ABSTRACT

R2RML is used to specify transformations of data available in relational databases into materialised or virtual RDF datasets. SPARQL queries evaluated against virtual datasets are translated into SQL queries according to the R2RML mappings, so that they can be evaluated over the underlying relational database engines. In this paper we describe an extension of a well-known algorithm for SPARQL to SQL translation, originally formalised for RDBMS-backed triple stores, that takes into account R2RML mappings. We present the result of our implementation using queries from a synthetic benchmark and from three real use cases, and show that SPARQL queries can be evaluated as fast as the SQL queries that would have been generated by SQL experts if no R2RML mappings had been used.

1. INTRODUCTION

Making relational database content available as RDF has played a fundamental role in the emergence of the Web of Data. Several approaches have been proposed in this direction since the early 2000s, focusing on the creation of mapping languages, models, and supporting technology to enable such transformations (e.g., R2O [2], D2R [3], Triplify [1]). In September 2012, the W3C RDB2RDF Working Group released the R2RML W3C Recommendation [9], a language to specify transformations of data stored in relational databases into materialised or virtual RDF datasets, so that relational databases can be queried using SPARQL, and several R2RML-aware implementations have been reported [19].

The R2RML specification does not provide any formalisation of the SPARQL to SQL query translation process that needs to be followed by R2RML-aware query translators, and the aforementioned implementations do not describe the formalisations of their query translation algorithms. In any case, some attempts have been done in the past to provide such a formalisation. For instance, Garrote and colleagues [11] provide a draft of the transformation of SPARQL SELECT queries to SQL based on the query translation algorithm defined in [7]. As their focus is to provide an R2RML-based RESTful writable API, the query translation algorithm that they describe is limited. For example, projections and conditions for those queries are built only for variable components of the triple pattern. However, a triple pattern may involve URIs and constants in their components, and these have to be taken into account when translating SPARQL queries. Rodríguez-Muro and colleagues have recently presented Quest/Ontop [15], which translates R2RML mappings and SPARQL queries into a set of Datalog rules, where optimizations based on query containment and Semantic Query Optimisation are applied, before transforming them into SQL queries. However, the process of how R2RML mappings are translated into Datalog rules has not been described at the moment of writing. Unbehauen and colleagues [18] define the process of binding triple patterns to the mappings and the process of generating column groups (a set of columns) for every RDF term involved in the graph pattern. Using the calculated bindings and column groups, they define how to translate each of the SPARQL operators (AND, OPTIONAL, FILTER, and UNION) into SQL queries. However, the function joinCond(s1,s2), in which two patterns are joined, is not clearly defined. In fact, one fundamental aspect that distinguishes SPARQL queries from SQL queries is the semantics of the joins, as discussed in [8, 7], which corresponds to the treatment of NULL values in the join conditions. None of the aforementioned systems explain how to deal with R2RObjectMap mappings, where a triples map is joined with another triples map (parent triples map), and consequently the generated SQL query has to take into account the logical source of the parent triples map and join condition specified in the mapping.

Furthermore, the performance of virtual RDF datasets based on RDB2RDF mapping languages has not always been satisfactory, as reported by Gray et. al. [12]. This experience has also been confirmed empirically by us in projects like Répener [17], BizkaiSense1, or Integrate2, for which we have had access to the data sources and mappings in languages like D2R. In all cases, some of the SQL queries produced by the translation algorithm could not be evaluated (e.g., too many joins are involved) or their evaluation takes too much time to complete, what makes their use in a virtual RDF dataset context unfeasible. The main reason for this is that the resulting queries are not sufficiently optimised to be efficiently evaluated over the underlying database engines.

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1 http://www.tecnologico.deusto.es/projects/bizkaisense/
2 http://www.fp7-integrate.eu
In the context of RDF data management, some works [7, 10] have focused on using relational databases as the backend for triple stores, with the idea of exploiting the performance provided by relational database systems. They normally work with schema-oblivious layouts (using a triple table layout containing columns corresponding to the triple elements) or vertical partitioned layouts (using different tables to store triples corresponding to the same predicate). They provide translation algorithms from SPARQL to SQL using the aforementioned layouts specifically designed to store triple instances. This is out of the scope of our work.

While R2RML specifies custom mappings defined by users, it comes with a companion standard called Direct Mapping [14], which defines the RDF representation of data in a relational database. The generated RDF instances will have their classes and properties reflecting the structure and contents of the relational database. Using Direct Mappings, Sequeda and colleagues [16] define the process of translating SPARQL into SQL efficiently, by removing self-joins and unnecessary conditions.

