**BioMed Xplorer**

*Exploring (Bio)Medical Knowledge using Linked Data*

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**Keywords:** BioMed Xplorer, Disease Related Information, Semantic Web, Knowledge Base Ontology, Visualization, Provenance Data, Medical Knowledge, External Data Source, RDF, Graph, Knowledge Exploration.

**Abstract:** Developing an effective model for predicting risks of a disease requires exploration of a vast body of (bio)medical knowledge. Furthermore, the continuous growth of this body of knowledge poses extra challenges. Numerous research has attempted to address these issues through developing a variety of approaches and support tools. Most of these tools however, do not sufficiently address the needed dynamism, lack intuitiveness in their use, and present a rather scarce amount of information usually obtained from a single source. This research aims to address the aforementioned gaps through the development of a dynamic model for (bio)medical knowledge, represented as a network of interrelated (bio)medical concepts, and integrating disperse sources. To this end, this paper introduces BioMed Xplorer, presenting a model and a tool that enables researchers to explore biomedical knowledge, organized in an information graph, through a user friendly and intuitive interface. Furthermore, BioMed Xplorer provides concept related information from a multitude of sources, while also preserving and presenting their provenance data. For this purpose a RDF knowledge base has been created based on a core ontology which we have introduced. Results are further experimented with and validated by some domain experts and are contrasted against the state of the art.

1 **INTRODUCTION AND RESEARCH APPROACH**

The (bio)medical field is vast and dynamic, with knowledge developing rapidly as a result of continuously ongoing research. Within this field, extensive research is conducted into identifying risk factors of diseases as well as assessing their effect on the presence and associated severity of a disease. The available knowledge from this research on risk factors enables researchers to develop models for risk prediction, which might be used by practitioners to assess someone’s risk on developing a particular disease. Conventional methods for developing such models for risk prediction would involve identifying the risk factors and their effects from the ever-evolving body of (bio)medical knowledge. Achieving this aim would thus involve checking vast amount of scientific publications for relevant statements regarding factors that might affect a disease. This, however, is a cumbersome and costly activity, especially when considering the fact that the U.S. National Library of Medicine’s (NLM) bibliographic database MEDLINE, as of today, contains over 22 million citations, over 750,000 of which were added in 2014 (U.S. National Library of Medicine, 2015b), and that these numbers have grown exponentially (Hunter and Cohen, 2006). As a result of the sheer size and continuous growth of the body of (bio)medical knowledge, exploration of this body of knowledge, as well as finding the relevant knowledge for inclusion in models for risk prediction, becomes increasingly challenging for researchers, potentially causing an information overload (Hunter and Cohen, 2006; Lu, 2011).

Numerous researchers have reckoned this problem and have attempted to address it from different perspectives (Lu, 2011; Cohen and Hersh, 2005), for example through the development of comprehensive visualizations that represent knowledge extracted from (bio)medical publications (Plake et al., 2006; Rebholz-Schuhmann et al., 2007; Tao et al., 2005; Kilicoglu et al., 2008; Bodenreider, 2000).

Even though the visual nature of these knowledge representation and visualization tools provides them with great expressive power, four common shortcomings can be identified among them, being: i) their restricted scope, focusing just on a particular sub-domain of the (bio)medical field, ii) their lack of in-
tuitiveness and rather sharp learning curve, iii) the scarce amount of information represented, solely limited to names and identifiers, lacking descriptions or definitions, while these are available externally, and iv) the fact that they are either no longer active (AliBaba, PGviewer), or do not work properly (EBIMed). From these shortcomings it thus becomes clear that there is a need for a meaningful representation of the available (bio)medical knowledge that: a) is intuitive, and b) represents information from multiple sources. As such the following research question can be conceived:

*Can we develop a model of the (bio)medical knowledge that is available from large, disperse, heterogeneous, and dynamic sources across the web?*

In order to address this research question a five-phase research approach has been designed, consisting of the following phases: 1) State of the Art Assessment, 2) Data Source Characterization and Selection, 3) Data Preprocessing and Ontology Design, 4) Data Interlinking and Fusion with external sources, and 5) Model Visualization.

