

Omics Markup Language (ISO/DIS 21393) is a data exchange format that is designed to facilitate exchanging omics data around the world without forcing changes of any database schema.

- Status of development: Normative
- Further Information: [https://www.iso.org/standard/70855.html](https://www.iso.org/standard/70855.html)

5.1.2.6. **OpenAPI Specification**

The OpenAPI Specification, previously known as the Swagger Specification, is a specification for machine-readable interface files for describing, producing, consuming, and visualizing RESTful web services.

- Status of development: Normative

5.1.2.7. **Schema.org**

Schema.org is a collaborative, community activity with a mission to create, maintain, and promote schemas for structured data on the Internet, on web pages, in email messages, and beyond.

- Status of development: Normative
- Further Information: [http://schema.org](http://schema.org)

5.1.2.8. **XML**

Extensible Markup Language (XML) is a markup language that defines a set of rules for encoding documents in a format that is both human-readable and machine-readable.

- Status of development: Normative
5.1.3. Standard data element sets

5.1.3.1. CCE - Common Condition of use Elements

This provides a core list of non-directional, atomic ‘concepts’ that specify types of use of an asset (e.g., the where, why, by whom, when etc). It aims to provide the basis for a common set of ‘conditions of use’ statements that resources (e.g., biobank, datasets, collections) can optionally assert, so that such information can be compiled, exchanged, aligned, and used for discovery and access decision making. These statements can be formulated as Digital Use Condition (DUC) constructs, and this is what is being employed presently to assess the utility of CCEs. Version 1.0 of CCE comprises 20 elements, applicable to registries and biobanks, which has been alpha- and beta-tested and is now being prepared for publication.

- Status of development: Trial use
- Further Information: https://docs.google.com/document/d/1EjmI2DFKcN5DMJ6qllgAjaP2r35Nmi7RQmHrioQ8fhg/edit?usp=sharing

5.1.3.2. CDE - Common Data Elements

Widely used minimum set of data fields to be collected by EU Rare Disease registries comprising a set of 16 common data elements, released by the EU RD Platform aiming at increasing interoperability of RD registries. The CDEs ensure basic utility of RD registry data and help with standardization of datasets and basic interoperability in the sense of having the same types of data elements across registries. However, this provides no guarantee that the type of information is harmonized across registries by a uniform data model or that query APIs are uniform. This is achieved by making CDEs available in the form of the machine readable CDE semantic model (see 5.1.4.1) and providing a common query API.

- Status of development: Normative

5.1.4. Standards on data models

5.1.4.1. Basic Formal Ontology (BFO) and The Open Biological and Biomedical Ontology (OBO) Foundry ontologies

The BFO is focused on the task of providing the upper ontology as the foundation for all OBO Foundry ontologies that cover specific domains of scientific research, as for example in biomedicine. There are many OBO domain ontologies that together cover large parts of known biomedical reality. Among them HPO (Human Phenotypes), MONDO (Diseases), NCIT (Cancer and many associated concepts), and OBIB (biobanks). OBO foundry poses a set of restrictions and best practices to guarantee rigour and interoperability between OBO ontologies.

- Status of development: Normative

5.1.4.2. Clinical Data Interchange Standards Consortium (CDISC)
The Clinical Data Interchange Standards Consortium (CDISC) develops and advances data standards to transform incompatible formats, inconsistent methodologies, and diverse perspectives into a framework for generating clinical research data.

- Status of development: Normative
- Further Information: https://www.cdisc.org/standards

5.1.4.3. bioCADDIE Data Tag Suite (DATS)
DATS, which stands for DAta Tag Suite, is a data description model designed and produced to describe datasets being ingested in DataMed, a prototype for data discovery developed as part of the NIH Big Data 2 Knowledge bioCADDIE project.

- Status of development: Normative
- Further Information: https://datatagsuite.github.io/docs/html/

5.1.4.4. DUC – Digital Use Conditions
This provides a standardised structure for expressing statements that specify conditions of use or consent. It comprises a Header (metadata) section, and a body of at least one statement. Each statement refers to a non-directional, atomic ‘type of use’ concept (see CCE), termed a ‘Condition Term’, with an optional ‘Condition Detail’ modifier that gives further details of that instance of that type of use. This is then given directionality via a ‘Rule’ that states whether that form of use is Obligated, Permitted or Forbidden, and a ‘Scope’ attribute that defines whether the condition of use applies to the ‘whole of the asset’ or ‘part of the asset’. Version 1.0 of DUC using CCE has been alpha- and beta-tested and will now be piloted as a basis for representing CCE statements to underpin VP discovery activities. DUC is compatible with other solutions that are used in the field such as DUO (Data Use Ontology) and effort is underway to express DUC Rules in ODRL (Open Digital Rights Language – A W3C standard) as a base ontology.

- Status of development: Trial use
- Further Information: https://docs.google.com/document/d/1EjmI2DFKcN5DMJ6qll2AJqP2z35Nmi7RQmHrioQ8thg/edit?usp=sharing
  https://docs.google.com/spreadsheets/d/1P23mP1ZCicLpq50allg0yusMWOW8Hewglzp8TKNQ-s/edit?usp=sharing

5.1.4.5. JRC Common Data Element Semantic Data Model
Semantic Model for the set of 16 Common Data Elements (CDEs) for Rare Diseases Registration released by the EU RD Platform to increase interoperability of RD registries (see 5.1.3.1). It maps the CDEs and qualified relationships between them to standard
ontologies. The basis for each data element module is a semantic design pattern defined by the Semanticscience Integrated Ontology (SIO). The model aims to represent, in a globally understood, machine readable language, what the values of CDE data records in registry datasets mean: a uniform way for computers to ‘understand’ data records across multiple registries. By annotating data records with the model at source, e.g., an RD registry, data records become globally linkable (interoperable) and machine actionable (‘ontologised data’). The model is defined in RDF, similar to the other semantic models in this document (e.g., DCAT2, the EJP RD Metadata model, aforementioned ontologies). Serialisations in other formats are possible at the expanse of machine-readable semantics. It should be noted that the access conditions that apply to the original data should also be applied to the ‘ontologised data’.