The contribution of our paper is threefold: first, we extend an existing SPARQL to SQL query rewriting algorithm [7], originally proposed for RDBMS-backed triple stores, by considering R2RML mappings. This implies allowing the algorithm to work with arbitrary layouts, such as the ones normally used in legacy systems. This work is complementary to [11] and provides an alternative to the one presented in [18] and [15]. Second, we implement and evaluate several versions of our algorithm (with different types of optimisations), using queries from the BSBM benchmark [4], a widely used synthetic benchmark for RDB2RDF systems, and show that the evaluation of such rewritten SPARQL queries is generally as fast as the corresponding native SQL queries specified in the benchmark. Third, we report our experience with queries from real projects, which show performance issues when using state-of-the-art RDB2RDF systems, and use our implementation to produce a better translation that allows them to be evaluated in an appropriate time.

The paper is structured as follows. In the following section, we review the R2RML language and Chebotko’s approach to SPARQL-to-SQL query rewriting and data translation. In Section 3 we formalise our extension to consider R2RML mappings. In Section 4 we describe our evaluation using the BSBM synthetic benchmark, and three positive experiences of applying our approach in real case projects. Finally, we present our conclusions and future work in Section 5.

2. BACKGROUND: R2RML AND CHEBOTKO’S QUERY TRANSLATION

2.1 R2RML

As discussed in the introduction, R2RML [9] is a W3C Recommendation that allows expressing customized mappings from relational databases to RDF. An R2RML mapping document \( M \), as shown in Figure 1, consists of a set of \( \text{rr}:\text{TriplesMap} \) classes. The \( \text{rr}:\text{TriplesMap} \) class consists of three properties; \( \text{rr}:\text{logicalTable} \), \( \text{rr}:\text{subjectMap} \), and \( \text{rr}:\text{predicateObjectMap} \).

\[ \text{triples} = \{ \text{subject}, \text{predicate}, \text{object} \} \]

- \( \text{rr}:\text{logicalTable} \) specifies the logical table to be mapped, whether it is a SQL table, a SQL view, or an R2RML view.
- \( \text{rr}:\text{subjectMap} \) specifies the target class and the URI generation form.
- \( \text{rr}:\text{predicateObjectMap} \) specifies the target property and the generation of the object via \( \text{rr}:\text{objectMap} \), whose value can be obtained by constant \( \text{rr}:\text{constant} \), column \( \text{rr}:\text{column} \) or template \( \text{rr}:\text{template} \). Linking with another table is accommodated by the use of \( \text{rr}:\text{refObjectMap} \), which specifies the parent triples map and the join conditions.

![Figure 1: An overview of R2RML, based on [9]](http://www.w3.org/ns/r2rml)

Appendix 6 gives an example of a mapping document that maps Product and Offer tables to \( \text{ex:Product} \) and \( \text{ex:Offer} \) respectively, which we will use throughout this paper to illustrate our definitions and algorithms.

2.2 Chebotko and colleagues’ approach

Our approach is based on the formalisation and algorithm from Chebotko and colleagues [7]. Other RDB2RDF query translation approaches lack a clear formalisation of the approach (e.g., [2], [3]) or they are not focused on RDF generation and/or SPARQL querying (e.g., [6]).

Chebotko’s approach proposes the use of a set of mappings \( (\alpha, \beta) \) and functions \( \text{genCondSQL}, \text{genPRSQL}, \text{genname} \) to translate SPARQL into SQL. Its formal definition is available at [7]. We now give an example of how these mappings and functions work in the absence of R2RML mappings. In Section 3 we provide the formal definition of these mappings and functions with R2RML mappings.

Example 1. Without R2RML Mappings. Consider a triple table \( \text{Triples}(s, p, o) \) that uses its columns to store the subject, predicate, and object of RDF triples. A user poses a SPARQL query with the triple pattern \( \text{tp1} = :\text{Product1 rdfs:label} \text{?label} \).

- Mapping \( \alpha \) returns the table that holds the triples that correspond to this triple pattern \( \text{tp1} \). That is, \( \alpha(\text{tp1}) = \text{Triples} \).
- Mapping \( \beta \) returns the column that corresponds to each triple pattern position \( (\text{sub}, \text{pre} \text{ or obj}) \). That is, \( \beta(\text{tp1, sub}) = s, \beta(\text{tp1, pre}) = p, \text{and} \beta(\text{tp1, obj}) = o. \)
3. AN R2RML-BASED EXTENSION OF CHEBOTKO'S APPROACH

We have seen from the previous example how the algorithm defined in [7] is used to translate SPARQL queries into SQL queries for RDBMS-backed triples stores. As mentioned in the Introduction section, this approach has to be extended so that all the functions and mappings used in the algorithm take into account user-defined R2RML mappings. Before we present the detail of how to extend the algorithm, we give an illustrative example that is comparable to the previous example.