Completion of these five phases delivers a system with an architecture that is shown in Figure 1. As one might notice, the architecture consists of four core modules, each of which corresponds to one of the major design and development stages. The components of these modules will be gradually defined in the corresponding sections, as such fully describing the system architecture.

The remainder of this paper is structured according to the five phases that were outlined above, with each section elaborately discussing a particular phase of the research. In section 2, the characterization and selection of data sources for inclusion in the model is described. This is followed by a discussion of the data preprocessing and ontology design in section 3, whereas section 4 covers the fusion and interlinking of the data. The visualization of the model is subsequently discussed in section 5, while the work is validated in section 6. Finally, section 7 concludes the paper.

## 2 DATA SOURCE CHARACTERIZATION AND SELECTION

Central to the development of a model is the data that eventually will be represented in the model and thus needs to be utilized for building and populating the model. With the research question in mind, the identification, and subsequent selection, of data sources that provide disease related information, pertaining to, for example, symptoms, inheritability, and genetics of a disease, thus are the first key steps in the development process of the disease related information model. A search for disease related information results in a wide variety of structured (i.e. standardized terminologies or vocabularies, ontologies, and databases) and unstructured (e.g. websites (U.S. National Library of Medicine, 2015c; WebMD, LLC, 2015)) data sources. Data from unstructured sources requires conversion to a structured format, for example using Natural Language Processing (NLP) techniques, and thus cannot be directly incorporated into the disease related information model. As a result we have decided to only incorporate structured data sources. It is essential to designate a primary data source for the development of the disease model as the available structured data sources for disease related information overlap in terms of covering the same information in different formats and presentations.

A disease model that is represented in a network-like format consists of two components, namely concepts and relationships among these concepts. Concepts can be sourced from standardized terminologies, or from ontologies. Some well-known terminologies in the biomedical field are the *International Classification of Diseases (ICD)* (World Health Organization, 2015), *Medical Subjects Headings (MeSH)* (U.S. National Library of Medicine, 2015a), and *Systematized Nomenclature of Medicine Clinical Terms (SNOMED-CT)* (International Health Terminology Standards Development Organisation, 2015), whereas the *National Cancer Institute Thesaurus (NCIt)* (U.S. National Cancer Institute, 2015b), the *Disease Ontology* (Institute for Genome Sciences - University of Maryland School of Medicine, 2015), and the *Gene Ontology* (Ashburner et al., 2000) are among the frequently used ontologies within the biomedical field. Instead of sourcing concepts from one or multiple individual terminologies, one can source the concepts from the *Unified Medical Language System (UMLS) Metathesaurus* (U.S. National Library of Medicine, 2015e) or the *National Cancer Institute Metathesaurus (NCIm)* (U.S. National Cancer Institute, 2015a), both of which integrate, among many others, the aforementioned sources into a single terminology. Using these metathesauri provides the opportunity of broadening the scope of the concepts that are covered and, as such, expanding the knowledge base of the model by using concepts represented in the majority of separate terminologies. Therefore, the use of either the UMLS or NCIm to define concepts in the model is preferred over the use of separate terminologies.
Relationships, on the other hand, can also be sourced from the UMLS and NCIm. More extensive relationships, however, can be obtained from the Online Mendelian Inheritance in Man (OMIM) database (Johns Hopkins University, 2015), MalaCards (Weizmann Institute of Science, 2015), or SemMedDB (Kilicoglu et al., 2012). Considering the overarching aim of this research in aiding (bio)medical researchers in their knowledge explorations efforts, and due to the fact that (bio)medical knowledge originating from peer-reviewed literature is considered trustworthy and rich, relationships directly derived from (bio)medical literature are selected as the primary relationships in the model. To this end, SemMedDB is thus selected as the primary source, presenting disease related information, for incorporation into the developed model. This choice is further motivated by the fact that SemMedDB is considerably larger (containing over 70 million statements) than the other identified sources containing disease related information. Finally, the broad scope, covering terms across the entire biomedical domain, also played a role in the choice for SemMedDB.