- Status of development: Normative
- Further Information: https://github.com/ejp-rd-vp/CDE-semantic-model

5.1.4.6. OMOP (OHDSI object model)

The OMOP Common Data Model allows for the systematic analysis of disparate observational databases. The concept behind this approach is to transform data contained within those databases into a common format (data model) as well as a common representation (terminologies, vocabularies, coding schemes), and then perform systematic analyses using a library of standard analytic routines that have been written based on the common format.

- Status of development: Normative

5.1.4.7. Resource Description Framework (RDF) and Linked Data

The Resource Description Framework is a World Wide Web Consortium (W3C) Recommendation for representing information in the Web and expressing machine readable descriptions of resources. The building blocks are subject-predicate-object triples, where the elements may be hyper-links (typically Uniform Resource Identifiers, URIs), blank nodes, or data-typed literals (e.g., the values in registry records). RDF triples form graphs that express meaning for computers. The Web Ontology Language (OWL), used by most biomedical ontologies, is defined in RDF. By using URIs for connecting nodes and edges, RDF builds on web standards that have proven to scale to globally federated networks. RDF is closely associated with the Linked Data Principles: (i) use URIs as names for things; (ii) use HTTP URIs so that people can look up those names; (iii) provide useful information when someone looks up a URI, using the standards (RDF, SPARQL’); (iv) include links to other URIs, such that they can discover

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5 Rajaram Kaliyaperumal, Mark D. Wilkinson, et al., Semantic modelling of Common Data Elements for Rare Disease registries, and a prototype workflow for their deployment over registry data. https://doi.org/10.1101/2021.07.27.21261169
7 https://www.w3.org/TR/sparql11-overview/
more things. In summary, the properties and purpose of RDF make it the default backbone for machine readable semantic models, including those used in the VP. NB RDF should not be confused with file formats: RDF is a data model. RDF graphs can be serialised and exchanged in various formats (XML, JSON-LD, Turtle, etcetera). The W3C-recommended ‘SPARQL’ language is the default query language for RDF graphs.

- Status of development: Normative
- Further Information: https://www.w3.org/RDF/

5.1.5. Standards on data ontology, terminology and vocabulary

5.1.5.1. Anatomical Therapeutic Chemical (ATC)
The Anatomical Therapeutic Chemical (ATC) Classification System is used for the classification of active ingredients of drugs according to the organ or system on which they act and their therapeutic, pharmacological and chemical properties. It is controlled by the World Health Organization Collaborating Centre for Drug Statistics Methodology (WHOCC) and was first published in 1976.

- Status of development: Normative
- Further Information:
  - https://www.who.int/tools/atc-ddd-toolkit/atc-classification
  - https://bioportal.bioontology.org/ontologies/ATC

5.1.5.2. Data Catalogue Vocabulary (DCAT)
Data Catalog Vocabulary (DCAT) is an RDF vocabulary designed to facilitate interoperability between data catalogues published on the Web. By using DCAT to describe datasets in catalogues, publishers increase discoverability and enable applications to consume metadata from multiple catalogues. It enables decentralized publishing of catalogues and facilitates federated dataset search across catalogues. Aggregated DCAT metadata can serve as a manifest file to facilitate digital preservation

- Status of development: Normative
- Further Information: https://www.w3.org/TR/vocab-dcat-2/

5.1.5.3. Data Use Ontology (DUO)
DUO is an ontology which represent data use conditions. DUO allows to semantically tag datasets with restriction about their usage, making them discoverable automatically based on the authorization level of users, or intended usage. It is a GA4GH approved standard

- Status of development: Normative
  https://obofoundry.org/ontology/duo.html
5.1.5.4. **FAIRplus recipe on how to choose controlled vocabulary**

Using ontologies to represent the data items in a data set is a great way to comply with the FAIR principles. The main purpose of this recipe is to provide guidance on how to select the most suitable semantic artefacts given a specific research context in general, and when it comes to life and biomedical sciences projects, their main themes, i.e. risk assessment, clinical trial, drug discovery or fundamental research.

- Status of development
- Further information: [https://faircookbook.elixir-europe.org/content/recipes/interoperability/selecting-ontologies.html](https://faircookbook.elixir-europe.org/content/recipes/interoperability/selecting-ontologies.html)

5.1.5.5. **Gene Ontology (GO)**

The Gene Ontology (GO) knowledgebase is the world’s largest source of information on the functions of genes. This knowledge is both human-readable and machine-readable, and is a foundation for computational analysis of large-scale molecular biology and genetics experiments in biomedical research.

- Status of development: Normative
- Further information: [http://geneontology.org](http://geneontology.org)

5.1.5.6. **GENO**

Genotype Ontology, an integrated ontology for representing the genetic variations described in genotypes, and their causal relationships to phenotype and diseases.

- Status of development: Draft
- Further Information:
  - [https://www.ebi.ac.uk/ols/ontologies/geno](https://www.ebi.ac.uk/ols/ontologies/geno)
  - [https://bioportal.bioontology.org/ontologies/GENO](https://bioportal.bioontology.org/ontologies/GENO)

5.1.5.7. **HGNC (HUGO)**

HUGO Gene Nomenclature Committee is the resource for approved gene nomenclature, and is part of the CDE for rare disease registration (see 5.1.4.1).

- Status of development: Normative
- Further Information: [https://www.genenames.org/](https://www.genenames.org/)

5.1.5.8. **Sequence Variant Nomenclature (HGVS)**

The HGVS Nomenclature is a set of recommendations when describing sequence variant in a consistent and unambiguous manner to facilitate the report and exchange of information on the analysis of a genome. It is an IRDiRC Recognized Resource and is part of the CDE for rare disease registration (see 5.1.4.1).