3.1 Illustrative Example

Example 2. With R2RML Mappings. Consider a table Product(nr, label) that is mapped into an ontology concept bsbm:Product. The column nr is mapped as part of the URI of the bsbm:Product instances and column label is mapped to the property rdfs:label. The same triple pattern tp1 = :Product rdfs:label ?placeb in the previous example produces the following mappings/functions:

- Function genCondSQL filters the table Triples returned by α(tp1) so that only those records that match the triple pattern tp are returned. genCondSQL(tp1, β) = (s = :Product1 AND p = rdfs:label).
- Function name generates alias for each triple pattern element. That is, name(Product1) = iri_Product1, name(rdfs:label) = iri_rdfs_label, and name(label) = var_plabel.
- Function genPRSQL projects the β column as the alias name of each triple pattern element. That is, genPRSQL(tp1, β, name) = {nr AS iri_Product1, 'rdfs:label' AS iri_rdfs_label, label AS var_plabel}.
- Function trans(tp1) finally returns the SQL query using the values returned by the previous mappings and functions. That is, trans(tp1, α, β) = SELECT s AS iri_Product1, 'rdfs:label' AS iri_rdfs_label, label AS var_plabel FROM Product WHERE nr = 1 AND label IS NOT NULL.

Figure 2 illustrates our two examples on using the Chebotko’s algorithm in the context of triple stores and R2RML-based query translation.

Next, describe how we extend the translation function of a triple pattern in Chebotko’s approach taking into account R2RML mappings. We then go into more detail explaining the mappings/functions used in the translation algorithm5.

3.2 Function trans under M

Definition 1. Given the set of all possible triple patterns \( TP = (IV) \times (IV) \times (IVL) \) and \( tp \in TP \), mappings \( \alpha \) and \( \beta \), functions name, genCondSQL, and genPRSQL, \( \text{trans}(tp) \) generates a SQL query that can be executed by the underlying RDBMS to return the answer for gp.

Listing 1: \( \text{trans}(tp) \) under M

```
trans3\((\text{TP, } \alpha, \beta, \text{name, Queries}) \) ->
getTriplesMaps(\( \text{TP, M, TMList} \)).
transTMList(\( \text{TP, } \alpha, \beta, \text{name, TMList, Queries} \)).
transTMList(\( \text{TP, } \alpha, \beta, \text{name, TMListTail, Queries} \)).
createUnionQuery(QHead, QTail, Queries).
transTM(\( \text{TP, } \alpha, \beta, \text{name, TM, QHead} \)).
transTM(\( \text{TP, } \alpha, \beta, \text{name, TM, QHeadTail} \)).
createUnionQuery(QHead, QTail, Queries).
transPreURLList(\( \text{TP, } \alpha, \beta, \text{name, TM, PreURL} \)).
\text{getPreURLsMaps}(\( \text{TP, TM, PreURLList} \)).
transPreURLList(\( \text{TP, } \alpha, \beta, \text{name, TM, PreURLList, Queries} \)).
createUnionQuery(QHead, QTail, Queries).
transPreURL(\( \text{TP, } \alpha, \beta, \text{name, TM, PreURL} \)).
\text{getPreURLsMaps}(\( \text{TP, TM, PreURL} \)).
\text{buildQuery}(\text{PrePart, FromPart, WherePart}).
```

The algorithm (see Listing 16) for translating a triple pattern \( TP = (IV) \times (IV) \times (IVL) \) can be explained as follows:

- A triple pattern can be mapped to several triples maps in the mapping document, thus all results from the possible mappings will be put in a single UNION query (line 1-7).

5While the order of presenting the mappings/functions is not important, we start by explaining \( \text{trans}(tp) \), so as to facilitate understanding.

6We use Prolog-syntax to present our listings.

7For compactness, we write IRI as I, Variable as V, Literal as L, and combinations of these letters.

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For the sake of simplicity, we omit the aliases that have to be prefixed on each column name.

4
The same process occurs if the predicate part of the triple map is unbounded. Then the variable of the predicate part will be grounded according to available mappings, and results will be merged as a UNION query (line 11-20).

In the case where the triples map is defined and the predicate of the triple pattern is bounded, \( \text{trans}(tp) \) builds a SQL query using the result of auxiliary mappings \( (\alpha, \beta) \) and functions \( \text{genCondSQL}, \text{genPRSQL} \) (line 22-26). Those mappings and functions are explained in the following subsections.

The translation function \( \text{trans} \) for other patterns (AND, OPTIONAL, UNION, FILTER) follows the algorithm described in [7].

