

Towards a Query Rewriting Algorithm over Proteomics XML Resources

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Abstract

Querying and sharing Web proteomics data is not an easy task. Given that, several data sources can be used to answer the same sub-goals in the Global query, it is obvious that we can have many candidates' rewritings. The user-query is formulated using Concepts and Properties related to Proteomics research (Domain Ontology). Semantic mappings describe the contents of underlying sources. In this paper, we propose a characterization of query rewriting problem using semantic mappings as an associated hypergraph. Hence, the generation of candidates' rewritings can be formulated as the discovery of minimal Transversals of an hypergraph. We exploit and adapt algorithms available in Hypergraph Theory to find all candidates rewritings from a query answering problem. Then, in future work, some relevant criteria could help to determine optimal and qualitative rewritings, according to user needs, and sources performances.

Keywords: Proteomics, Ontology, XML, Trees, Semantic Web, Query Rewriting, minimal Transversals.

1. INTRODUCTION

The rapid progress of biotechnologies and the multiple genome project [7], [5] about organisms and diverse species have generated an increasing amount of proteomic data stored in many sources [17] available and publicly accessible on the Web. They contain data about metabolic pathways, protein 3D structures, DNA Sequences, organisms, diseases, and so on. Many biological questions require that data from several data sources are queried, searched, and integrated. The first step in this process of Data Integration is the Query rewriting which consists of reformulating a global query in several specific local queries. The query rewriting problem has recently received significant attention because of its relevance to a wide variety of data

management problems [14]: query optimization, maintenance of physical data independence, data integration, and data warehouse design.

Semantic mappings can be used to adapt a global query expressed in terms of a Domain Ontology in terms of specific local sources. In fact, as several sources could provide expected resources, more than one could be relevant to rewrite the global query.

The motivation of this work is to investigate how to generate candidate rewritings when user-query is posed over XML Sources, and mappings are expressed in LAV Approach. To achieve this goal, we characterize the query rewriting problem as minimal Transversals Discovery from an associated Hypergraph. From this set of generated rewritings, we would compute an optimal and best quality rewriting based on some defined and relevant criteria. We illustrate an intuitive execution of the rewriting algorithm proposed, using a scenario of Proteomics data sources.

The following two XML sources contain data that are semantically similar, but are described with autonomous and heterogeneous schemas. They both represent the same proteomics data, but not identically. They give an idea of differences in terms of terminologies, structures and contents.

```
<PROTEIN_SET>
<PROTEIN>
<ACCESSION>P26954</ACCESSION>
<ENTRY_NAME>IL3B_MOUSE</ENTRY_NAME>
<PROTEIN_NAME>Interleukin-3 receptor
class II beta chain [Precursor]
</PROTEIN_NAME>
<GENE_NAME>CSF2RB2</GENE_NAME>
<ORGANISM taxonomy_id="10090">Mus
musculus</ORGANISM>
</PROTEIN>
...
</PROTEIN_SET>
```

```
<PROTEIN_BASE>
<PROTEIN ACC_NUMBER="P26954 ">
<ENTRY>IL3B_MOUSE</ENTRY>
<PROTEIN_NAME>Interleukin-3 receptor
class II beta chain [Precursor]
</PROTEIN_NAME>
<GENE>CSF2RB2</GENE>
<ORGANISM >
<TAX_ID>10090</TAX_ID>
<NAME> Mus musculus</NAME>
</ORGANISM>
</PROTEIN>
...
</PROTEIN_BASE>
```

FIGURE 1: XML Data Source 1

FIGURE 2: XML Data Source 2

We can remark also some semantics heterogeneities, like ACC_NUMBER attribute in Source 2 which is equivalent with ACCESSION Element in Source 1.

Although both data sources contain semantically similar proteomic data, the simple user query "Which are proteins that are encoded by the gene named by CSF2RB2 in Mus Musculus Organism? " need to be formulated quite differently with existing XML query languages, like Xquery or XPath for both sources.

So, flexible, user-centric and semantic strategies of discovering relevant sources are needed to compute optimal and best quality rewriting, according to suitable criteria.