- Status of development: Normative
Further Information: http://varnomen.hgvs.org

5.1.5.9. HPO-ORDO ontological module (HOOM)
HOOM is a module that qualifies the relationship between a clinical entity and phenotypic abnormalities according to its frequency of occurrence in the disease population and further qualifiers such as “pathognomonic sign” and “diagnostic criterion”. It is based on Orphanet knowledge base of annotations of rare diseases by its phenotypes, using Human Phenotype Ontology (HPO)

- Status of development: Normative
- Further Information: http://www.orphadata.org/cgi-bin/index.php#hoommodal

5.1.5.10. Human Phenotype Ontology (HPO)
The Human Phenotype Ontology (HPO) provides a standardized vocabulary of phenotypic abnormalities encountered in human disease. It is an IRDiRC Recognized Resource and is part of the CDE for rare disease registration (see 5.1.4.1). The HPO is currently being developed using the medical literature, Orphanet, DECIPHER, and OMIM.

- Status of development: Normative
- Further Information: https://hpo.jax.org/app/

5.1.5.11. International Classification of Diseases (ICD)
International Classification of Diseases developed and maintained by the World Health Organisation (WHO), integrated in the WHO’s Family of International Classifications (WHO-FIC). ICD is historically used for statistical reports on mortality and morbidity and is the leading classification used in health information systems across the world in its different versions and national adaptations. Current ICD international versions in use are ICD-10 and ICD-11.

- Status of development: Normative
- Further Information: https://icd.who.int

5.1.5.12. International Classification of Functioning, Disability and Health (ICF)
Part of the WHO’s Family of International Classifications (WHO-FIC), ICF is the WHO framework for measuring health and disability at both individual and population levels. It is recommended in the CDE for rare disease registration (see 5.1.4.1).

- Status of development: Normative
- Further Information: https://www.who.int/standards/classifications/international-classification-of-functioning-disability-and-health
5.1.5.13. International Classification of Diseases for Oncology (ICD-O)

Part of the WHO’s Family of International Classifications (WHO-FIC), ICD-O is a multi-axial classification of the site, morphology, behaviour, and grading of neoplasms. It is used principally in tumour or cancer registries for coding the site (topography) and the histology (morphology) of neoplasms, usually obtained from a pathology report. Current version is ICD-O-v3.

- Status of development: Normative
- Further Information: https://www.who.int/standards/classifications/other-classifications/international-classification-of-diseases-for-oncology

5.1.5.14. Informed Consent Ontology (ICO)

Informed Consent Ontology (ICO) is a community-based ontology in the domain of informed consent. It is an OBO library ontology and developed by following the OBO Foundry principles.

- Status of development: Normative
- Further Information:
  - https://obofoundry.org/ontology/ico.html
  - https://bioportal.bioontology.org/ontologies/ICO

5.1.5.15. LOINC

Logical Observation Identifiers Names and Codes. The international standard for identifying health measurements, observations, and documents.

- Status of development: Normative
- Further Information: https://loinc.org/

5.1.5.16. MedDRA - Medical Dictionary for Regulatory Activities Terminology

Rich and highly specific standardised medical terminology to facilitate sharing of regulatory information internationally for medical products used by humans.

- Status of development: Normative
- Further Information: https://www.meddra.org/

5.1.5.17. MeSH - Medical Subject Headings

MeSH is a comprehensive controlled vocabulary for the purpose of indexing journal articles and books in the life science.

- Status of development: Normative
- Further Information: https://meshb.nlm.nih.gov/
5.1.5.18.  **NCIT - National Cancer Institute Thesaurus**
A vocabulary for clinical care, translational and basic research, and public information and administrative activities.

- Status of development: Normative
- Further Information:
  o  [https://ncithesaurus.nci.nih.gov/ncitbrowser/](https://ncithesaurus.nci.nih.gov/ncitbrowser/)
  o  [https://bioportal.bioontology.org/ontologies/NCIT](https://bioportal.bioontology.org/ontologies/NCIT)

5.1.5.19.  **Online Mendelian Inheritance in Man (OMIM)**
Online Mendelian Inheritance in Man is an online catalogue of human genes and genetic disorders. It is an IRDiRC Recognized Resource.

- Status of development: Normative
- Further Information: [https://omim.org](https://omim.org)

5.1.5.20.  **Orphanet nomenclature of rare diseases (ORPHAcodes) and Orphanet ontology of rare diseases (ORDO)**
Orphanet nomenclature of rare diseases (ORPHAcodes) and its enriched ontological format Orphanet Rare Diseases Ontology (IRDiRC Recognized Resource). Orphanet nomenclature is a standardised vocabulary allowing semantic annotation of rare disease diagnosis and is part of the CDE for rare disease registration (see 5.1.4.1). ORDO includes alignments between ORPHAcodes and other medical terminologies (ICD-10, OMIM, MedDRA, UMLS, MeSH), gene-disease relationships, disease epidemiological data by geographical location. The Orphanet nomenclature of rare diseases and its alignments with other terminologies files, including SNOMED CT, are also released in XML and JSON formats in Orphadata.org, recognised ELIXIR Core Data Resource. A dedicated API and a Data Visualisation tool are also available.
Orphanet nomenclature and ORDO are multilingual resources.

- Status of development: Normative
- Further Information:
  o  [www.orphadata.org](http://www.orphadata.org)
  o  [http://bioportal.bioontology.org/ontologies/ORDO](http://bioportal.bioontology.org/ontologies/ORDO)

5.1.5.21.  **PROV Ontology**
It provides a set of classes, properties, and restrictions that can be used to represent and interchange provenance information generated in different systems and under different contexts

- Status of development: Normative
- Further Information: [https://www.w3.org/TR/prov-o/](https://www.w3.org/TR/prov-o/)
5.1.5.22. Semanticscience Integrated Ontology (SIO)
The Semanticscience Integrated Ontology (SIO) is an ontology to facilitate biomedical knowledge discovery. SIO features a simple upper level comprised of essential types and relations for the rich description of arbitrary (real, hypothesized, virtual, fictional) objects, processes and their attributes. SIO specifies simple design patterns to describe and associate qualities, capabilities, functions, quantities, and informational entities including textual, geometrical, and mathematical entities, and provides specific extensions in the domains of chemistry, biology, biochemistry, and bioinformatics.