3 DATA PREPROCESSING

Due to the large amounts of heterogeneous and dynamic information that is nowadays available across a multitude of sources, relational databases are considered to be less than ideal for storing and instantiating knowledge representations of information with such nature (Hendler, 2014). Linked data, on the other hand, provides a promising solution to this issue as it is able to cope with such large amounts of dynamic and heterogeneous information (Berners-Lee et al., 2001). To this end we therefore aim to develop our model using Semantic Web technologies. According to (Berners-Lee et al., 2001; Antoniou and Van Harmelen, 2004) the Semantic Web consists of three main components, being i) labeled graphs that encode meaning by representing concepts and the relations among them, and are usually expressed as (subject-predicate-object) triples in RDF, ii) Uniform Resource Identifiers (URIs) to uniquely identify the items in the datasets as well as to assert meaning, which is reflected in the design of RDF, and iii) ontologies to formally define the relations that can exist among data items. In order to develop our model using the Semantic Web, the existence of these three components needs to be ensured. Processing the data in SemMedDB such that these three components exist, is therefore the main aim of the preprocessing stage.

3.1 Ontology Design

In order to be able to generate labeled graphs from a relational database, such as SemMedDB, and to ensure the use of URIs, an ontology needs to be developed that represents the desired data structure of these graphs. This ontology should define the data items, as well as the relations among them, that are aimed to be represented. Considering that the planned model should represent the statements, and their provenance data, in SemMedDB as a RDF graph, it is key for the ontology to closely resemble SemMedDB’s database design. Prior work has been conducted in this area by (Tao et al., 2012). In their work, (Tao et al., 2012) aimed to optimize the organization and representation of Semantic MEDLINE data (SemMedDB) for translational science studies by reducing redundancy.
through the application of Semantic Web technologies. This is achieved by representing the concepts and associations in SemMedDB as RDF. Despite successfully decreasing the redundancy of the information in SemMedDB, two shortcomings can be identified in the ontology that was developed by (Tao et al., 2012). First of all, the ontology represents a limited amount of information compared to the information that is available in SemMedDB. This, in turn, impedes the ability to incorporate external resources into the model since among the information from SemMedDB that is omitted are unique identifiers that are required to retrieve the appropriate entities from these external sources. The second shortcoming is the lack of reuse of terms defined in existing vocabularies, which is one of the founding principles of the Semantic Web (Shadbolt et al., 2006). In (Tao et al., 2012), the developed ontology defines all terms used, whereas equivalent classes might already exist in other vocabularies in the Web of Data. Such reuse would facilitate the linking of data to a Web of Data, which is an overarching goal of the Semantic Web (Berners-Lee et al., 2001).

Despite the limitations of the ontology developed in (Tao et al., 2012), this ontology is considered as a starting point as well as an opportunity to improve on and extend upon. To this extent, the BioMed Xplorer Ontology is developed that addresses the identified shortcomings by representing most of the information contained within SemMedDB, as well as by reusing as much terms from existing vocabularies or ontologies as possible. The BioMed Xplorer Ontology is developed in the Web Ontology Language (OWL2) and is published on a Persistent Uniform Resource Locator (PURL) (Weibel et al., 1996) domain. Such a locator allows the underlying Web address of a resource to change while not affecting the availability of the systems that depend on this resource. The BioMed Xplorer Ontology is shown in Figure 2.

RDF Reification. Considering that the provenance data in SemMedDB applies to statements as a whole, reification is necessary in order to represent this provenance data in the ontology. (Tao et al., 2012) also recognized this need, however, they did not use the RDF Reification vocabulary as outlined in (World Wide Web Consortium et al., 2014). The BioMed Xplorer Ontology on the other hand implements the RDF reification vocabulary.