The rest of the paper is structured as follows. Section 2 gives a brief survey on our related work regarding existing query rewriting approach in Mediators. Section 3 discusses the basic concept of Knowledge representation. The hypergraph-based semantic query rewriting algorithm is presented in Section 4. The last section 5 draws the conclusions and future work.

2. RELATED WORK

One of the first LAV systems that allow the integration of XML is AGORA [21]. But AGORA still makes an extended use of the relational model: Although it offers an XML view for relational and XML data, this view is translated into a generic relational schema, XML resources are described as relational views over this schema and XQuery expressions are translated to standard SQL queries, which are then decomposed and evaluated.

Information Manifold [19] also follows a local-as views approach. In this system, the global schema is a flat relational schema, and Description Logics are used to represent hierarchies of classes. The sources are expressed as relational views over this schema. Query rewriting is done

by the Bucket algorithm which rewrites a conjunctive query expressed of the global schema using the source views. It examines independently each of the query sub-goals and tries to find rewritings but loses some by considering the sub-goals in isolation.

STyX system [12] already uses a domain Ontology as global schema language and translates an OQL-like global query language to XQuery expressions on the heterogeneous XML sources. STyX maps XPath expressions to ontology concepts. In [1], a data integration system whereby XML sources are mapped into a simple ontology (supporting inheritance and roles), is discussed. In [18], authors have also used to integrate XML heterogeneous data sources. Their work consists to map XML schema constructs to concepts. The main difference to STyX is the approach to semantic mapping. Although Lehti's approach is not as flexible and powerful as using XPath mappings, it is in principle able to detect inconsistencies in the mapping with the help of a description logic reasoner [2].

Other data integration approaches that use an Ontology as Global Schema are either based on an extended data warehouse. Semantic mediation in C-Web [2] is based on thesauri. In Xyleme mediator [8], the global schema is a set of abstracted DTDs which are terms Trees according to domain vocabularies such as Culture or Tourism. Both follow GLAV (GAV and LAV together) because correspondences between Mediator vocabulary and Sources vocabularies are expressed by simpler path mappings.

Recently, with the development of Semantic Web, mediation systems have been developed. Project Piazza [15] proposes an infrastructure based on Peer-to-Peer (like a decentralized mediator) for RDF and OWL data integration.

According to the query rewriting algorithm, you can refer to [14] for a large survey on the Query rewriting problem. Our approach is inspired by WS-CatalogNet's semantic-driven Algorithm. In this work, [3] have developed a novel and more advanced query rewriting techniques for flexible and effective E-Catalogs selection.

3. REPRESENTING KNOWLEDGE

In order to rewrite semantically a global query, it is essential to make a choice of an adequate abstraction model for local sources and to express in a common formalization language all available knowledge. This last case concerns the domain ontology, the semantic mappings, and the user query.

The proteomics sources which we are working with are stored and available as XML Documents according to their XML Schemas. XML [6] is presently becoming the standard for the exchange of biological data sources. So, the reason for the use of XML Sources for the data Integration is obvious. XML Schemas [24] are more suitable than DTDs for expressing the syntax, structural, cardinality and typing constraints required by proteomics data. We propose to abstract sources XML Schemas as unordered Trees, and we try to propose a specification language based on description logic (DL) [2] formalization and reasoning (Trees Logics).

3.1 Trees abstract model

We know that XML Schemas are special XML Documents. Various models have been proposed to represent XML Documents. The W3C proposed a generic model named Document Object Model (DOM). In this model, presented in the current section, XML Documents can be abstracted as Trees. Our motivation using this way of abstraction is to further exploit some achieved and well known results on Trees Embedding Problem [24] as knowledge semantic retrieval in an integration framework.

3.2 Logic-based Trees descriptions

To provide a semantic formalization, necessary for rigorous characterization of proteomic queries and knowledges, we propose to use a description language of hierachic structures such as Trees, based on Logics and called *Trees-Logics*.

Many researchers have addressed the question of using logics over Trees. In [9] and [10] authors have translated XQuery global queries into local conjunctive queries over Trees in a Data integration processes. In [24] a language called ApproXQL, which exploits among others logical

operators, to formalize more richer and expressive requests, has been developed. These requests expressed as conjunctive queries could be illustrated and interpreted as Trees. Then, in the paper [13] authors have studied the complexity and the expressive power of conjunctive queries over Trees.