- Status of development: Normative

- Further Information:
  - https://github.com/MaastrichtU-IDS/semanticscience

5.1.5.23. SNOMED CT
SNOMED Clinical Terms is a systematically organized computer processable collection of medical terms providing codes, terms, synonyms and definitions used in clinical documentation and reporting. SNOMED CT is considered to be the most comprehensive, multilingual clinical healthcare terminology in the world. The primary purpose of SNOMED CT is to encode the meanings that are used in health information and to support the effective clinical recording of data with the aim of improving patient care. SNOMED CT provides the core general terminology for electronic health records. SNOMED CT comprehensive coverage includes clinical findings, symptoms, diagnoses, procedures, body structures, organisms and other aetiologies, substances, pharmaceuticals, devices and specimens.

- Status of development: Normative

- Further Information: https://www.snomed.org/

5.1.6. Standards on data discovery
5.1.6.1. Automatable Data Discovery and Access Matrix (ADA-M)
Automatable Data Discovery and Access Matrix (GA4GH Standard) Comprehensive information model that provides the basis for producing structured metadata “Profiles” of data access regulatory conditions (e.g., Informed consent information in a machine-readable format).

- Status of development: Normative

- Further Information: https://github.com/ga4gh/ADA-M

5.1.6.2. Beacon-1
The Beacon protocol defines an open standard for biomedical data and sample discovery, developed by members of the Global Alliance for Genomics & Health.

- Status of development: Normative
Further Information: https://github.com/ga4gh-beacon/specification

5.1.6.3. Beacon-2 API
GA4GH API specification adapted for the EJP-RD context (via data field/value and filter element specifications) to query metadata and/or record level information relating to resources description, diagnoses, genes, phenotypes, samples, demographics, and asset availabilities. Validated extensions enable interaction with many other areas of standardisation in GA4GH. It is one way that federated queries may be structured.

- Status of development: Normative
- Further Information: https://beacon-project.io/

5.1.6.4. Bioschemas
Bioschemas aims to improve the Findability on the Web of life sciences resources such as datasets, software, and training materials. It does this by encouraging people in the life sciences to use Schema.org markup in their websites so that they are indexable by search engines and other services.

- Status of development: Normative
- Further Information: https://bioschemas.org/

5.1.7. Standards on data exchange mechanisms

5.1.7.1. DOIP - Digital Object Interface Protocol
The Digital Object Interface Protocol (DOIP) is a simple, but powerful conceptual protocol for software applications (“clients”) to interact with “services” which could be either the digital objects or the information systems that manage those digital objects.

- Status of development: Normative
- Further Information: https://www.dona.net/doi_pv1doc

5.1.7.2. FHIR - Fast Healthcare Interoperability Resources
Standard for exchanging healthcare information electronically.

- Status of development: Normative
- Further Information: https://www.hl7.org/fhir/overview.html
5.1.7.3. **HL7 FHIR4FAIR – FHIR Implementation Guide**

The HL7 FHIR4FAIR – FHIR Implementation Guide aims to provide guidance on how HL7 FHIR can be used for supporting **FAIR health data** implementation and assessment to enable a cooperative usage of the HL7 FHIR and FAIR paradigms. Other kinds of health-related artefacts, such as clinical guidelines, algorithms, software, models are out of scope. The Guide aims to allow researchers to make available under specified conditions of use set of data, derived from a data source, that have been collected and consolidated for a specific purpose; and to allow researchers and data scientists to look for and access previously collected data sets to answer specific questions.

- Status of development: Normative
- Further Information and official definitions: [HL7.FHIR.UV.FHIR-FOR-FAIR\FHIR for FAIR Home Page - FHIR v4.3.0](https://www.hl7.org/standards/fhir/v4.3.0/index.html)

5.1.7.4. **ISO /AWI TR 24305 Health informatics - Guidelines for implementation of HL7/FHIR based on ISO 13940 and ISO 13606**

This standard which is currently under development will provide guidelines on how to implement FHIR based on ISO 13940 (Health informatics — System of concepts to support continuity of care) and ISO 13606 (Health informatics – Electronic health record communication).

- Status of development: Draft
- Further Information and official definitions: [https://www.iso.org/standard/78390.html](https://www.iso.org/standard/78390.html)

5.1.7.5. **mzML/mzIdentML**

ProteomeXchange Consortium was established to provide globally coordinated standard data submission and dissemination pipelines involving the main proteomics repositories, and to encourage open data policies in the field.

- Status of development: Normative
- Further Information:
  - [http://www.psidev.info/mzidentml](http://www.psidev.info/mzidentml)
  - [www.proteomexchange.org](http://www.proteomexchange.org)

5.1.7.6. **PhenoPackets**

This standard (ISO/WD 4454) enables the exchange of clinical phenotype related information between information systems within EJP RD and with other project (such as Solve-RD), not least to/from ERN registries, RD-NEXUS discovery tools, Linked Data Platforms, and GPAP. Its scope includes data on individuals and family/pedigree information. It is compatible with ontologies such as the HPO, ORDO and OMIM and with genetic data. It is a GA4GH approved standard.

A semantic model of PhenoPackets has been developed within EJP RD.

- Status of development: Normative
• Further Information: http://phenopackets.org

5.1.7.7. **REST**

Representational State Transfer (Design Guide). Representational State Transfer (REST) is an architectural style for distributed hypermedia systems, describing the software engineering principles guiding REST and the interaction constraints chosen to retain those principles, while contrasting them to the constraints of other architectural styles.

- Status of development: Normative
- Further Information: https://en.wikipedia.org/wiki/Representational_state_transfer

5.1.8. **Standards on security, authentication, and authorisation**

5.1.8.1. **GA4GH Passport**

A standard for a global federated data sharing network that allows the searching, and subsequent -optional- processing of the results in a cloud environment.