3.3 Mapping \( \alpha \) under \( M \)

Definition 2. Given the set of all possible triple patterns \( TP = (IV) \times (I) \times (IVL) \), a set of relations \( REL \), and a set of all possible mappings \( M \) defined in R2RML, a mapping \( \alpha \) is a many-to-many mapping \( \alpha : TP \times M \rightarrow REL \), where given a triple pattern \( tp \in TP \) and a mapping \( m \in M \), \( \alpha^m(tp) \) returns a set of relations that generate all the triples that match \( tp \).

Listing 2: \( \alpha \) under \( M \)

<table>
<thead>
<tr>
<th>Line</th>
<th>Expression</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>( \alpha^m(TP, TM, PreURI, [AlphaSub, AlphaPreObj]) := )</td>
</tr>
<tr>
<td>2</td>
<td>( \alpha_{\text{ub}}(TM, AlphaSub, AlphaPreObj) )</td>
</tr>
<tr>
<td>3</td>
<td>( \alpha_{\text{preobj}}(TP, TM, PreURI, AlphaPreObj) )</td>
</tr>
<tr>
<td>4</td>
<td>( \alpha_{\text{preobj}}(TM, TM.logicalTable) )</td>
</tr>
<tr>
<td>5</td>
<td>( \alpha_{\text{reobj}}(TP, TM, PreURI, AlphaPreObj) := )</td>
</tr>
<tr>
<td>6</td>
<td>( \text{getPredicateObjectMap}(TM, PreURI, POMap) )</td>
</tr>
<tr>
<td>7</td>
<td>( \text{ROMap = POMap.refObjectMap} )</td>
</tr>
<tr>
<td>8</td>
<td>( \alpha_{\text{preobj}}(TM, PreURI, ROMap, AlphaPreObj) )</td>
</tr>
<tr>
<td>9</td>
<td>( \alpha_{\text{preobj}}(TM, PreURI, ROMap, ROMap.parentTriplesMap.logicalTable) )</td>
</tr>
</tbody>
</table>

The algorithm for computing mapping \( \alpha \) under a set of R2RML mappings is provided in Listing 2. The output of this algorithm is used as the FROM part of the generated SQL query. This algorithm is divided into two main parts; calculating \( \alpha \) for the subject (\( \alpha^m_{\text{ub}}(TM) \)) (line 5), and for the predicate-object part \( \alpha_{\text{reobj}}(POMap) \) (line 13).

- The function \( \alpha^m_{\text{ub}}(TM) \) returns the logical table property \( \text{logicalTable} \) of a triples map \( TM \) (line 5).
- A logical table from a triple may be joined with another logical table through the \( \text{refObjectMap} \) property. This case is handled by an auxiliary function \( \alpha_{\text{reobj}}(POMap) \) that retrieves the \( \text{predicateObjectMap} \) property \( POMap \) of \( TM \) that corresponds to the predicate of the triple pattern \( tp \), and then returns the parent logical table of \( \text{refObjectMap} \) property \( ROMap \) (line 7-13).

The output of mapping \( \alpha \) is a set of logical tables with the result from the two auxiliary functions (line 1-3). Table 1 presents some examples of mapping \( \alpha \) results.

3.4 Mapping \( \beta \) under \( M \)

Definition 3. Given a set of all possible triple patterns \( TP = (IV) \times (I) \times (IVL) \), a set of positions in a triple pattern \( POS = \{\text{sub}, \text{pre}, \text{obj}\} \), a set of relational attributes \( \text{ATTR} \), and a set of all possible R2RML mappings \( M \), a mapping \( \beta \) is a many-to-one mapping \( \beta^m : TP \times POS \times M \rightarrow \text{ATTR} \), if
given a triple pattern \( tp \in TP \), a position \( pos \in POS \), and \( m \in M \), \( ATTR \) is a relational attribute whose value may match the triple pattern \( tp \) at position \( pos \).

The algorithm for computing the mapping \( \beta \) under \( M \) is provided in Listing 3 and some examples of \( \beta \) results can be seen on Table 1. The output of mapping \( \beta \) is used in the functions \( genPRSQL \) and \( genCondSQL \) to select the relational attribute corresponding to the triple pattern position:

- If the position \( pos \) is subject, then the auxiliary function \( \gamma_{m}^{S} \) returns the corresponding relational attributes attached to the subject map \( SM \) of \( TriplesMap \) (line 1-2, 8-9).

- If the position \( pos \) is predicate, then the function \( \gamma_{m}^{P} \) is returned as a constant. (line 3-4, 11).

- If the position \( pos \) is object, then the function \( \gamma_{m}^{O} \) (line 13-16) checks whether the argument \( POMap \) contains a Reference Object Map \( ROMap \).
  - If \( POMap \) contains \( ROMap \), the parent triple map \( ParentTriplesMap \) of \( ROMap \) is retrieved and the relational attributes corresponding to the subject map of \( ParentTriplesMap \) are returned (line 20-21).
  - Otherwise, the corresponding relational attributes of object map \( objectMap \) are returned (line 18-19).