As the statements contained in SemMedDB relate two UMLS concepts to each other, both the subject and object of an rdf:Statement instance are modelled as instances of a Concept class. The concepts are related to each other through one of 58, relationships that are identified by SemRep (U.S. National Library of Medicine, 2015d). The predicate of an rdf:Statement instance therefore is modelled as one of 58 instances of the Relation class. This set of relations consists of two disjunctive subsets, with one subset containing 31 relationships derived from the UMLS Semantic Network, such as "causes", and the other subset containing the remaining 27 relationships, which are negated versions of the relationships in the first subset, such as "neg-causes", referring to "does not causes" (Kilicoglu et al., 2012). Relationships belonging to the negated subset are prefixed with "NEG", whereas all other relationships are considered to belong to the subset of affirmed relationships. These two subsets of relations are represented in the BioMed Xplorer Ontology as two subclasses of the Relation class, being the AffirmedRelation and the NegatedRelation classes respectively.

The provenance data in SemMedDB includes both the sentences from which a statement is derived, as well as the publications in which these sentences occur. Reification of the statements enables the assertion of this provenance data to their respective statements. To this end, sentences are represented as instances of the Sentence class, which are related to the rdf:Statement class through a derivedFrom property. The articles in which these statements and sentences are contained, are represented as instances of an Articles class, which are related to the rdf:Statement class through a source property. Furthermore, sentences are related to articles through the partOf property, indicating that a sentence is part of an academic article. In addition to the object properties, relating classes to each other, discussed in this section, a number of datatype properties, associating data values (such as identifiers) to classes, are asserted to each of the classes in the BioMed Xplorer Ontology as well. Collectively, these properties aim to represent as much information from SemMedDB in the ontology as possible.

Vocabulary Reuse. The BioMed Xplorer Ontology aims to reuse as much existing classes and properties as possible. To this extent all elements of the ontology, which include the classes and both the object and datatype properties, except elements from the RDF or RDFS namespaces, have been checked for the presence of already defined equivalent concepts or properties in existing vocabularies. This has been accomplished by making use of the online RDF vocabulary search and lookup tool vocab.cc (Institute of Applied Informatics and Formal Description Meth-
ods - Karlsruhe Research Institute, 2015) that allows one to enter any term, and returns any classes and properties that (partially) match the term. In general the highest ranked term that corresponds to the role of the term in the BioMed Xplorer Ontology (e.g. class or property) is selected for reuse in the BioMed Xplorer. In the end, the search for existing terms lead to the incorporation of terms from three existing vocabularies, being i) the Bibliographic Ontology (D’Arcus and Giasson, 2015), ii) the Dublin Core Metadata Terms (Dublin Core Metadata Initiative (DCMI), 2015), and iii) the Simple Knowledge Organization System (World Wide Web Consortium, 2015).

4 DATA INTERLINKING & FUSION

The ontology developed in section 3.1 defines the desired data structure for the developed model. Generating the labeled graphs from the SQL in SemMedDB, however, requires a mapping that specifies how the data in the database is matched and converted to the appropriate class instances, properties, and property values specified in the ontology. Such a mapping can be developed using D2RQ, a declarative language for describing mappings between relational databases, RDF(S), and OWL ontologies (Bizer and Seaborne, 2004). The developed D2RQ mapping files have been made available online\(^2\). A mapping file enables RDF applications to access relational databases as virtual RDF graphs through the companion tool D2R Server (Bizer and Cyganiak, 2006). These virtual RDF graphs can subsequently be queried using the SPARQL protocol, with the D2RQ mapping translating the SPARQL queries to SQL queries, and translating the query results back to RDF. Both D2RQ and D2R are jointly available in the D2RQ Platform (Bizer and Seaborne, 2004). With the developed mapping file, the data in SemMedDB can be interlinked as RDF triples according to the specified ontology, as such surfacing and populating the actual disease related information model. Furthermore, the combination of the ontology and the use of RDF ensures the ability to link to the data in the information model from external datasets, through the URIs assigned to instances and properties.