So, we believe strongly that it is possible to describe data Trees with a suitable subset of logical formalisms [2]. We want to exploit all these works in order to provide a logical description of hierarchical structures, such as Trees and consequently Paths in particular. In our final integration framework, both the phases of Trees generation and their specification in *Trees-Logics* are totally transparent for the user. We precise that Description Logics [2] are a family of logics which were developed for modeling complex hierarchical structures and to provide a specialized reasoning engine to do inferences on these structures.

Due to the space limitation, we could not give more details on *Trees-Logics* and so this paper will focus only on the query rewriting problem.

4. QUERY REWRITING

In this section, we begin by presenting an abstraction of our approach for query rewriting. Then, we show that using some hypergraph Theory results can help generate candidate rewritings. Therefore we present the Classical algorithm to compute minimal Transversals of a hypergraph. Finally, we illustrate an execution of this algorithm to find candidate rewritings, given concrete case of bio-query reformulation.

We recall that the main goal of query rewriting phase is to reformulate a Global query Q expressed as *Trees-Logics* over Domain ontology, into Local queries Q_j that are expressed in terms of Local schemas. This operation is realized using semantic mappings pre-calculated and stored on the mediator. Semi-automatic detection of semantic mappings has no impact on the processing time of the user query.

The concrete algorithm showing how these semantic mappings are calculated is out of scope of this paper.

The domain Ontology is abstracted as a Tree and expressed using the defined language, *Trees-Logics*. The knowledge domain concerns proteomics research including concepts such Protein Family, 3D Structures, Coding Genes, Motifs, Domains, amino-acids sequences, Active Sites, Binding Sites, Enzymes, Chains, Chemical Bonds, ... and their relative properties, so we call the ontology by *O'proteomics*. Due to space constraints, we could not give more details on *O'proteomics*.

The Ontology constitutes a support for user query formulation and gives an idea of which concepts, it is possible (but not obliged) to find or retrieve in the underlying Proteomics sources. Therefore, the first initiative consists of determining the part of the query that cannot be answered by available proteomics sources.

4.1 Logic-based Trees descriptions

We represent by the following couple $Sch' O = (O' \text{ proteomics}, M \text{ mappings})$, the set of semantic knowledges about our domain of interests, which is proteomics.

The concepts annotations, defined in *O'proteomics* will serve to enrich Global query before rewriting process. The semantic mappings will show query answering capabilities of the underlying sources.

Given a Global query Q and the knowledges couple $Sch' O$, our rewriting approach consist to determine two sub-queries Q_{valide} and $Q_{invalide}$. Explicitly, we shall calculate:

- $Q'' = Q_{invalide}$ having a size as minimal as possible. The Size of a query is the number of atomic goals that it contains. Sub-Query Q'' cannot be answered by underlying sources, at the moment of the sending of the Global query Q . This initial operation has the role of cleaning up Q of domain concepts/properties which are

not yet available, as Web proteomics registered resources. So, no processing will be realized on Q'' , in the future.

- $Q' = Q_{valide}$ is the part of Q that will be rewrite using semantic mappings M of $Sch' O$. Sub-query Q' can be answered by registered sources. Our final goal is to propose an intelligent subdivision of $Q' = Q_{valide}$ into sub-queries Q'_1, Q'_2, \dots, Q'_m with $1 \leq m \leq n$, n is the number of sources available in the integration while m denotes the number sources which are necessary to provide an answer to the query Q . So, we might find the set $Q' = \{(Q'_j, m_j)\}$ of couples (Q'_j, m_j) such as Q'_j be an atomic subdivision of Q' that will be answered by mapping m_j .

We can easily remark that several rewritings can be proposed, we will call them Candidates rewritings. In fact, more than one source, and so mapping could provide the same resources researched.

The algorithm receives as input a global query Q , a schema $Sch' O$ and generate as output all candidates' rewritings $r_i(Q)$.

4.2 Hypergraph-based Algorithm

In practice, Global queries are expressed like conjunctive queries using *Trees-Logics*. So, a rewriting Q' is a suitable conjunction of constraints. These constraints might be checked by all final answers of the global query Q , because they constitute an indication of resources that may be retrieved from adequate sources.