The auxiliary function \( genCondSQL \) under \( M \) checks the term type of \( tp.subject \). If \( tp.subject \) is an IRI (line 6-11), then the identifier of this subject is obtained by a function \( inverseExpression \), a function that takes an IRI and returns only the corresponding values that can be related to database values. For example, \( inverseExpression \) will return 1 or 3847 for IRIs corresponding values that can be related to database values. Only in case that the value is used in some filter expressions (e.g., \( FILTER(\{\text{bound}(\text{?label})\}) \)) that we must remove the IS NOT NULL expression so that \( genCondSQL \) returns NULL, what will be interpreted as unbounded values.

- If \( tp.object \) is a variable, two cases are evaluated:
  - In the case that PredicateObjectMap doesn’t have any Reference Object Map, then the SQL expression stating that the \( \beta \) of PredicateObjectMap IS NOT NULL is returned (line 21-23). This is to guarantee that we do not return NULL values. Only in case that the value is used in some filter expressions (e.g., \( FILTER(\{\text{bound}(\text{?label})\}) \)) that we must remove the IS NOT NULL expression so that \( genCondSQL \) returns NULL, what will be interpreted as unbounded values.

  - If \( RefObjectMap \) of PredicateObjectMap is specified, then a SQL expression that is the \( JoinCondition \) property from \( RefObjectMap \) is returned (line 24-26).

- If \( tp.predicate \) is a literal, then a SQL equality expression of \( tp.object \) and the output of \( \beta \) for PredicateObjectMap is returned (line 28-30).

3.5 Function \( genCondSQL \) under \( M \)

Definition 4. Given a set of all possible triple patterns \( TP \), a mapping \( \beta \), and a set of all possible mappings \( M \) defined in R2RML mapping document, \( genCondSQL \) generates a SQL expression that is evaluated to true if and only if \( tp \) matches a tuple represented by relational attributes \( \beta_{m}^{S}(tp,sub), \beta_{m}^{P}(tp,pre), \) and \( \beta_{m}^{O}(tp,obj) \) where \( tp \in TP \) and \( m \in M \).

The algorithm of function \( genCondSQL \) under \( M \) is provided in Listing 4. The output of this function is used as the WHERE part of the generated SQL query. This function calls and returns the result of two auxiliary functions: \( genCondSQL_{sub}^{m} \) and \( genCondSQL_{preobj}^{m} \).
The algorithm for computing genPRSQL under \(\mathcal{M}\) is provided in Listing 5. The outputs of this function are used in the SELECT part of the generated SQL as the select items of the SQL queries together with their aliases, which are generated by function \(\text{name}\). As defined in \[7\], given a term in IVL, a function \(\text{name}\) generates a unique name such that the generated name conforms to the SQL syntax for relational attribute names. In our examples, \(\text{name}(t)\) generates term type appended by the name of the term. For example, the result of \(\text{name}(\text{?lb})\) is \(\text{var}_{\text{lb}}\) and the result of \(\text{name}(\text{hasOffer})\) is \(\text{iri}\_\text{hasOffer}\). The function genPRSQL projects all the \(\beta\) attributes and assigns them unique aliases generated from the name function (line 16, 19, and 25). Some examples of \(\text{genPRSQL()}\) results can be seen in Table 1.

### 3.6 Function \(\text{genPRSQL under } \mathcal{M}\)

**Definition 5.** Given a set of all possible triple pattern \(\mathcal{TP}\) and \(\mathcal{tp} \in \mathcal{TP}\), a mapping \(\beta\), a function \(\text{name}\), and a set of all possible mappings \(\mathcal{M}\) defined in \(\mathcal{R}\) and \(\mathcal{m} \in \mathcal{M}\), \(\text{genPRSQL}\) generates SQL expression which projects only those relational attributes that correspond to distinct \(\mathcal{tp}\.\text{subject}, \mathcal{tp}\.\text{predicate}, \mathcal{tp}\.\text{object}\) and renames the projected attributes as \(\beta^\mathcal{M}(\mathcal{tp}, \text{subject}) \rightarrow \text{name}(\mathcal{tp}\.\text{subject}), \beta^\mathcal{M}(\mathcal{tp}, \text{predicate}) \rightarrow \text{name}(\mathcal{tp}\.\text{predicate}), \beta^\mathcal{M}(\mathcal{tp}, \text{object}) \rightarrow \text{name}(\mathcal{tp}\.\text{object}).

