

# Identifiers.org virtual SPARQL endpoint for IRI schemes conversion

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## ABSTRACT

**Motivation:** On the Semantic Web, in life sciences in particular, data is often distributed via multiple resources. Each of these sources is likely to use their own IRI (International Resource Identifier) for conceptually the same resource or database record. This multitude of identifiers introduces a barrier when executing federated SPARQL queries between different datasets.

**Results:** We introduce a virtual SPARQL endpoint provided by Identifiers.org, using its Registry's knowledge about IRI patterns, to automatically translate identifiers from one IRI pattern to another, making querying across resources easier and more robust. This endpoint is fully SPARQL 1.1 compliant.

**Availability:** This IRI translating endpoint is available at <http://identifiers.org/services/sparql> and is free to use for all.

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## 1 INTRODUCTION

The use of Semantic Web technologies such as the Resource Description Framework (RDF; Manola and Miller (2004)) for publishing data is becoming popular in the life sciences. For example, the Bio2RDF (Belleau *et al.* (2008)) data mashup is one such resource built upon the data from various publicly available biological databases translated into RDF and linked together. More recently, EMBL-EBI launched the EBI RDF platform (Jupp *et al.* (2014)) in order to expose some of the datasets it produces in RDF.

Bio2RDF uses normalized URIs (using in its own domain name) to support federated queries across datasets. Datasets that are part of the EBI RDF platform follow a set of rules regarding the URIs they can use, resulting in a consistent set of URIs used across all participating resources. Hence, while these two platforms provide linked datasets which can be easily queried individually, the use

of two different URI schemes for what are essentially the same concepts makes it difficult to perform cross platform queries using SPARQL (Prud'hommeaux and Seaborne (2008)).

SPARQL, a query language for RDF which supports complex queries, can be used to merge data distributed over multiple RDF resources. To enable querying across different platforms, there is a need to rewrite IRIs on the fly. String manipulation, supported by SPARQL, may be used for this purpose but requires users to know in advance the IRI types being used in each resource, making it a cumbersome and inefficient solution.

Identifiers.org (Juty *et al.* (2012)) provides resolvable persistent IRIs used to identify individual records (based on the existing entity identifiers assigned directly by the data providers). IRI assignment is reliant upon an underlying Registry which contains information about individual datasets (data collection), the resources (or physical locations) which distribute them, and the different IRIs used to identify and access the individual records. For example, "plasma membrane" from the Gene Ontology (Ashburner *et al.* (2000)) is available from AmiGO (Carbon *et al.* (2009)), QuickGO (Binns *et al.* (2009)), Bio2RDF, OBO PURLs, etc.

To resolve cross query issues, Identifiers.org has developed a virtual SPARQL endpoint to support IRI conversion between linked datasets using different IRI schemes. Consequently, this provides an automatic system for linking and querying ever growing number of life science data resources.

## 2 METHODS

The OpenRDF Sesame SPARQL engine (Visser *et al.* (2014)) was extended to translate specific query patterns to allow IRI translation. Specifically, the service has been designed to answer queries requesting IRIs and making use of the predicate `owl:sameAs`. When such queries are detected, the system determines from which data collection the identified entity originates, extracts from the Registry the alternative

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root URIs for this concept, regenerates complete IRI alternative forms and returns them. The main integration point within the SESAME engine is the *getStatements* method of a class extending *org.openrdf.query.algebra.evaluation.TripleSource*.

The database underlying Identifiers.org's Registry was extended in order to record alternative IRI schemes, and IRI-specific conversion patterns. This has additionally required a curatorial effort to ensure the necessary information has been stored in the Registry.

This is a "virtual" SPARQL endpoint, directly driven by the database underpinning the Registry, since a triple store would not have been able to provide the on demand conversion services required. The service is exposed at <http://identifiers.org/services/sparql>. LODestar (Jupp (2014)), used by the EBI RDF platform, has been deployed to provide a query interface.

### 3 RESULTS

The Identifiers.org virtual SPARQL endpoint can be used to list all alternative URI schemes available for a given URI. It can also be used to check whether two IRIs describes the same concept (although it is not a mapping service between entities provided by different datasets). Example queries can be found at the endpoint page URL.

With the Identifiers.org SPARQL service, it is possible to avoid using string manipulation within a SPARQL query (figure 1) and does not require users to know which IRI patterns are being used within a given resource.

The query below illustrates how the service can be used to integrate resources using different IRI schemes. In this example, the resources are the EBI RDF platform (which uses IRIs of the form "<http://identifiers.org/go/GO:0006915>" for GO terms) and Bio2RDF (which uses IRIs of the form "<http://bio2rdf.org/go:0006915>" for GO terms). The query lists the cross-references to Gene Ontology terms present in the computational model "Edelstein1996 - EPSP ACh event" (BIOMD0000000001) from BioModels Database (Li *et al.* (2010)). It is executed from the BioModels' SPARQL endpoint (<http://www.ebi.ac.uk/rdf/services/biomodels-sparql>) and connects to Identifiers.org to retrieve the alternative IRIs for the retrieved GO terms. Those are then passed to the Bio2RDF SPARQL endpoint to retrieve the description of each GO term.

```
SELECT DISTINCT ?species ?annotation ?description WHERE {
  <http://identifiers.org/biomodels.db/BIOMD0000000001>
    sbmlrdf:species ?species .
  ?species bqbio:isVersionOf ?annotation .

  SERVICE <http://identifiers.org/services/sparql>{
    ?annotation owl:sameAs ?otherURIs .
  }

  SERVICE <http://bioportal.bio2rdf.org/sparql>{
    ?otherURIs dcterms:description ?description .
  }
}LIMIT 10
```

**Fig. 1.** Using Identifiers.org virtual SPARQL endpoint to transform Identifiers.org IRIs into Bio2RDF IRIs.

### 4 DISCUSSION

There are a number of mapping services aimed to connect identifiers from one database to another. However, these are conceptual mappings, for instance relating a gene identifier to its protein product. In contrast, Identifiers.org records all access URLs and IRI-based identification schemes used to identify individual entities from a single data collection.

Also, while the produced endpoint is SPARQL 1.1 compliant, its virtual nature (not based on an actual triple store) means the queries available are limited to IRIs using the `owl:sameAs` predicate in a basic graph pattern; otherwise no results will be returned. The correct SPARQL solution would be to return the infinite possible IRI translations, but although a technically correct solution, would not be useful in practice. Instead we take the approach that Identifiers.org SPARQL endpoint is a remote service not under direct control by the user, and in our case it is under control of entity that deletes everything in the graph when such a query arrives and restores it after the query is answered.

This solution provides users the illusion of a valid SPARQL service that can translate all IRI patterns recorded in the Identifier.org Registry as if all these queries were materialised, at a fraction of the cost.

### 5 CONCLUSION

This IRI conversion service, provided by Identifiers.org as a virtual SPARQL endpoint, enhances the federated query capacities of SPARQL 1.1 to effectively combine information from multiple heterogeneous datasets.

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