In order to provide a characterization of our query rewriting problem, we give an alternative formulation of the rewriting formalization.

Given a Global Query Q and the semantic knowledges couple $Sch' O$, query rewriting consists to compute two sub queries Q_{valide} and $Q_{invalid}$ on the basis of mappings set M , such as:

$$Q = Q_{valide} \wedge Q_{invalid} \quad (1)$$

We are searching for all candidates rewritings, formulated as the conjunction of constraints:

$$Q_{valide} = Q' = \bigwedge_{i=1}^m C_i \quad (2)$$

All constraints C_i are logical representation of the user specific needs. Finally, our motivation is to answer the fundamental question which is to find, given Q a new query called rewriting expressed by $Q_{valide} = Q' = \bigwedge_{i=1}^m C_i$ such as Q' denotes as much as possible the resources expected by query Q ?

We have said that several and alternatives rewritings are possible, due to the fact that more than one mapping could be used to reformulate an atomic goal of the Global query. From this point of view, rewriting problem which requires generating all candidates' rewritings can be characterized as a current Hypergraph Theory problem of computing all minimals Transversals of an Hypergraph. Generate a Transversal Hypergraph consists of generate all minimals Transversals.

4.2.1 Definition of Hypergraph [16].

An Hypergraph H is an ordered pair $H = (V, E)$ where $V = \{v_1, v_2, \dots, v_n\}$ is a finite set of elements and $E = \{E_1, E_2, \dots, E_m\}$ is a family of subsets of V such that:

- $E_i \neq \emptyset$, ($i = 1, \dots, m$)

$$\text{- } \bigcup_{i=1}^m E_i = V$$

The elements of V are called nodes while the elements of E are called hyperedges of the hypergraph H . A hypergraph can be seen as a generalization of a graph where the restriction of an edge having only two nodes does not hold.

4.2.2 Definition of Transversals [1].

Let $H = (V, E)$ be an hypergraph. A set $T \subseteq V$ is called a Transversal of H if it intersects all its hyperedges, i.e., $T \cap E_i \neq \emptyset, \forall E_i \in E$. A **Transversal** T is called minimal if no proper subset T' of T is a transversal of H . The **Transversal Hypergraph** $Tr(H)$ of an hypergraph H is the family of all minimal transversals of H .

From a rewriting query problem, we need to give a mathematical characterization, by defining an associated Hypergraph $H_{Q,M}(V, E)$, built as follows:

- For every mapping m_i , describing a local concept from M , as a logical function of O'proteomics global concepts, we associate a vertice V_{m_i} in the hypergraph $H_{Q,M}(V, E)$ and $V = \{V_{m_i}, i \in [1, n]\}$.
- For every constraint C_i of the Global query Q , we associate an hyperedge E_{C_i} in the hypergraph $H_{Q,M}(V, E)$. To simplify, we suppose that all these constraints are describing atomics goals. So, each hyperedge E_{C_i} is a set of mappings, calculated by considering those mappings which are relevant to answer these goals.

A classical algorithm to compute minimal transversals of an hypergraph is proposed and available in [20]. Many papers [11] have discussed about algorithm of generation of Hypergraph Transversal, which is a set of minimal transversals. One of the first results remains Berge's Algorithm [4], but several variants have been proposed in order to deal with the algorithm complexity [22].

Now, we present our Query rewriting algorithm called *Q-Candidates'Finder*, which integrates the better and efficient complexity of the classical Algorithm:

```

Input: A Query  $Q$  and Sch'O = (O'proteomics, M mappings)
Output: The set of candidates rewriting such as  $Q_{candidates} = \{(Q_{valide}, Q_{invalid})\}$ 
1: Build the associated Hypergraph  $H_{Q,M}(V, E)$ 
2: Compute  $Q_{invalid} = \bigwedge_{i=1}^k C_i$  such as  $C_i$  is not provided by any mappings in  $M$ .
3: Build the associated Hypergraph  $H^*_{Q,M}(V, E^*)$ 
4:  $Q_{candidates} = \emptyset$ 
5: Generate the Hypergraph Transversal of  $H^*_{Q,M}(V, E^*)$ 
   - Let be HypTransv - Using the Classical Algorithm [Mannila, 1994]
6: For all edge  $X = \{V_{m_1}, V_{m_2}, \dots, V_{m_p}\} \in HypTransv$  do
7:    $Q_{valide} = r(Q) = Q' = \{(Q'_j, m_j), j \in [1, p]\}$ 
   where  $Q'_j$  is a subdivision of  $Q_{valide}$  that will be answered by the mapping  $m_j$ .
8:    $Q_{candidates} = Q_{candidates} \cup Q_{valide}$ 
9: End For
10: Return  $Q_{candidates}$ 