### Listing 5: \(\text{genPRSQL under } \mathcal{M}\)

```sql
1  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
2  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
3  genPRSQL^\mathcal{M}(\alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
4  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
5  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
6  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
7  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
8  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
9  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
10  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
11  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
12  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
13  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
14  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
15  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
16  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
17  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
18  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
19  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
20  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
21  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
22  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
23  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
24  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
25  genPRSQL^\mathcal{M}(\mathcal{TP}, \alpha, \beta, \text{name}, \mathcal{TM}, \mathcal{PreURI}, \text{Result}) :=
```

### 3.7 Query Rewriting Optimizations

The translation mappings/functions that we have presented so far can be evaluated directly over relational database systems (RDBMS). Taking into account that many of these RDBMS have already implemented a good number of optimisations for their query evaluation, most of the resulting queries can be evaluated at the same speed as the native SQL queries what would have been generated using SQL directly. However, there are RDBMS that need special attention (e.g., MySQL, PostgreSQL), because the resulting queries contain some properties that affect the capabilities of the database optimizer. For this reason, whenever it is possible, we remove non-correlated subqueries from the resulting queries by pushing down projections and selections (as advocated by \[10\]). We also remove self-joins that occur in the rewritten queries such as when the graph patterns contain alpha mappings coming from the same table(s). Those translations are illustrated in Listing 6.

### Listing 6: Examples of translation queries

```sql
1  SELECT DISTINCT \(\mathcal{pr}\), \(\mathcal{productLabel}\), \(\mathcal{productComment}\), \(\mathcal{productProducer}\)
2  FROM \{\}
3  \(\mathcal{SELECT p1\.pr, p1\.productLabel, p1\.productComment, p1\.productProducer}
4  FROM product p1\}
5  \(\mathcal{SELECT p2\.pr, p2\.productLabel, p2\.productComment, p2\.productProducer}
6  FROM product p2\}
7  \(\mathcal{SELECT p3\.pr, p3\.productLabel, p3\.productComment, p3\.productProducer}
8  FROM product p3\}
9  \(\mathcal{WHERE p1\.nr = p2\.nr AND p2\.nr = p3\.nr}
10  \(\mathcal{AND p1\.nr IS NOT NULL AND p2\.nr IS NOT NULL)
11  \(\mathcal{AND p2\.nr IS NOT NULL AND p3\.nr IS NOT NULL}
12  \(\mathcal{AND p1\.nr IS NOT NULL AND p3\.nr IS NOT NULL})
13  \(\mathcal{SELECT \mathcal{pr}, \mathcal{productLabel}, \mathcal{productComment}, \mathcal{productProducer})
14  \}
15  \(\mathcal{FROM \mathcal{product p}}\}
16  \(\mathcal{WHERE p1\.nr = p2\.nr AND p2\.nr = p3\.nr}
17  \(\mathcal{AND p1\.nr IS NOT NULL AND p2\.nr IS NOT NULL)
18  \(\mathcal{AND p2\.nr IS NOT NULL AND p3\.nr IS NOT NULL})
19  \(\mathcal{AND p1\.nr IS NOT NULL AND p3\.nr IS NOT NULL})
```

### 4. EVALUATION

We separate our evaluations into two categories, using a synthetic benchmark and using three real projects. Their details are available in the following subsections. Our query translation algorithm has been implemented in our latest version of Morph, which is available as a Java/Scala open-source project in Github\(^7\). The query translation types supported by Morph are the naïve translation queries (C), which

\[\text{https://github.com/priyanta/morph}\]

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\(^7\)https://github.com/priyanta/morph
are the result of the query rewriting algorithm described in Section 3, together with three variants of it; with subquery elimination (SQE), self-join elimination (SJE), and both types of eliminations (SQE+SJE).

### 4.1 Synthetic Benchmark Evaluation

The BSBM benchmark [5] focuses on the e-commerce domain and provides a data generation tool and a set of twelve SPARQL queries together with their corresponding SQL queries generated by hand. The data generator is able to generate datasets with different sizes containing entities normally involved in the domain (e.g., products, vendors, offers, reviews, etc). For the purpose of our benchmark, we work with the 100-million triple dataset configuration.

All these queries have been evaluated on the same machine, with the following configuration: Pentium E5200 2.5GHz processor, 4GB RAM, 320 GB HDD, and Ubuntu 13.04. The database server used for the synthetic benchmark queries is PostgreSQL 9.1.9. We normalize the evaluation time over the native evaluation time. We have run all queries with 20 times with different parameters, in warm mode run. The resulting sets of queries together with query plans generated by PostgreSQL 9.1.9, and the resulting query evaluation time are available at [http://bit.ly/15XSdDM](http://bit.ly/15XSdDM).

