Natural Language Question Analysis for Querying Biomedical Linked Data

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Abstract. Biomedical knowledge is disseminated in knowledge bases which become increasingly available on the Web. While using biomedical Linked Data is crucial, life-science researchers may have difficulties using SPARQL language. Interfaces based on Natural Language question-answering are recognized to be suitable for querying knowledge bases. In this paper, we propose a method for translating natural language questions into SPARQL queries. We use Natural Language Processing tools, semantic resources and the RDF triples description. We designed a four-step method which linguistically and semantically annotates the question, performs an abstraction of the question, then builds a representation of the SPARQL query and finally generates the query. The method is designed on 50 questions over 3 biomedical knowledge bases used in the task 2 of the QALD-4 challenge framework and evaluated on 27 new questions. It achieves good performance with 0.78 F-measure on the test set. The method for translating questions into SPARQL queries is implemented as a Perl module and is available at http://search.cpan.org/\texttt{~thhamon/RDF-NLP-SPARQLQuery/}.

Keywords: Natural Language Processing, SPARQL, biomedical domain, semantic resources, frames

1 Introduction

The knowledge bases (KB) recording biomedical knowledge are becoming increasingly available on the Web. Such life-science bases usually focus on a specific type of biomedical information: chemical, pharmacological and target information on drugs in Drugbank \cite{drugbank}, clinical studies in ClinicalTrials.gov\textsuperscript{1}, drugs and their side effects in Sider \cite{sider}, etc.

Nowadays, the connections between these KB are crucial for obtaining a more global and comprehensive view on the links between different biomedical

\textsuperscript{1} http://clinicaltrials.gov/
components. They are also required for inducing and producing new knowledge from the already available data. Particularly, the creation of fine-grained links between the existing KB related to drugs is a great challenge that is being addressed by the project Linked Open Drug Data (LODD) for instance. The knowledge recorded in the KB and dataset interlinks are represented as RDF triples, on the basis of which the linked data can then be queried through a SPARQL end-point. However, typical users of this knowledge, such as physicians, life-science researchers or even patients, cannot manage the syntactic and semantic requirements of the SPARQL language neither can they manage the structure of various KB. This situation impedes the efficient use of KB and the retrieval of useful information. Therefore, it is important to design friendly interfaces that mediate the technical and semantic complexity of the task and provide simple approaches for querying the KB. The main challenge is then to design the optimal methodology for an easy and reproducible rewriting of natural language questions into SPARQL queries. In the following, the term question means the natural language expressions uttered by human users to formulate their information need, and the term query designates the same expression formalised in the SPARQL syntax and semantics.

We present in this paper a novel method to translate natural language questions into SPARQL queries over biomedical Linked Data. The method is based on the use of Natural Language Processing (NLP) tools and resources to enrich question with linguistic and semantic information. Questions are then translated into SPARQL with a rule-based approach. We design our approach on the 50 questions proposed by the task 2, Biomedical question answering over interlinked data, of the QALD-4 challenge, and evaluate it on 27 newly defined questions. A sample of the new test set is given at Sect. 5.1. We work with three KB (Drugbank, Diseasome, and Sider described in Sect. 4). The paper is structured as follows. Section 2 presents the related work. We describe the proposed method in Sect. 3 and, then, the semantic resources available and developed for enriching the questions in Sect. 4. The evaluation of the method is presented in Sect. 5.

2 Related Work

Querying Linked Data requires the definition of end user interfaces which hide the underlying structure of the KB as well as the SPARQL syntax. While [6] identify three ways for querying Linked Data (Knowledge-Based Specific Interface, Graphical Query Builder and Question-Answering System), it has been recognized that Natural Language interfaces are the most suitable. Thus, the authors have demonstrated that, for querying the KB and the Semantic Web

2 http://www.w3.org/wiki/HCLSIG/LODD
3 http://greententacle.techfak.uni-bielefeld.de/~cunger/qald/index.php?x=task2&q=4
4 The new set with 27 questions is available at the following URL: http://perso.limsi.fr/hamon/Files/QALD/qald-4 biomedical additional test.xml
data, the use of full and standard sentences is preferred to the use of keywords, menus or graphs [6].