In order to achieve complete data fusion with external sources, as such creating a truly Linked Data model, the data in the disease related information model should be linked to related entities or instances in external (RDF) data sources. This can be achieved by setting RDF links between the data in the model and these external data sources (Berners-Lee et al., 2009). One common way of setting such links between data sets is through the owl:sameAs property, which indicates that two linked individuals refer to the same thing (Dean et al., 2004). Establishing these links subsequently enables the incorporation of data from the external data sources into the disease related information model. Key to this data fusion process is the identification of external data sources containing instances that are equivalent to the instances in the developed model. The search for these data sources containing equivalent instances has been facilitated by searching the Linked Open Data cloud\(^3\) for unique standardized instance identifiers. Among these identifiers in SemMedDB are the UMLS Concept Unique Identifier (CUI), the Entrez-Gene ID, and the OMIM identifier for concepts, as well as the PubMed Identifier (PMID) for publications. The search of the

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\(^2\)The mapping files are available online from: https://goo.gl/1yD0WO

\(^3\)For details see http://lod-cloud.net/
Linked Open Data cloud for (bio)medical RDF data sources that represent either (bio)medical concepts, identified by one of the aforementioned identifiers, or publications, identified by the PubMed identifier, returned two main external data sources that could be fused with the data in SemMedDB: Linked Life Data (Momtchev et al., 2009), and Bio2RDF (Belleau et al., 2008).

5 BIOMED XPLORER UI

Assisting (bio)medical researchers in their knowledge exploration efforts can be achieved by enabling them to intuitively explore the body of (bio)medical knowledge. To this end it is therefore imperative to visualize the developed disease related information graph, representing this body of knowledge, that incorporates other disease related information gathered and aggregated from disperse sources across the Web. With this in mind three key requirements for the BioMed Xplorer UI can be imagined, being that it should: i) be usable and intuitive (e.g. supported by an appropriate visualization paradigm), ii) concisely represent provenance data (e.g. the publications as well as sentences from which statements are derived), and iii) represent information from multiple sources (e.g concept summaries and definitions). Based on these identified requirements, BioMed Xplorer has been developed and made available on the Web. The BioMed Xplorer UI supports the visualization of the information and has been developed in JavaScript in combination with the d3.js and jQuery libraries.

The data visualized by the BioMed Xplorer UI is obtained from BioMed Xplorer’s back-end, which consists of a Virtuoso triple store containing the disease related information model in RDF. This triple store provides a built-in SPARQL endpoint that can be queried by BioMed Xplorer using the SPARQL (Harris et al., 2013) protocol. Efficient query handling has been achieved by the development of a caching mechanism.

The developed user interface has three key features being: i) a graph-based visualization, ii) the exploration of concept information, and iii) the exploration and assessment of relationships. Each of these three features will be briefly discussed in the remainder of this section.

Exploring Concept Information. Within the BioMed Xplorer UI concept information can be explored through concept summaries, which can be opened by clicking on concepts, and concept overviews (as shown in Figure 4), which can be opened by choosing to show details in a concept summary. Concept information includes a wide range of information available from within the model as well as from external sources, such as Linked Life Data and Bio2RDF.

Exploring and Assessing Relationships. Relationships between concepts can be explored in the BioMed Xplorer UI through statement summaries, which can be opened by clicking on an edge, and statement overview, which can be opened by choosing to show details in a relationship summary. Within statement overviews, a wide range of statement information is available, as is shown in Figure 5. Among the available information is: the complete statement, two aggregates of the available provenance data, a brief overview of the source and target concepts of the relationship, the sentences from which the statement is derived at a publication level, as well as the details of the publications.

6 VALIDATION

Keeping the key roles of both the BioMed Xplorer Ontology, as the the foundation for the knowledge base, and the BioMed Xplorer UI, as the visualization of this knowledge base in mind, the validation of these two outcomes is imperative. This validation aims to assess whether the proposed solutions successfully address the already identified gap as well as how the proposed solutions measure against existing work. To this end, a two folded validation of both the ontology and the visualization has been conducted. In this regard a comparison to prior work has been conducted first, the results of which are shown in Tables 1 and 2 respectively. Secondly, an evaluation has been performed by 6 experts in the field. Results of this expert evaluation showed that both the BioMed Xplorer Ontology and the BioMed Xplorer UI successfully sat-
Figure 3: A screenshot from BioMed Xplorer UI and its graph-based visualization of (bio)medical knowledge, representing (bio)medical concepts as nodes and their interrelationships as edges.