The BSBM SPARQL queries are designed in such a way that they contain different types of queries and operators, including SELECT/CONSTRUCT/DESCRIBE, OPTIONAL, UNION. In the same spirit, the corresponding SQL queries also consider various properties such as low selectivity, high selectivity, inner join, left outer join, and union among many others. Out of the 12 BSBM queries, we focus on all of the 10 SELECT queries (that is, we leave out DESCRIBE query Q09 and CONSTRUCT query Q12). We compare the native SQL queries (N), which are specified in the BSBM benchmark with the ones resulting from the translation of SPARQL queries generated by Morph. Although not included here, we also evaluated those queries using D2R 0.8.1 with the –fast option enabled. The reason why we do not include here the results from these evaluations is because in many queries (such as in Q2, Q3, Q4, Q7, Q8, and Q11), D2R produces multiple SQL queries and then the join/union operations are performed at the application level, rather than in the database engine, what prevents us from doing direct comparisons. Nevertheless, this approach is clearly not scalable (e.g., in Q07 and Q08 the system returned an error while performing the operations, while the native and the translation queries could be evaluated over the database system).

#### 4.1.1 Discussion

We can observe that all translation types (native, C, SQE, SJE, SQE+SJE) have similar performance in most of BSBM queries, ranging from 0.67 to 2.60 when normalized according to the native SQL queries. To understand this behaviour better, we analyzed the query plans generated by the RDBMS. Our observation tells us that in many of the queries (Q01, Q02, Q03, Q05, Q06, Q10, and Q11), C produces identical query plans to SQE’s, and SJE produces identical query plans to SQE+SJE’s. Additionally, SQE also produces identical query plans to SQE+SJE in Q07 and Q08. The reason for this is the capability of the database’s optimizer to eliminate non-correlated subqueries. The difference between the query plans produced by C/SQE and the ones produced by SJE/SQE+SJE is the number of joins to be performed. However, as the join conditions are normally performed on indexed columns, there is small overhead in terms of their performance. This explains why all the translation results have similar performance.

However, in some queries the translation results show significant differences, such as in Q04 and Q05.

- **BSBM SQL 4** contains a join between two tables (product and producttypeproduct) and three subqueries, two of them are used as OR operators. The SPARQL equivalent of this query is a UNION of two BGPs (a set of triple patterns). We note that the native query contains a correlated subquery and the generated query plan requires a table scan to find a specific row condition. The query plans generated by the translation algorithm, on the other hand, produce joins, instead of a correlated subquery, and the joins are able to exploit the indexes defined.

- **BSBM SQL 5** is a join of four tables (product, producttypeproduct, and productfeatureproduct). The size of table productfeatureproduct is significantly bigger than the table product (280K rows vs 5M rows). The generated query plan by the native query joins bigger tables (productfeatureproduct and productfeatureproduct) before joining the intermedi-
ate result with the smaller table (product and product). This join order is specified in the query itself. The join orders of the translation queries are different; C and SQE join based on the order of triple patterns in the graph, SJQ and SQE+SJE join based on the smaller tables first (which is an effect of the self-joins elimination process).

These explain why the translation queries perform better than the native queries in Q04 and Q05 and in fact show that the native queries proposed in the benchmark should have been better optimised when proposed for the benchmark.

4.2 Real Cases Evaluation

While the BSBM benchmark is considered as a standard way of evaluating RDB2RDF approaches, given the fact that it is very comprehensive, we were also interested in analysing real-world queries from projects that we had access to, and where there were issues with respect to the performance of the SPARQL to SQL query rewriting approach. In all the cases, we compare the queries generated by D2R Server with –fast enabled with the queries generated by Morph with subquery and self-join elimination enabled. All the resulting queries together with their query plans are also available at http://bit.ly/15XSdDM.

4.2.1 BizkaiSense Project

The BizkaiSense project is an effort to measure various environmental properties coming from sensors that are deployed throughout Greater Bilbao, Spain. Some of the most common queries that are needed in this project are:

- Q01 obtains all observations coming from a particular weather or air quality station together with the time of the observation. We set 100, 1000, and 10000 as the maximum number of rows to be returned.
- Q02 extends Q01 by returning the sensor results generated from those observations obtained. We set 100, 1000, and 10000 as the maximum number of rows to be returned.
- Q03 returns the average measures by property for a given week in a given station (with the units of measure).
- Q04 returns the average measures by property for a given week in a given station (without the units of measure).
- Q05 returns the maximum measure in all the stations for each property in a given day (returning also the station in which it happened).
- Q06 returns the maximum measure in all the stations for each property in a given day (without the station in which it happened).
- Q07 returns the maximum measure in all the stations for a given property in a given day (returning the station in which it happened) -avoiding the use of the "MAX" clause.