Also, another work [9] proposes a Question-Answering system (AutoSPARQL) based on active supervised machine learning independent from the KB: the SPARQL query model is learnt from Natural Language questions. The authors report that 50 questions are successfully processed with the system.

Most of existing approaches usually rely on patterns or templates for translating the natural language questions into SPARQL queries. Thus, modular patterns are designed to parse questions [11]. Their main purpose is to model the questions for the SPARQL query generation where first keywords are detected and then relations between them are identified. The method is tested on 160 movie-related questions. The existing model requires four query patterns instead of twelve patterns proposed in a previous work.

[14] observe that Question-Answering systems offer a good compromise between expressivity and intuitiveness, and also propose a template-based system relying on NLP tools and semantic resources for processing natural language questions. The application of the system on 50 questions from DBpedia proposed by the QALD-2 challenge gives competitive results with 0.62 average F-measure obtained with 39 questions (average recall is 0.63 and average precision is 0.61), but shows a low coverage (11 questions are not covered by the templates).

Up to now the design of friendly user interfaces is mainly addressed for general KB [4][7]. However, as domain-specific Linked Data are more and more available, the studies on these data begin to show. Thus, we can mention the work which aims at translating medical questions issued from a journal into SPARQL queries [1]. The method combines the SVM machine learning-based approach to extract the characteristics of the questions (named entities, relations) with patterns to generate the SPARQL queries. The evaluation is carried out on 100 questions and the corresponding queries tested on clinical documents. The method achieves a 0.62 precision when querying the documents. Recently, the Question Answering over Linked Data (QALD-4) challenge proposes a task\(^5\) dedicated to the retrieval of precise biomedical information in linked KB with questions expressed in natural language.

Our approach is close to the one proposed by [14]: we use NLP tools in order to linguistically enrich the questions. The main difference is that we use information issued from the Linked Data resources to semantically annotate the questions and to define the frames (i.e., linguistic representations of the RDF schema) in order to model and to build the SPARQL queries.

3 Question translation into SPARQL Query

To translate natural language questions into SPARQL queries, we design a four-step rule-based method, that relies on NLP methods, semantic resources and the RDF triple description (see Fig. 1). Natural language questions are enriched

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\(^5\) Biomedical question answering over interlinked data, http://greententacle.techfak.uni-bielefeld.de/~cunger/qald/index.php?x=task2&q=4
This information is used to abstract the questions and to identify relevant information (Sect. 3.2), to build the corresponding SPARQL query representations (Sect. 3.3) and to generate the SPARQL queries (Sect. 3.4). The same process is used for translating questions provided by training and test sets. The method for translating questions into SPARQL queries is implemented as a Perl module and is available at http://search.cpan.org/˜thhamon/RDF-NLP-SPARQLQuery/.

Fig. 1. The global architecture of the system (the processing steps are in yellow, the resources in blue). Yellow boxes represent the processing steps which are further detailed in figures 2 to 6. Blue boxes describe the resource used for the processing of the questions and queries.
3.1 Linguistic and Semantic Annotation of Questions

The annotation step aims at associating a linguistic and semantic description to the words and terms of the questions (see Fig. 2).

Fig. 2. Linguistic and semantic annotation process. Yellow boxes represent the steps of the linguistic and semantic analysis of questions. Blue boxes indicate the resources used for the semantic annotation.

First, numerical values (such as numbers and solubility values) are identified with a named entity recognizer. Then, the parsing of the questions consists in: word segmentation, part-of-speech tagging and lemmatization of words with TreeTagger [12]. Semantic entities, i.e. terms with associated semantic types representing their meaning, are identified with the TermTagger Perl module. This term recognition relies on semantic resources (see Sect. 4) to recognize semantic entities such as disease names, side effects, etc. However, because the semantic resources often suffer from low coverage [3, 10], we also use the term extractor YATEA7 [2] to improve the coverage of our method. YATEA performs shallow parsing of the POS-tagged and lemmatized text by chunking it according to syntactic frontiers (pronouns, conjugated verbs, typographic marks, etc.) in order to identify noun phrases. Then, parsing patterns are recursively applied and provide parsed terminological entities, usually noun phrases relevant for the targeted domain. These parsing patterns have been manually defined during previous

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6 http://search.cpan.org/~thhamon/Alvis-TermTagger/
7 http://search.cpan.org/~thhamon/Lingua-YaTeA/
work. They take into account the morpho-syntactic variation and reflect basic syntactic dependencies within terminological entities. Each term is represented as a syntactic tree, and sub-terms are also considered as terms in the current configuration. No semantic types are associated with the terms extracted by YATEA. Figure 3 illustrates the linguistic and semantic annotation of questions.