- Status of development: Normative
- Further Information: https://github.com/ga4gh-duri/ga4gh-duri.github.io/blob/master/researcher_ids/ga4gh_passport_v1.md

5.1.8.2. **GA4GH Visa**

An assertion from a Passport Visa Assertion Source organization that is bound to a Passport Visa Identity and signed by a Passport Visa Issuer service whose signature is verifiable via its public key.

- Status of development: Normative
- Further Information: https://github.com/ga4gh-duri/ga4gh-duri.github.io/blob/master/researcher_ids/ga4gh_passport_v1.md#passport-visa

5.1.8.3. **Open ID Connect**

OpenID Connect is the third generation of OpenID technology. It is an authentication layer on top of the OAuth 2.0 authorization framework.[82] It allows computing clients to verify the identity of an end user based on the authentication performed by an authorization server, as well as to obtain the basic profile information about the end user in an interoperable and REST-like manner. In technical terms, OpenID Connect specifies a RESTful HTTP API, using JSON as a data format.

- Status of development: Normative
- Further Information: https://en.wikipedia.org/wiki/OpenID#OpenID_Connect_(OIDC)
5.1.8.4. SAML 2.0

Security Assertion Markup Language 2.0 (SAML 2.0) is a version of the SAML standard for exchanging authentication and authorization identities between security domains. SAML 2.0 is an XML-based protocol that uses security tokens containing assertions to pass information about a principal (usually an end user) between a SAML authority, named an Identity Provider, and a SAML consumer, named a Service Provider. SAML 2.0 enables web-based, cross-domain single sign-on (SSO), which helps reduce the administrative overhead of distributing multiple authentication tokens to the user.

- Status of development: Normative
- Further Information: https://en.wikipedia.org/wiki/SAML_2.0
5.2. Tools

The following tools represent examples of tools that were agreed to be used within the VP. However, this list is not intended to be complete, since any tool that fulfils requirements specified in the VIPS can be included in the VP.

5.2.1.1. FAIR Genomes – ELSI Framework

The FAIR Genomes ELSI Framework is a framework that defines Guidelines for secondary use of Data. The framework defines which elements from Informed consent must be annotated with ontology terms to aid in making it FAIR, which includes making the informed consent machine-readable. With machine-readable ontology terms, the data use conditions are made explicit and machine-readable. It supports the researcher in finding the right data for her/his research study by making it easier to find the data and making it easier to learn what uses are allowed. The ontology terms describe information about the informed consent given, and they do not provide access to the data.

In addition to the machine-readable ontology terms, this framework also describes roughly what else is needed for FAIR data sharing to adhere to legal regulations, and ethical principles and guidelines.

- Status of development: Normative
- Further Information: ELSI rapport FAIR Genomes okt21 v4.pdf (health-ri.nl)

5.2.1.2. FAIR Genomes – Metadata scheme

We have developed the FAIR Genomes metadata scheme based on a national consensus that we achieved during this project among stakeholders in the Netherlands. This schema contains the data elements needed to ease the sharing of NGS data. Where possible, we have learned from international initiatives such as European Joint Program for Rare Disease (EJP-RD), GA4GH, Solve-RD, European Bioinformatics Institute, FAIR sharing, and existing large public databases, where we have adapted existing elements to the FAIR Genomes scheme. This semantic scheme can be used to generate blueprints for different data capture systems and makes data exchange interoperable. With the help of prototypes, we have now shown that this approach is feasible. In addition, tools have been developed to assist with this implementation. The scheme and technology will be further developed in collaboration with (inter)national partners, and we therefore conclude that we have laid a solid foundation for FAIR Genomes-compatible production systems for FAIRification in practice. In addition, by involving stakeholders, we have increased awareness of the FAIR principles among healthcare professionals, and we expect our efforts to contribute to significantly more reuse of NGS data between Dutch institutes. FAIR genomes are a starting point for increasing European semantic interoperability genome data, e.g., as a promising best practice within the 1+ Million Genomes and Beyond 1 Million Genomes Initiatives.

- Status of development: Normative
- Further Information: https://github.com/fairgenomes/fairgenomes-semantic-model
5.2.1.3. FAIR Genomes – Metadata codebook

The FAIR Genomes scheme and codebooks version 1.2 consists of 9 modules that contain 112 metadata elements and 85351 coded lookups in total. It is freely available at https://github.com/fairgenomes/fairgenomes-semantic-model/tree/v1.2. The 9 modules are: Study, Personal, Leaflet and consent form, Individual consent, Clinical, Material, Sample preparation, Sequencing and Analysis. The data elements have preferred value types such as integer, string or date. There is also a special element value type called 'lookup' that refer to codebooks of all user-selectable options. All modules, elements and lookups are defined using ontology terms to prevent ambiguity in the meaning of the concepts used. In addition, these definitions allow computer-readable formats to be created. Also, using the 'reference' value type, modules may refer to other modules as their source, following the typical flow of an NGS analysis in diagnostics or research. For instance, Sample preparation is performed on a Material taken from a Person. Not all terms needed to build the FAIR Genomes codebooks were present in existing ontologies. Therefore, to complete the schema and lookups, we defined 743 new ontology terms. Of these terms, 740 are new lookup values for data-use modifiers, institutes, NGS kits, tissue pathological state, and sequencing instrument models, while three terms represent new element definitions.

- Status of development: Normative
- Further Information: FAIR Genomes metadata schema - Datasets (nictiz.nl)

5.2.1.4. FAIR Genomes – VARDA

To facilitate accurate estimation of variant frequencies, Varda stores all relevant information including data needed to discriminate between no variant called from sufficient data (read coverage) and no variant called because of missing data (no coverage). Additionally, by storing this information it is possible to combine data from many different sources in one database (e.g., gene panel analysis, Whole Exome Sequencing (WES) and Whole Genome Sequencing (WGS)) without compromising the accuracy of calculated frequencies. Data is added to Varda via a bulk import mechanism. Per sample, the client uploads the full list of variants and a list of covered regions. Ideally, this is done automatically from within the data analysis pipeline. For this purpose, a reference implementation of a pipeline module is made available. Querying is done via an application programming interface (API). Preferably, this API is used in the interpretation stage from within a graphical user interface like Agilent Alissa or LOVD+.