Figure 4: BioMed Xplorer UI’s Concept Overview for “Malignant Neoplasms”.

isfy the identified requirements, with average grades of a 7.8 and 7.6 out of 10 respectively. Details of the expert evaluation of both the BioMed Xplorer Ontology and the BioMed Xplorer UI are provided in Tables 3 and 4.

Comparison to Related Work. There are five main knowledge representation and visualization tools identified that attempt to address similar challenges associated with exploring the body of (bio)medical knowledge through the representation and visualization of the knowledge contained within scientific publications. Among these tools are: AliBaba (Plake et al., 2006), EBIMed (Rebholz-Schuhmann et al., 2007), PGviewer (Tao et al., 2005), Semantic MEDLINE (Kilicoglu et al., 2008), and the Semantic Navigator (Bodenreider, 2000). Due to the close correspondence between the aims of these tools and the aims of our research, these five tools are considered as the base for comparison to BioMed Xplorer. A
Figure 5: BioMed Xplorer UI's Statement Overview for "Malignant Neoplasms" part of "Rattus Norvegicus".

Table 2: Comparison of BioMed Xplorer UI with five knowledge visualization tools developed in prior work.

<table>
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<th></th>
</tr>
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<tbody>
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<td>Scope</td>
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<td>Limited</td>
<td>Limited</td>
<td>Biomedical</td>
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<td>Biomedical</td>
</tr>
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<td>No</td>
<td>No</td>
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<td>Yes</td>
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<td>Visualization paradigm</td>
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<td>Tree</td>
<td>Graph</td>
<td>Graph</td>
<td>Graph</td>
</tr>
<tr>
<td>Concept categorization</td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Incorporation of links to external sources</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
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</tr>
<tr>
<td>Incorporation of data from external sources</td>
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<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Presentation of concept related information</td>
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<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Incorporation of provenance data</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Table 1: Comparison of BioMed Xplorer Ontology with the ontology developed by (Tao et al., 2012)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Tao et al., 2012</th>
<th>BioMed Xplorer Ontology</th>
</tr>
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<tr>
<td>RDF Reification</td>
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<td>Yes</td>
</tr>
<tr>
<td>Vocabulary reuse</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Links to external data sources</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Number of related data-items captured</td>
<td>4</td>
<td>17</td>
</tr>
<tr>
<td>Provenance data captured</td>
<td>Publications</td>
<td>Publications and sentences</td>
</tr>
</tbody>
</table>

A comparison of the characteristics of these five selected tools is provided in Table 2.

AliBaba acts as an interactive tool that graphically summarizes the associations between concepts from a rather limited sub-domain of the (bio)medical field, namely between cells, diseases, drugs, proteins, species, and tissues. AliBaba extracts these concepts and the associations between them from scientific publications that match a PubMed query.

Semantic MEDLINE provides similar functionality in a broader domain as it uses concepts in the UMLS Metathesaurus as its base. These concepts, and their relationships, are extracted, respectively identified, from the complete MEDLINE database, and, similar to AliBaba, subsequently presented as a graph. The Semantic Navigator also employs a graph-based format. In this tool, the graph is used to represent the semantic structure of the UMLS, and as such enables users to visually explore the concepts in the UMLS as well as their relationships.

Contrary to the graph-based format employed by AliBaba, Semantic MEDLINE, the Semantic Navigator, and BioMed Xplorer for visualizing (bio)medical
knowledge, EBIMed and PGviewer make use of two alternative visualization paradigms. On the one hand, EBIMed identifies relationships between a set of (bio)medical concepts extracted from publications that match a MEDLINE query, and visualizes these in a tabular format. The concepts represented in EBIMed stem from the (bio)medical subdomain consisting of proteins, Gene ontology annotations, drugs, and species. On the other hand, PGviewer employs tree visualization, with the purpose of clustering, in order to present relationships from the genotype and phenotype subdomain that are stored both in structured, such as MEDLINE, and (unstructured) textual databases, such as the OMIM.