```

FIGURE 3: Q-Candidates' Finder Algorithm

4.3 Q-Candidates' Finder Illustration

To illustrate the proposed rewriting approach, let us consider the following mappings (L.A.V approach). As the domain Ontology is characterized as a Tree, semantic mappings might express subsumption (**Sub**) and equivalence (**Eq**) relations that exist between Local XML Schemas, also abstracted as Trees, and the Ontology's Tree model. We suppose in order to simplify this illustration that we have only simple paths mappings (and so, no sub-trees) between Concepts, according to 1:1 cardinality. For every registered proteomics' sources, we provide the **LHS** (Left Hand Side expressing Local Concepts/Properties), the **TYPE** (the type of mappings), and the **RHS** (Right Hand Side, expressing Ontology Concepts/Properties) of the current mapping.

	LHS	TYPE	RHS
<i>O'proteomics</i> Ontology	Gene (Genes, Proteins, Species, Organisms) Proteine (IdProteins, Peptides, DevStadium)		
Mapping <i>m1</i> : Description Of Source I	S1_Gene (S1_GeneName, S1_ProteinName, S1_species, S1_organisms)	Eq	Gene (Genes, Proteins, Species, Organism)
Mapping <i>m2</i> : Description Of Source II	S2_Gene (S2_NomGenes, S2_NomProteine, S2_Especies, S2_Organismes)	Eq	Gene (Genes, Proteins, Species, Organisms)
Mapping <i>m3</i> : Description Of Source III	S3_TreeLife (S3_Species, S3_Genus...)	Sub	Gene (Species, Organisms)

TABLE 1: Mapping Table

We have just shown mappings which are relevant for the Query we shall process. Note once again, that we are considering corresponding paths of these Concepts/Properties in their abstract trees.

According to this mapping table, we can say that the ontology includes Concept Gene and Proteine with their relative Properties such as Genes, Proteins, and Species ... Mappings *m1* and *m2* show that Source I and Source II provide the properties such as Gene, Proteins, Species, and Organisms, while the mapping *m3* illustrate that Source III only provides properties such as Species and Organisms.

Let us consider now the following query which is expressed over the domain ontology, *O'proteomics*:

What are the genes which proteins could have a peptide Signal and for which, it is assumed that they are expressed at Tardive Shizont stadium for the Plasmodium falciparum?

4.3.1 Hypergraph Construction

Intuitively, *Q* can be expressed like a conjunction of the following constraints:

$$\begin{aligned} Q = & C_{\text{Genes}} \wedge C_{\text{Proteins}} \wedge C_{\text{Peptides}} \wedge C_{\text{DevStadiums}} \\ & \wedge C_{\text{Organisms}} \wedge C_{\text{Species}} \end{aligned} \quad (3)$$

In practice, the user will formulate this request by using an user-friendly graphic interface, and the generation of its Trees-Logics version is done automatically. He will choose the Concept Gene and indicate for each property Genes, Proteins, Species, Organisms, Peptides, DevStadiums the expected values.

The associated hypergraph $H_{Q,M}(V,E)$ consists of the following sets of vertices and edges:

$$V = \{V_{S2_Gene}, V_{S1_Gene}, V_{S3_TreeLife}\} \quad (4)$$

and

$$E = \left\{ \begin{array}{l} E_{C_Genes}, E_{C_Proteins}, E_{C_Peptides}, \\ E_{C_DevStadiums}, E_{C_Organisms}, E_{C_Species} \end{array} \right\} \quad (5)$$

We could see this illustration using a Sets' Theory point of view. We materialize all query constraints as Sets that contain the providers' mappings.