Fig. 3. Example of question pre-processing (question #22 of the QALD-4 test set). The blue boxes represent the word and semantic entities. The subscript texts are the Part-of-Speech tags and the bracketed subscript texts are the semantic types associated with the semantic entities.

3.2 Question Abstraction

The question abstraction step aims at identifying the relevant elements within questions and at building the representation of these elements (see Fig. 4). It relies on the linguistic and semantic annotation associated with the question words in the previous step.

Fig. 4. Question abstraction process. Yellow boxes represent the abstraction steps of questions. Blue boxes indicate the resources used.
Before the identification of relevant elements, the annotations are post-processed in order to disambiguate the semantic annotations. Indeed, the annotated semantic entities may receive conflicting or concurrent semantic types, while the post-processing permits to select those entities and semantic types that may be useful for the next steps. As part of this post-processing, we keep larger terms which do not include other semantic entities. Also, we manually defined rewriting rules on the training set in order to adjust (modify or delete) the semantic types associated with a given entity according to the context. Other rules may also modify or delete the entity itself according to its context. For instance, the semantic entity interaction has to be rewritten into interactionDrug1 if its context contains mention of drugs, but it has to be rewritten into foodInteraction if its context contains a term with the semantic type food. On the whole, we defined 44 contextual rewriting rules based on the vocabulary used in questions from the training set and on the documentation of KB, mainly the one from Drugbank\textsuperscript{8}. Besides the rewriting rules, an additional disambiguation of the annotations is also performed during the SPARQL query construction step when the arguments of the predicate or the question topic are connected because they share the same semantics.

For performing the question abstraction, we identify information related to the query structure:

1. Definition of the Result form: the question is scanned for identifying words expressing the negation, e.g. *no*, and its scope, the aggregation operation on the results, e.g. *number* for *count*, *mean* for *avg* or *higher* for *max*, and specific result forms such as boolean queries (*ASK*). Information related to negation or conjunction marks, aggregation operators, and requirement on specific result forms is recorded and will be used at the end of the *query construction* step or during the *query generation* step. In the example from figure 3, no such information is found.

2. Identification of the Question topic: the first semantic entity with a given expected semantic type is considered to be the question topic (this assumption is also verified on the test set). The expected semantic types are those provided by the RDF subjects and objects issued from the resources. This information will be used during the *query construction* step. The question topic of our example is identified as *drug*.

3. Identification of Predicate and Argument: we use linguistic representations of the RDF schema i.e. frames which contain one predicate and at least two elements with associated semantic types. In that respect, the potential predicates, subjects and objects of frames are identified among the semantic entities and recorded in a table (entries are the semantic types of the elements and refer to linguistic, semantic and SPARQL information associated with these elements). Subjects and objects are fully described in the table with inflected and lemmatized forms of words or terms, the corresponding SPARQL types and indicators on their use as object or subject of a predicate.

\textsuperscript{8} http://www.drugbank.ca/documentation
Concerning the predicates, only the semantic types of their arguments are instantiated. Subjects and objects can be URI, RDF typed literals (numerical values or strings) and extracted terms (these are considered as elements of regular expressions). In the example from figure 3, the predicate state with the expected arguments drugbank/drugs and Gas/String is recognized.

4. Scope of negation and conjunctions: the argument and predicate in the neighbourhood of negation and conjunctions are identified. These elements are recorded as negated or coordinated.

Figure 5 presents a graphical representation and abstraction of question 22 from the QALD-4 test set.