The five tools identified from prior work, in summary, thus have two main shortcomings. On the one hand they focus on a particular subdomain of the (bio)medical field (AliBaba, EBIMed, PGviewer) and, as such, inhibit the exploration of the body of (bio)medical knowledge. On the other hand they employ an alternative visualization paradigm (EBIMed and PGviewer) that is less focused on the visual representation of knowledge. The BioMed Xplorer UI overcomes these shortcomings, as such improving over most tools developed in prior work, through its broad scope, aiming to cover the complete (bio)medical domain, and its graph-based visualization. More specifically, BioMed Xplorer can be considered to be on par with Semantic MEDLINE and the Semantic Navigator as these two tools both focus on the entire (bio)medical field as well as employ a graph-based paradigm for visualizing (bio)medical knowledge. The three aforementioned tools are furthermore available on the Web, whereas AliBaba, EBIMed, and PGviewer are no longer available. The position of BioMed Xplorer is further reinforced by the fact that it is the only tool that is based on RDF, which improves its ability to handle large amounts of heterogeneous data from disparate sources. Other tools, on the other hand, are based on traditional relational databases, such as such inhibiting their ability to incorporate data from additional external sources into these tools.

In addition to representing (bio)medical knowledge through statements that relate two (bio)medical concepts to each other, the presentation of concepts and statements related information is also of great importance, as it provides background knowledge about the concepts involved in the statements or about the statements themselves. To this end, BioMed Xplorer is on par with all of the other tools considering the presentation of statement related information. This information typically includes the complete statement itself, including its source and target concepts, the type of the statement, as well as the provenance data associated with the statements in terms of the abstract or sentences, and publications from which the statements were derived. Such provenance data is provided by all the tools included in the comparison, ex-

<table>
<thead>
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<th>Statement</th>
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<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
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<tbody>
<tr>
<td>The ontology is capable of representing (statements of) biomedical knowledge</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>The ontology models (statements of) biomedical knowledge appropriately</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>The ontology is capable of representing the provenance data associated with (statements of) biomedical knowledge</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>The ontology models provenance data associated with (statements of) biomedical knowledge appropriately</td>
<td>0</td>
<td>1</td>
<td>1</td>
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<tr>
<td>The ontology globally fits its purpose</td>
<td>0</td>
<td>0</td>
<td>2</td>
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</tr>
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</table>

Table 4: Frequency distribution of the five point Likert-scale scores for evaluating the BioMed Xplorer UI. A score of 1 indicates disagreement and 5 indicates agreement.
cept for the Semantic Navigator as this tool solely represents the relationships stored in the UMLS. Occasionally, the statement related information might also include aggregates of the provenance data, such as the number of sentences and publications from which a particular statement is derived, as is the case for Semantic MEDLINE and BioMed Xplorer. Whereas the BioMed Xplorer is on par with the other tools in relation to the presentation of statement related information and the incorporation of provenance data, it in fact improves over these tools on the presentation of concepts related information. The concepts related information presented in BioMed Xplorer UI extends well beyond the conventional information that is incorporated. While, tools such as EBIMed and the Semantic Navigator do not present any of such concepts related information at all, data items such as (semantic) types, synonyms, and parts of publications that mention the particular concept are presented by AliBaba, PGviewer, and Semantic MEDLINE. BioMed Xplorer extends this further through the incorporation of a wide range of cross-identifiers of concepts, a definition, and a range of data items pertaining to the clinical features, diagnosis, inheritance, pathogenesis, and genetics of a disease from OMIM, if available. The presentation of this wide range of concept related information in BioMed Xplorer is partially facilitated through the incorporation of data from external (structured) data sources, including Linked Life Data and Bio2RDF, which demonstrates its superiority compared to the other tools developed in prior work. Among these other tools, the incorporation of information from such external sources is either largely absent (such as in Semantic MEDLINE and Semantic Navigator), or limited to the inclusion of information from PubMed (such as in AliBaba, EBIMed, and PGviewer). Links to external data sources, usually in the form of cross-references to standardized terminologies, on the other hand, are commonly used by the tools developed in prior work, with only PGviewer and the Semantic Network lacking such cross references.