The corresponding SPARQL queries can be seen in Listing 7. All queries generated by D2R and Morph are evaluated on a machine with the following specification: Intel(R) Xeon(R) E5640 2.67GHz x 16, 48GB RAM, Ubuntu 12.04 LTS - 64 bits and MySQL 5.5 with a 10 minutes timeout. The evaluation time of those queries (in some cases with a limit in the number of returned bindings) is shown in Figure 4. We note that the database server failed to evaluate the queries Q03, Q05, Q06, and Q07 generated from D2R Server and in all cases, the queries generated by Morph take significantly less time to be evaluated compared to the ones generated by D2R Server.

4.2.2 RÉPENER Project

RÉPENER [17] is a Spanish national project that provides access to energy information using semantic technology called SEiS. Two of the most common queries that are consulted through SEiS are the following:

- Q01 retrieves all buildings and their climatezone and building life cycle phase
- Q02 retrieves all buildings and their climatezone, building life cycle phase, and conditioned floor area.

The corresponding SPARQL queries are shown in Listing 8. All queries generated by D2R and Morph are evaluated on a machine with the following specification: Intel Core 2 Quad Q9400 2.66 GHz, 4 GB RAM, Windows 7 Professional 32 bits, and MySQL 5.5 as the database server. As shown in Figure 5, all queries are successfully evaluated by the database server, with queries generated by Morph being 2-3 faster than those generated by D2R server.

4.2.3 Integrate Project

Integrate is an FP7 project for sharing and integrating clinical data using HL7-RIM [13], a model that represents

![Figure 4: BizkaiSense - query evaluation time (in seconds)](image)

![Figure 5: SEiS - query evaluation time (in seconds)](image)
entities and relationships commonly involved in clinical activities. Some commonly-used queries in this project are:

- Q01/Q02/Q03 are similarly structured, with a code that specifies whether to obtain tumor size, tumor stage, or pregnant women.
- Q04 obtains multiple participants who have been treated with Antracyclines.
- Q05 obtains demographic information (people that are older than 30 years old).

The corresponding SPARQL queries are shown in Listing 9. All queries generated by D2R and Morph are evaluated on a machine with the following specification: Intel QuadCore processor 2 GHz, 4 GB of RAM, 126GB SSD HD, Ubuntu 64-bits operating system and MySQL 5.6. The evaluation time can be seen in Figure 6. For Q01/Q02/Q03, the queries generated by D2R could not be evaluated by the database server because they returned an error message saying that too many tables needed to be joined. For the other queries, the queries generated by Morph were evaluated in less time than the ones generated by D2R.

Figure 6: Integrate - query evaluation time (in seconds)

5. CONCLUSION AND FUTURE WORK

In this paper we have shown that the query translation approaches used so far for handling RDB2RDF mapping languages may be inefficient. For instance, systems like D2R normally produce multiple queries to be evaluated and then perform the join operations in memory rather than in the database engine, or generate SQL queries where the number of joins inside the query is so large that the underlying database engine cannot evaluate them, producing an error.

We have proposed an extension of one of the most detailed approaches for query rewriting, Chebotko's, which was not originally conceived for RDB2RDF query translation but for RDBMS-backed triple stores. Now we consider R2RML
There is still room for improvement in our work, which we will address in the near future. For instance, some of the optimised translated queries perform better than the native ones, due to the introduction of additional predicates, what is common in the area of Semantic Query Optimization (SQO). In this sense, we will deepen in the design of our algorithm so as to take into account other SQO techniques that can be useful in query translation. For instance, predicate introduction may speed up the query evaluation process by exploiting more indexes. Our evaluation setup will be also made available as a service for other researchers to use, so that they can evaluate their R2RML query rewriting implementations with low effort in a number of RDBMSs.

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6. MAPPING DOCUMENT EXAMPLE

```sql
#Q01/Q02/Q03
SELECT DISTINCT ?subject WHERE {
  FILTER (isType(?subject) = "Patient")
}
WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}
```

#Q01 to obtain tumor size of patients, code value is 263605001
#Q02 to obtain tumor stage of patients, code value is 58790005
#Q03 to obtain tumors that are/have been pregnant, code value is T7386006
#Q4 to retrieving multiple participants who have been treated
#with Anticancer drugs

SELECT WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

#Q05 to obtain patient demographics information: people older than 30 years

SELECT DISTINCT ?subject ?birthdate WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

#Q06 to obtain images

WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

#Q07 to obtain information where a diagnosis are based in

WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

#Q08 to obtain diagnosis

WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}

#Q09 to obtain observation

WHERE {
  ?subject rdfs:isDefinedBy ?value.
  FILTER (isType(?value) = "Code")
}
7. REFERENCES