- Status of development: Normative
- Further Information: https://vkgl.molgeniscloud.org/

5.2.1.5. FAIR Genomes – Mutalyzer

A method that automatically deals with the HGVS intricacies and outputs correct unambiguous variant descriptions is of high necessity for consistent variant dissemination. In this context, the Mutalyzer tool suite was created to assist geneticists in applying the HGVS guidelines in databases and literature by providing the means for automatic checking and correction of HGVS variant descriptions. In Mutalyzer 3, the third iteration of this tool suite, a set of independent programming libraries was
developed to enable the implementation of variant disambiguation functionality in locally running pipelines. Additionally, a Web API was developed to enable on-line systems, like databases, to incorporate Mutalyzer functionality. Finally, a user friendly website is available for ad hoc analyses.

- Status of development: Normative
- Further Information: https://mutalyzer.nl/

5.2.2. Authentication and Authorization Infrastructure

5.2.2.1. LifeScience AAI

The LifeScience Authentication and Authorisation Infrastructure is a common authentication and authorisation service for the 13 European life science research infrastructures. The service is managed by the life sciences community and operated by the e-infrastructure, including GEANT and EGI.

The Life Science AAI issues a new identifier called a Life Science ID to a user who registers to the Life Science AAI and accepts its usage policy. Users authenticate using authentication providers, such as their home universities or the Life Science AAI's Hostel IdP which can be linked to their Life Science ID. It's also possible to use an ORCID, LinkedIn, or Google account. Multiple accounts can be linked to a single LS AAI account, and will be recognized as the same person at the end service. To cater services with specific assurance needs, Life Science AAI supports an assurance framework and will provide a step-up authentication service in the near future.

- Status of development: Normative
- Further Information: https://elixir-europe.org/about-us/commissioned-services/identity-access

5.2.3. Pseudonymisation

5.2.3.1. EUPID

European Unified Patient Identifier (EU Standard) International unique global identifier systems for patients (EUPID) - an EJP-RD supported method for encrypting patient IDs, to help protect patient identity whilst still being able to track and connect patient records (available via ERDRI). Allows to count patients among European registries avoiding double counts.

- Status of Development: Normative
- Further Information: https://eupid.eu/

5.2.3.2. SPIDER

ERDRI:spider (Secure Privacy-preserving Identity management in Distributed Environments for Research) pseudonymisation tool generates pseudonyms for RD patients. In addition, it allows linking and transferring RD patients’ data across registries without revealing patients’ identities.
Using ERDRI.spider pseudonymisation tool researchers can combine structured data and create patient cohorts for studies and research without exposing RD patients’ privacy to risk. While accessing ERDRI.spider services, sensitive patient data is never communicated to the service provider (EU RD Platform). See the SPIDER presentation video for more details.

- Status of Development: normative
- Further Information: SPIDER pseudonymisation tool (europa.eu)

5.2.4. Mapping / Alignment Services

5.2.4.1. Data Model Alignment Service

The CDE semantic model was built to represent, in ‘ontologised’ linked data form, the CDEs defined by JRC for RD registries. The aforementioned data standards are commonly used for health data and some of them are already adopted by ERN registries. Mapping of the CDE semantic model to/from the 3 data standards (CDISC, ODHIS OMOP and FHIR) is therefore needed. Therefore, a data model alignment service is envisioned comprising a mapping table for CDE terms and scripts for transforming data into the standards, and vice versa. The CDE model uses ontological terms from various ontologies to represent the CDEs. These ontological terms are to be mapped to the ontologies or terminologies used in the 3 standards, presented in a mapping table.

- Status of development: Draft
- Further Information: see VIPS

5.2.4.2. OxO

OxO is a service for finding mappings (or cross-references) between terms from ontologies, vocabularies, and coding standards. OxO imports mappings from a variety of sources including the Ontology Lookup Service and a subset of mappings provided by the UMLS.

- Status of Development: Normative
- Further information: https://www.ebi.ac.uk/spot/oxo/

5.2.4.3. The FAIR Evaluator

The FAIR Evaluator is a tool that assembles automated tests of individual FAIR principles and applies them to a digital resource (e.g., a registry metadata record) to determine the resource’s level of compliance with the FAIR Principles. A public version of The Evaluator is available to test open-access resources. A commercial version is available for deployment inside sensitive data spaces, or for cases where the result of the evaluation should not be made public.

- Status of Development: Normative
5.2.5. Consent and use conditions

5.2.5.1. CCE / DUC Profile Creation Tool [ULEIC]
An online form that enables users to created DUC structured ‘Profiles’ for CCE elements, to support the testing and adoption of these standards. The Leicester version of this tool is a standalone service that does not require (but does permit) users to save a copy of the created Profile on the server, with creation date and version. These can be retrieved later and further edited. Download formats include JSON, CSV and TXT. Users can only view and access Profiles that they themselves created.