For the validation of the ontology, the ontology developed by (Tao et al., 2012) is considered as the base to which our developed ontology is compared. A comparison of the characteristics of the two ontologies is provided in Table 1. As is clear from this table, the ontology developed in this research improves the ontology developed by (Tao et al., 2012) on a number of aspects, which will be further discussed below, as such contributing to the validation of the ontology developed in this research. As was discussed in section 3.1, reification has been applied in both ontologies to allow triples to involve a particular (bio)medical statement, as a whole, into another statement, and thus enable meta-statements: statements about statements. This can be achieved by treating a statement, relating two (bio)medical concepts to each other through a relation, as a separate entity to which the subject, the predicate, and the object of the original statement are assigned using an object property. The ontology developed by (Tao et al., 2012) performs this by making use of the Association class in combination with the has_name, has_predicate, and haso_name properties. BioMed Xplorer Ontology, on the other hand, makes use of the official RDF reification vocabulary that uses the rdf:Statement class in combination with the rdf:subject, rdf:predicate, and rdf:object properties. Additionally, the use of this official RDF reification vocabulary also contributes to the reuse of existing vocabularies, one of the key principles of Semantic Web (Shadbolt et al., 2006). To further promote this base principle of the Semantic Web, the developed ontology, in addition to the use of the RDF reification vocabulary, makes extensive use of existing classes and properties from other vocabularies. This is a considerable improvement over the ontology developed by (Tao et al., 2012), as the reuse of existing vocabularies, aside from the RDF vocabulary, is not present in their ontology.

Since the purpose of the developed model is to enable researchers to explore the body of (bio)medical knowledge as well as its (disease) related information, the amount of information captured by the ontology is of great importance. To this extent, the ontology developed by (Tao et al., 2012) can be considered as rather limited due to the fact that there is no direct evidence of the incorporation of any (disease) related information beyond the three datatype properties assigning names to concepts and relations, as well as identifiers to publications. The ontology developed in our research improves on this point by associating 17 data-type properties to the ontology classes that can be used to capture a wide range of (disease) related information. This is further facilitated by the incorporation of RDF links to the equivalent resources in external data sources, including Linked Life Data and Bio2RDF, which contain a wealth of (disease) related information. No such links are incorporated in the ontology developed by (Tao et al., 2012).

Finally, the developed ontology extends the ontology developed by (Tao et al., 2012) by incorporating the sentences from which the represented statements are derived, in addition to those publications from which these sentences are a part, as a component of the provenance data that is associated to the statements. The incorporation of these sentences provides additional value to the disease related informa-
tion model as it enables the presentation of the direct source of a particular statement, as opposed to the presentation of solely the publication from which a statement is derived.

7 CONCLUSION

Two common shortcomings among (bio)medical knowledge discovery representation and visualization tools are the scarcity of the information that is represented, usually coming from a single source, as well as the lack of intuitiveness. To address this gap, this research aimed at developing a dynamic model representing (bio)medical knowledge, available from disparate sources across the Web, as a network of inter-related (bio)medical concepts, while incorporating Semantic Web technologies to deal with large amounts of dynamic and heterogeneous information.

To achieve this goal, a five phase research approach has been followed, consisting of: 1) State of the Art Assessment, 2) Data Source Characterization and Selection, 3) Data Preprocessing and Ontology Design, 4) Data Interlinking and Fusion with external sources, and 5) Model Visualization. Completion of these phases resulted in the development of the BioMed Xplorer Ontology, providing a foundation of the knowledge base, and the BioMed Xplorer UI, acting as a visualization of the knowledge base.

Future work will focus on implementing key indicators, representing the importance of instances, to more efficiently regulate which concepts and statements are presented to the user. To this end, indicators such as the degree of concepts, or number of sentences or publications from which a statement is derived, might be used. A second point of future work will focus on extending BioMed Xplorer’s functionality with extensive filtering options, as such enabling the user to view important, or less important, concepts and statements based on key indicators.

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REFERENCES