![Graphical representation of question abstraction](image)

**Question Topic**

**Aggregation operator:**

**Question topic:** drug (drugbank/drugs)

**Predicates:**

<table>
<thead>
<tr>
<th>Frame</th>
</tr>
</thead>
<tbody>
<tr>
<td>drugbank/drugs</td>
</tr>
<tr>
<td>state Gas/String</td>
</tr>
</tbody>
</table>

**Arguments:**

<table>
<thead>
<tr>
<th>Semantic type</th>
<th>Word</th>
<th>SPARQL type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas/String</td>
<td>gaseous</td>
<td>STRING(Gas)</td>
</tr>
</tbody>
</table>

**Fig. 5.** Example of question abstraction (question#22 of the QALD-4 test set)

3.3 Query Construction

The objective of the query construction step is to connect previously identified elements and to build a representation of the SPARQL graph pattern (introduced by the keyword WHERE). Figure 6 presents the architecture of the query construction. Thus, the predicate arguments are instantiated by URIs associated with the subjects, objects, variables, and numerical values or by strings. For each question, we perform several connections:

1. The question topic is connected to the predicate(s). A variable is associated with the question topic and the predicate arguments that match the semantic type of the question topic. Note that at the end of this stage, the question topic may remain non-associated to any predicate. In the example from
Fig. 6. Query construction process. Yellow boxes represent the construction steps of queries. Blue boxes indicate the resource used.

figure 3, the variable "v0" represents the association between the question topic and the subject (with the expected type drugbank/drugs) of the predicate state.

2. The predicate arguments are connected to subjects and objects identified during the question abstraction: they concern elements referring to URIs. Moreover, each predicate within the conjunction scope is duplicated and its arguments are also connected to the subject and objects if needed;

3. The predicates are connected between them through their subjects and objects. The connection between two predicates is then represented by a variable;

4. The predicates from different datasets are connected. We use the sameAs description to identify URIs referring to the same element. New variables are defined to connect two predicates;

5. The remaining question topic is connected to arguments of the sameAs predicate;
6. The arguments corresponding to the **string** type are connected with the extracted terms. We assume these arguments will be related to the string matching operator **REGEX**. Thus, the terms are considered as string expressions.

At this point, the predicate arguments which remain unassociated are replaced by new variables in order to avoid empty literals. Finally, the negation operator is processed: the predicates are marked as negated and the arguments within the negation scope are included in a new predicate **rdf:type** if required.

At this stage, each question is fully translated into a representation of the SPARQL query. Figure 7 illustrates the construction of the query corresponding to question 22 of the QALD-4 test set.

![Diagram of query construction](image)

**Fig. 7.** Example of query construction (question #22 of the QALD-4 test set)

### 3.4 Query Generation

The SPARQL query representation built during the *query construction* step is used to generate the SPARQL query string. The process is composed of two parts:

1. The generation of the result form which takes into account the expected type of the result form (**ASK** or **SELECT**), the presence of aggregation operators and the variable associated with the question topic;
2. The generation of the graph pattern. This part consists in the generation of strings for representing each RDF triple and the filtering if the predicates are negated. But when aggregation operators are used, we also need to recursively generate sub-queries for computing the subsets of expressions,
before their aggregation. In the example from figure 3, the predicate `state` is replaced by the corresponding URI and its object is replaced by the string `gas`.

The SPARQL queries have been submitted to a SPARQL end-point\(^9\) and answers are collected for the evaluation. Figure 8 presents the generated query which corresponds to question 22 of the QALD-4 test set.

\[
\text{SELECT DISTINCT } ?v0 \\
\text{WHERE } \{ \\
?v0 <http://www4.wiwiss.fu-berlin.de/drugbank/resource/drugbank/state> "Gas". \\
\}
\]

**Fig. 8.** Example of the query generation (question#22 of the QALD-4 test set)

### 4 Definition of the Semantic Resources

The method described above relies on the existing biomedical resources that provide information on the semantic entities (Sect. 4.1), but also on additional resources specifically collected and built to support the method (Sect. 4.2).

#### 4.1 Domain-specific Resources

To process the set of questions from QALD-4, we used three biomedical resources:

- **Drugbank**\(^{10}\) is dedicated to drugs [15]. It merges chemical, pharmacological and pharmaceutical information from other available KB. We exploited the documentation\(^{11}\) of this resource to define the rewriting rules and regular expressions for the named entity recognition.

- **Diseasome**\(^{12}\) is dedicated to diseases and genes linked among them by known disorder/gene associations [5]. It provides a single framework with all known phenotypes and disease gene associations, indicating the common genetic origin of many diseases. We exploited the RDF triples and the documentation of the resource to define the rewriting rules.