- Status of development: Trial use
- Further Information: https://duc.le.ac.uk/

5.2.5.2. CCE / DUC Profile Creation Tool [UMCG]
An online form that enables users to created DUC structured ‘Profiles’ for CCE elements, to support the testing and adoption of these standards. The UMCG version of this tool is integrated into the MOLGENIS platform, and thereby provides a local service for groups whose registry is built on the MOLGENIS platform. Access requires login to their MOLGENIS account

- Status of development: Trial use
- Further Information: https://irdirc.molgeniscloud.org/menu/main/home

5.2.6. Record linkage services

5.2.6.1. EUPID
See above (Tools / Pseudonymization / EUPID).

5.2.6.2. SPIDER
See above (Tools / Pseudonymization / SPIDER).

5.2.7. Resource/Data/Sample discovery and access
The following list of resource/data/sample discovery and access tools represents examples that are not intended to be complete. However, these tools represent references that can help by other tool providers to link their tools to the VP. Additionally, they might be used as hubs to the VP for other resource/data/sample providers that prefer to connect to those tools than to link with the VP directly. Detailed lists of resources are described in Del. 11.8 and 11.18.
5.2.7.1. **Beacon in a box**

Beacon-in-a-box is a reference implementation of the GA4GH Beacon-2 standard, and more specifically a specialisation of this according to the EJP-RD Beacon API. It is optionally available as a Dockerised image, and includes an Excel-based upload system customised to the relevant data/metadata, a MongoDB to hold the uploaded data, and a set of relevant Beacon-2 end-point services and filters that utilise (query across) the stored information. This allows adopters to easily establish the Beacon endpoints that can be queried by the VP Portal. For Level-1 queries, results present the custodians metadata as required by the DCAT standard used in EJP-RD. For Level-2 queries, results present counts of matching samples or records.

- Status of Development: Normative
- Further information: https://github.com/ejp-rd-vp/EJP-RD-Beacon-in-a-Box

5.2.7.2. **bio.tools**

bio.tools is an open source, open data registry of biological and biomedical software descriptions used to help researchers find, understand, utilise and cite the resources they need in their day-to-day work. Resources in bio.tools include everything from simple command-line tools and scripts, through to databases and complex, multi-functional analysis workflows. Resources are described in a rigorous semantics and syntax, providing end-users with the convenience of concise, consistent, and therefore comparable information. An entry in bio.tools is assigned a human-readable unique identifier which provides a persistent reference to the resource even after the resource is no longer accessible. In this way bio.tools preserves information and ensures citations are always available.

- Status of Development: Normative
- Further information: https://bio.tools

5.2.7.3. **Castor**

Castor is a web-interface data capture system that transforms and hosts your data, making it FAIR-at-source without requiring previous experience and programming knowledge. It allows connection to the VP in all levels, making your dataset discoverable and queryable. The costs associated with this tool vary according with the specific necessities of the study (EDC, eConsent, API), the number of patients that will be involved and the type of organization conducting the study (e.g., Biopharma, CRO, Academic Research).

- Status of Development: Normative
- Further information: https://www.castoredc.com/

5.2.7.4. **CDE in a box/FAIR in a box**

CDE in a box is a tool suite for generating, storing, and publishing common data elements (CDEs) according to the CDE semantic model. The suite performs a data-
uplifting activity - it takes CSV files as inputs and generates CDE model-compliant semantic RDF files and stores them in a secure triplestore. CDE in a box is composed of three major components: a triplestore to store ontologised version of the CDE dataset and its metadata, FAIR Data Point (FDP) software to publish the metadata of the CDE dataset, and a transformation service to convert CDEs that are provided to the CDE in a box as CSV files. FAIR in a box extends CDE in a box to include: a fully automated installer; an FDP loader that uses an MS Excel template to capture metadata facets and load them into an FDP; a metadata updater service, that keeps the FDP metadata aligned with each re-load of the CDE data layer; a pre-configured SSH proxy; a Beacon2 “individuals” API. The components used in the CDE/FAIR in a box are dockerized. Users can easily deploy these components on servers that have the docker engine installed.

- Status of Development: Normative
- Further Information:  
  https://github.com/ejp-rd-vp/FiaB

5.2.7.5. Combined RD-Nexus + CDE/FAIR-in-a-Box

The CDE/FAIR-in-a-box is a collection of software applications which enables creation, storing and publishing of "Common Data Elements" according to the CDE semantic model.

RD-Nexus is a flexible web-based data discovery tool that enables data discovery. The main advantage of data discovery using Cafe Variome/RD Nexus is to find data from a Federation of Networks.

The combination of these tools allows you to use the authentication service provided by RD-Nexus along with the semantic technologies provided by CDE/FAIR-in-a-box. The local implementation allows you to control your data better.

Can be implemented as a local implementation or it can be hosted by a pro

- Status of Development: Trial use
- Further information:

5.2.7.6. FAIR Data Point (FDP)

FAIR Metadata Endpoint Serves as a promoter of your registry, increasing its Findability by providing both human and machine-readable metadata. The FDP metadata follows the DCAT standard, which can easily be extended to carry other FAIR-required information such as the accessibility constraints for your data (e.g., the conditions of use, and contact person for obtaining access).

- Status of development: Normative
- Further Information:  
5.2.7.7. **FDP reference implementation**

This tool helps make your resource discoverable by making your metadata available, adding it to the FDP index. This makes your resource’s metadata appear in the VP. With FDP you can present your metadata in connection level 1 to the VP. Using the FAIR-in-a-box solution also provides limited non-authenticated Level 2 access – independent of Beacon2 – via the “grlc” tool that is distributed with FAIR-in-a-box. Grlc provides anonymous/aggregate record level data from pre-approved queries intended to be exposed without authentication. The FDP reference implementation (alone or via FAIR-in-a-box) can be either hosted on your own infrastructure, through an agreement with a non-commercial provider, or by one of several FAIR-compliant commercial hosting providers.

- Status of Development: Normative
- Further information: https://doi.org/10.1162/dint_a_00160
  https://doi.org/10.1162/dint_a_00161

5.2.7.8. **GRLC**

grlc is a lightweight server that takes SPARQL queries (stored in a GitHub repository, in your local filesystem, or listed in a URL), and translates them to Linked Data Web APIs. This enables universal access to Linked Data. Users are not required to know SPARQL to query their data, but instead can access a web API. Grlc, as originally designed, is not sufficiently secure to run over sensitive healthcare data.