We show graphically these sets of mappings:

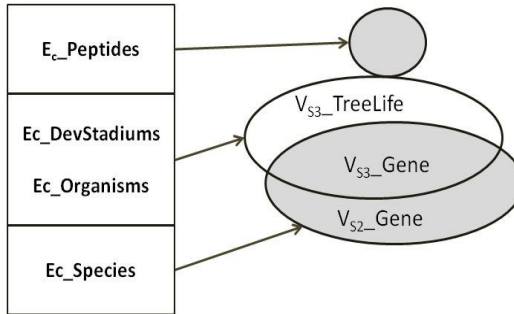


FIGURE 4: Associated Hypergraph of the Illustration

4.3.2 Determination of $Q_{invalid}$

$Q_{invalid}$ is the part of Q that cannot be answered by available sources. That means, we might constitute $Q_{invalid}$ with all Q 's constraints, which are characterized by empty hyperedges of hypergraph $H_{Q,M}(V,E)$.

We can easily see that no mapping provides the hyperedges DevStadiums and Peptides. These hyperedges are empty of associated mappings (Q-Candidates'Finder: Line 2). Hence:

$$Q_{invalid} = C_{Peptides} \wedge C_{DevStadiums} \quad (6)$$

4.3.3 Determination of Q_{valid}

Q_{valid} is the part of Q that can be answered by available sources. That means, we might constitute Q_{valid} with all Q 's constraints, which are characterized by non-empty hyperedges of hypergraph $H_{Q,M}(V,E)$.

So, we have:

$$Q_{valide} = C_{Genes} \wedge C_{Proteins} \wedge C_{Organisms} \wedge C_{Species} \quad (7)$$

In fact, Q_{valide} means that we could only answer the following request, given semantic mappings:
Give the genes which code all proteins, for the Plasmodium falciparum?

4.3.4 Determination of $Q_{candidates}$

From calculated above, we generate associated hypergraph $H^*_{Q,M}(V, E^*)$, (see Line 3).

Intuitively, according to Sets' Theory vision, finding all candidates rewriting suppose firstly to construct the Cartesian product of all sets of mappings. It is obvious that Elements of the Cartesian product are 4-uplets in our illustrative example. So, we must generate for each 4-uplet, an associated Set (These sets correspond to Transversals). In fact, it will be useful to use a minimal number of Sources that would be requested. This condition is guaranteed if we consider only associated sets that not contain another associated Set (These sets are minimals Transversals). That is why we call the Classical Algorithm (see Line 5). The maximal cardinality of our example Transversals is 4, it is the size of Q_{valide} .

From Sets' Theory point of view we can say that any set that contain V_{S1_Gene} and V_{S2_Gene} is a Transversal and constitutes possible rewritings of Q . The minimal Transversal $\{ V_{S1_Gene} \}$ and $\{ V_{S2_Gene} \}$ constitute two candidates rewritings. In our case, we could find 36 quadruplets, associated with 6 Transversals but only 2 are minimal Transversals.

5. CONCLUSION & FUTURE WORK

This paper deals with Global query rewriting, which consists in data integration context, to rewrite the global query expressed in terms of concepts and their properties defined in global schema domain ontology) into suitable terms of local data sources.

The Query rewriting process is based on semantic mappings. Our knowledge domain concern Proteomics research, and so we have proposed ontology according to interviews and talks with biologists and bio-informaticians, called O'proteomics. We provide a characterization of the Query rewriting problem based on Hypergraph Theory. We have presented the classical algorithm that computes all minimals Transversals, given an Hypergraph. We observe that those minimals Transversals correspond to Candidates rewritings of the Global Query.

Therefore, we need to better defined a logical formalism or language to specify the syntax and the semantics of data trees. It could be seen as a subset of Description Logics or based on Psi-terms formalism. After this essential choice, we will try to provide a prototype.

This paper shows briefly our current research that aims to provide a semantic framework to realize a Data Integration over XML bio-informatics sources on the Web. We will define some relevant criteria to rank candidates rewritings, necessary to select an optimal and qualitative rewriting $Q_{optimal}$. An efficient way for selecting best rewritings, iteration by iteration, will permit us to investigate the properties and the optimization of our algorithm. These relevant criteria could concern user preferences, quality of underlying sources, cost communication, etc...

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