- **Sider**\(^{13}\) is dedicated to adverse drug effects [8]. It contains information on marketed medicines and their recorded adverse drug reactions. Information

\(^9\) For our experiments, we use the SPARQL end-point provided by the QALD-4 challenge [vtentacle.techfak.uni-bielefeld.de:443/sparql](http://vtentacle.techfak.uni-bielefeld.de:443/sparql)

\(^{10}\) [http://www.drugbank.ca](http://www.drugbank.ca)

\(^{11}\) [http://www.drugbank.ca/documentation](http://www.drugbank.ca/documentation)

\(^{12}\) [http://diseasome.eu](http://diseasome.eu)

\(^{13}\) [http://sideeffects.embl.de](http://sideeffects.embl.de)
is extracted from public documents and package inserts. The available information includes side effect frequency, drug and side effect classifications as well as links to other data, such as drug-target relations.

The content of each resource is provided in specific format: RDF triples subject predicate object. In that respect, we also exploit their RDF schema to define frames (see below).

4.2 Additional Resources for the Question Annotation

On the basis of the RDF triples, frames are built from the RDF schemas in which the RDF predicate is the frame predicate, and subject and object of the RDF triples are the frame elements. This also includes the OWL sameAs triples. Several types of frame entities are isolated:

- As indicated, subject, object and predicate become semantic entities. They may occur in questions: in this way, the frames are the main resources for rewriting questions in queries.
- The vocabulary specific to questions is also built. It covers for instance aggregation operators, negation and types of questions.
- RDF literals, issued from named entity recognizer or term extractor, complete the resources. The RDF literals are detected with specifically designed automata that may rely on the source knowledge base documentation.

These entities are associated with the expected semantic types, which allows creating the queries and rewriting the RDF triples into SPARQL queries. In that respect, we can consider IRIs, strings, common datatypes or regular expressions when literals are expected.

Most of the entities are considered and processed through their semantic types, although some ambiguous entities (e.g. interaction or class) are considered atomically. For these, the rewriting rules are applied contextually to generate the semantic entities corresponding to the frames (see Sect. 3.2). When using the queries, the semantic types become variables and are used for connecting the edges of queries.

5 Experiments and Results

5.1 Training and Test question set

As the complexity of the QALD-4 training and test sets was unbalanced (e.g. the QALD-4 training set does not propose questions requiring the use of aggregation operators), we use the 50 questions of the QALD-4 training and test sets for fitting our system. Besides, similarly to the usual layout of challenge data, we design 27 new questions used for the evaluation. The questions of the new test are similar to the QALD-4 questions but may differ according to the semantic entities or the involved predicates. Our method is applied to this new test set without additional adaptations. Figure 9 presents a sample of questions from the new test set.
Which foods does fluvoxamine interact with?
Are there drugs that target the Probable arabinosyltransferase A?
Which genes are associated with subtypes of rheumatoid arthritis?
Which disease has the highest degree?
Which targets are involved in immune function?

5.2 Evaluation Metrics

The generated SPARQL queries are evaluated through their answers with following macro-measures [13]:

\[
M\text{-}precision = \frac{\sum_{i=1}^{\vert q \vert} TP(q)}{\frac{TP(q) + FP(q)}{\vert q \vert}} \quad M\text{-}recall = \frac{\sum_{i=1}^{\vert q \vert} TP(q)}{\frac{TP(q) + FN(q)}{\vert q \vert}}
\]

\[
M\text{-}F\text{-}measure = \frac{2 \times M\text{-}precision \times M\text{-}recall}{M\text{-}precision + M\text{-}recall}
\]

where \(TP(q)\) are the correct answers, \(FP(q)\) are the wrong answers and \(FN(q)\) are the missing answers for the question \(q\).

The use of macro-measures equally considers all the questions independently of the number of expected answers to the SPARQL queries.

5.3 Global Results

Table 1 presents the overall results obtained on the training and test sets. On the test set, the macro-F-measure is 0.78 with 0.81 precision and 0.76 recall while on the training set, the macro-F-measure is 0.86 with 0.84 precision and 0.87 recall. Each question is processed in less than 2 seconds on a standard computer (2.7GHz dual-core CPU and 4 Gb of memory). Most of the computing time is spent for the linguistic and semantic annotation of the questions.