A version of grlc that has been heavily modified to be safe to use within protected data spaces is available via the FAIR in a box deployment. The FiaB grlc instance is constrained to only be capable of executing a set of peer-reviewed, privacy-preserving, aggregation queries available from the World Duchenne Organization; no other queries can be executed. The FiaB SSL proxy can be easily extended to provide an SSL proxy over the grlc service endpoint, to provide additional security.

- Status of Development: Trial Use
- Further information: https://github.com/World-Duchenne-Organization/grlc-queries
  https://github.com/ejp-rd-vp/FiaB/tree/main/grlc

5.2.7.9. **I2b2 / tranSMART**

I2b2 and tranSMART are modular open source software for query, exploration and analysis of clinical, translational and genomics data.

- Status of Development: Normative
- Further information: https://i2b2transmart.org/
5.2.7.10. INFRAFRONTIER

INFRAFRONTIER is the European Research Infrastructure for the generation, phenotyping, archiving and distribution of model mammalian genomes. The core services of INFRAFRONTIER comprise the systemic phenotyping of mouse mutants in the participating mouse clinics, and the archiving and distribution of mouse mutant lines by the European Mouse Mutant Archive (EMMA). INFRAFRONTIER aids in rare disease research by providing access to nearly 1800 mouse strains (via EMMA) that are related to over 1400 distinct rare diseases. In addition, INFRAFRONTIER provides specialized services such as the generation of germ-free mice (axenic service) and training in state-of-the-art cryopreservation and phenotyping technologies. The INFRAFRONTIER/EMMA DB has been selected as a FAIRfied data resource by the international FAIRSharing Consortium.

- Status of Development: Normative
- Further information: https://www.infrafrontier.eu

5.2.7.11. MOLGENIS

MOLGENIS is a generic, open source and free to use data platform for researchers to accelerate scientific collaborations and for bioinformaticians. MOLGENIS enables its users to quickly create FAIR database online to find, capture, exchange, manage and analyse a wide diversity of scientific data. MOLGENIS is fully customizable: data structure, user interface and layout can be fully changed and custom (bioinformatics) scripts can be plug-in. It provides Excel/CSV based option to configure the tables, column and relations, and then provides generic APIs and user interfaces to query, upload and download data using REST, CSV, Excel. Also, MOLGENIS provides options to extend functionality using ‘apps’, i.e., small JavaScript+html applications that provide a rich and specialised user experience. It is built on industry standard java, JavaScript, REST, PostgreSQL and elasticsearch.

In EJP-RD, MOLGENIS is being used for BBMRI-ERIC Directory, RD-connect sample database, Sandbox/RD3, and IRDIRC consent database described above. In the broader EJP-RD community, MOLGENIS is used for development of rare disease patient registries for Ithaca, SKIN, CRANIO and Genturis. Beyond EJP-RD, MOLGENIS is also used for multi-centre cohort study and real-world evidence catalogues. All these systems gain interoperability potential into EJP-RD by having federated AAI, FAIR data point and REST APIs. This year, the MOLGENIS team has been piloting also semantic extensions that would further ease integration of MOLGENIS based resources into the virtual platform. MOLGENIS is available free for local installation (support via molgenis-support@umcg.nl) but is also provided as ‘SaaS’ (software as a service) to project partners in EJP-RD or for a fee for outside users.

- Status of Development: Normative
- Further information:
  - https://www.molgenis.org/
  - http://github.com/molgenis

5.2.7.12. REDCap

REDCap is a secure web application for building and managing online surveys and databases. While REDCap can be used to collect virtually any type of data in any
environment (including compliance with 21 CFR Part 11, FISMA, HIPAA, and GDPR), it is specifically geared to support online and offline data capture for research studies and operations. The REDCap Consortium, a vast support network of collaborators, is composed of thousands of active institutional partners in over one hundred countries who utilize and support their own individual REDCap systems.

- Status of Development: Normative
- Further information: [https://www.project-redcap.org/](https://www.project-redcap.org/)

5.2.7.13. **RD-Connect GPAP**

The RD-Connect GPAP is a sophisticated and user-friendly online analysis system for RD gene discovery and diagnosis. The RD-Connect GPAP is an IRDiRC recognized resource hosted at the CNAG-CRG. De-identified phenotypic data is collected using HPO, ORDO and OMIM ontologies through custom templates implemented through the RD-Connect GPAP-Phenostore module. Pseudonymized experiment data (exomes and genomes) and metadata are collected in the RD-Connect GPAP, and processed using a standardized analysis and annotation pipeline. Integrated genome-phenome results are made available to authorized users for prioritisation and interpretation of genomic variants in the RD-Connect GPAP. Raw genomic data is deposited at the EGA for long-term archive and controlled access.

- Status of Development: Normative
- Further information: [https://platform.rd-connect.eu](https://platform.rd-connect.eu)

5.2.7.14. **Rare Disease Networked Exploration of the UnSeen (RD-NEXUS)**

A fully featured, modular, secure platform that supports federated discovery networks, whilst being agnostic to the type of asset or data structures/models being interrogated to enable discovery queries. May be installed locally, or hosted elsewhere (e.g., at ULEIC), while all admin actions, data controllership, and datasets remain with the installation and the relevant PIs. Containerised versions available, optionally integrated with the CDE in a box software. Fully customisable in terms of query interfaces, user accounts and permissions, discoverable data sources, threshold counts for positive searches, response types per user (from simple links through to yes/no indicators, count responses, handoffs, and data provisioning) as well as the particular networks that each installation contributes to. It employs multiple query engines synchronously (SQL, ElasticSearch, Neo4J, BCFTools) and is compliant with all relevant global and EJP-RD standards. Can operate a proxy client to enable integration of triple-stores into federated discovery networks, and allows for interrogation of one or more datasets included in each RD-NEXUS installation (obfuscated if necessary) as well as connection to primary databases. Exceptionally powerful and flexible capabilities in terms of ‘similarity’ searching, with options to save, share and re-use queries.

- Status of Development: Normative
- Further Information: [https://github.com/Cafe-Variome/RDNexus](https://github.com/Cafe-Variome/RDNexus)