<table>
<thead>
<tr>
<th>Query set</th>
<th>Training (50 Q)</th>
<th>Test (27 Q)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Correct Queries</td>
<td>39</td>
<td>20</td>
</tr>
<tr>
<td>(M\text{-}precision)</td>
<td>0.84</td>
<td>0.81</td>
</tr>
<tr>
<td>(M\text{-}recall)</td>
<td>0.87</td>
<td>0.76</td>
</tr>
<tr>
<td>(M\text{-}F\text{-}measure)</td>
<td>0.86</td>
<td>0.78</td>
</tr>
</tbody>
</table>

Our method always proposes syntactically correct SPARQL queries for all natural language questions: 18 questions provide the exact expected answers,
two questions return partial answers, other questions return erroneous answers. On the training, 39 SPARQL queries (out of the 50 questions) are semantically correct and provide the expected answers. We can also observe that 6 questions return partial answers, and 5 questions return no answers.

**Error Analysis.** An analysis of the erroneous or partial answers shows that most of the errors are due to (i) the encoding of the `sameAs` predicate in the resources and (ii) the management of ambiguities in the questions.

Regarding the first type of errors, we observe that, although our method generates the correct SPARQL query, the SPARQL end-point does not return the expected answers. Besides, we observe that by switching the arguments of the `sameAs` predicate in the queries, the expected answers are returned. In that respect, we consider that the instances of this predicate do not encode the expected reflexivity of this relation in the source KB while our method assumes that the `sameAs` predicate is reflexive by definition.

As for the errors due to ambiguity, they mainly concern:

- the annotation of semantic entities. For instance, in the question *Which genes are associated with breast cancer?*, *breast cancer* is correctly annotated, while the reference assumes it concerns the semantic entity *Breast cancer-1*.
- the intended meaning of the terms in the questions. Semantic entities mentioned in some questions may refer to specific entities while in other questions they refer to the general ones. For instance, the semantic entity *anemia* in *What are enzymes of drugs used for anemia?* refers to all types of anemia (*Hypercholanemia*, *Hemolytic anemia*, *Aplastic anemia*, etc.), and not to the elements that contain the label *anemia*.

These two main problems could be solved by using regular expressions in SPARQL graphs rather than URIs. However, we must test the influence of this modification on each query.

Other erroneous answers happen during the question abstraction step when the question topics are wrongly identified or when the contextual rewriting rules are not applied. Errors also occur during the query construction step: the method may abusively connect predicate arguments and semantic entities or, on contrary, it may not consider all the identified semantic entities. Further investigations have to be carried out to solve these limitations.

Besides, during the design of queries, we had difficulties to express some constraints in SPARQL. For instance, the question *Which approved drugs interact with calcium supplements?* requires to define a regular expression with the term *calcium supplement* while this term is only mentioned in conjunction with other supplements (e.g. *Do not take calcium, aluminum, magnesium or Iron supplements within 2 hours of taking this medication.*). We assume that solving this difficulty requires a more sophisticated NLP processing of the textual elements of the RDF triples (parsing of the RDF textual elements, named entity and term recognition, identification of discontinuous terms and term variants, etc.).
Other limitations concern the updating of the KB and the change of their structure. In the former case, it is only required to rebuild the semantic resources used for identifying the semantic entities. In the latter case, the frames must be regenerated. This is an ongoing research work. Moreover, the addition of new resources such as Dailymed\textsuperscript{14} is also related to these two problems.

6 Conclusion

We proposed a rule-based method to translate natural language questions into SPARQL queries. The method relies on linguistic and semantic annotations of questions with NLP methods, semantic resources and the RDF triples description. We designed our approach on 50 biomedical questions proposed by the QALD-4 challenge, and tested it on 27 newly created questions. The method achieves good performance with 0.78 F-measure on the set of 27 questions.

Further work aims at addressing the limitations of our current method including the management of term ambiguity, the question abstraction, and the query construction. Moreover, to avoid the manual definition of the dedicated resources required by our approach (frames, specific vocabulary and rewriting rules), we plan to investigate how to automatically build these dedicated resources from the RDF schemas of the Linked Data set. It will also facilitate the integration of other biomedical resources such as Dailymed or RxNorm, and the use of our method in text mining applications.

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\textsuperscript{14} http://dailymed.nlm.nih.gov/


