Duplicate-Aware Federated Query Processing over the Data Web

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ABSTRACT

Over the last years, the Linked Data Web has developed into a large compendium of interlinked data sets from diverse domains. Due to this distributed character, several of these data sets contain either duplicate or complementary data. So far, several federated querying approaches have been developed to retrieve information from these data sources. Still, only little attention has been paid to the effect of duplicates and fragmented data on the ranking of data sources during the processing of federated queries. This work presents a novel approach federated querying based on min-wise independent permutation vectors applied to RDF predicates. Our approach allows evaluating the ranking of data sources accurately while taking into account duplicate data. By these means, our approach achieves significantly smaller mean squared errors that the state of the art with respect to the approximation of result set sizes as well as in the ranking of data sources.

Categories and Subject Descriptors

General Terms

Algorithms, Experimentation, Theory

Keywords

Federated queries, SPARQL, deduplication

INTRODUCTION (AN+MS)

Over the last years, the Linked Data Web has developed into a large compendium of interlinked data sets from diverse domains[14]. One of the central principles underlying

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the architecture of these data sets is the reuse of URIs and vocabularies as well as the linking of knowledge bases [20]. One of the results of this architectural choice is that certain queries can only be answered by retrieving information from several knowledge bases. This type of querying, called federated querying, is of central importance for manifold applications such as question answering, knowledge retrieval and data integration. In addition to the information necessary to answering queries being distributed, certain pieces of information (i.e., triples) can be found in several knowledge bases. For example, the name of movie directors can be found both in DBpedia and LinkedMDB. The authors of papers can be found in both the ACM and DBLP libraries. Similarly, we have noticed that $DrugBank^1$, $Neurocommons^2$ datasets are duplicated at DERI health Care and Life Sciences Knowledge Base³. In our further analysis of linked data cloud, it is noticed that duplicates are more common in life sciences domain. The possibility of the duplicates records between different life sciences dataset is shown in figure 1. We call triples that can be found in several knowledge bases across the Web of Data duplicates.

H.2.4 [Database Management]: Systems—Distributed databases While the importance of federated queries over the Web of Data has been stressed in previous work, the impact of duplicates has not received much attention. Yet (as we will show in the remainder of this paper), a duplicate-aware approach to query processing can lead to more time-efficient and effective algorithms for federated queries. In this paper, we address this drawback by presenting a duplicate-aware approach for query processing based on min-wise independent permutation (MIPs) vectors. Our approach is able to predict the amount of new information contained in a knowledge base with a higher accuracy than the state of the art, leading to a better performance with respect to source ranking and result set size optimization. In the rest of this paper, we aim to show experimentally that our approach outperforms the state-of-the-art by running it against two other approaches on a benchmark of 79 queries. We begin by giving a brief overview of the state of the art in federated query processing. In addition, we argue for the use of MIPS for the approximation of result size sets. Thereafter, we present our

 $^{^1}$ http://datahub.io/dataset/fu-berlin-drugbank

²http://neurocommons.org/page/RDF_distribution

³http://hcls.deri.org:8080/openrdf-sesame/ repositories/hclskb

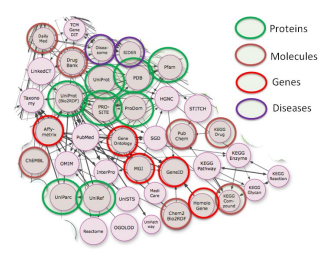


Figure 1: Possible duplicates in life sciences domain

duplicate-aware federated query processing approach. After an overview of the datasets used in this paper, we present experimental results which corroborate the superior efficiency of our approach. We conclude the paper with a discussion of our findings and an overview of future work.

2. RELATED WORK (MS)

2.1 Federated SPARQL queries

The research work on federated query processing over the Data Web can be categorized into two main directions:

- Service description/index-assisted approaches, which
 make use of the dataset summaries that have been
 collected in a pre-processing stage. These approaches
 lead to a more efficient query federation. However, the
 index needs to be constantly updated to ensure up-todate results. Also, the index size should not be large
 to ensure that it does not increase the overall query
 processing time.
- Index-free approaches, in which the query federation is performed without using any stored data summaries. This approach is less efficient however; the query response time is faster than index assisted approach.

Both of the above approaches can further be divided into two categories (a) Query federation with complete result retrieval (b) Query federation with partial (sufficient) record retrieval. In our case, we are focusing on index-assisted which can be used both for complete and partial record retrieval.

Heiner et. al[1] describes how to extend the Sesame RDF repository to support distributed SeRQL queries over multiple Sesame RDF repositories. They use a special index structure to determine the relevant sources for a query. Quilitz and Leser[2] uses the theoretical knowledge of[1] to develop the first federated query engine (named DARQ) for remote RDF data sources. DARQ uses service descriptions for relevant data source selection. A service description describes the capabilities of a SPARQL endpoint. They used query rewriting mechanism based on[8] and a cost-based optimization algorithm to reduce the query processing time with the

minimum bandwidth usage. DARQ is compatible with any SPARQL endpoint that implements the SPARQL standard. Their experimental results show that the optimization algorithm can greatly reduce the query processing time.

Andreas et al.[3] proposed a solution similar to DARQ using a mediator approach. All the SPARQL endpoints need to register first with the mediator using HTTP POST requests with an RDF document attached. The mediator continuously monitors the SPAQL endpoints for any dataset changes and updates the service descriptions automatically. Unlike DARQ, the service descriptions remain up-to-date all time

Olaf et al.[4] present a new approach for federated query processing over the Web of Data. Their approach discovers data that might be relevant for answering a query during the query execution itself. The discovery of relevant data is accomplished by traversing RDF links. They used an iterator-based pipeline and a URI prefetching approach for efficient query execution. Both DARQ and [3] are not able to discover relevant data sources by the query itself.

The previous techniques cannot answer all types of queries due to index or service description limitations. Umbrich et al.[5], [6] proposed a Qtree-based index structure which summarizes the content of data source for query execution over the Web of Data. They concluded that this approach is able to handle more expressive queries and return more complete results than previous approaches. Schwarte et al.[7] proposed an index-free query federation for the Web of Data. Their approach gives a reasonably fast data retrieval as compared to all the previous techniques. However, since they do not keep any statistics, it works only for limited data sources.

Li and Heflin[15] build a tree structure which supports federated query processing over multiple heterogeneous sources. They make use of an OWL reasoner to answer queries over the selected sources and their corresponding ontologies. Kaoudi et al.[16] propose a federated query technique on top of distributed hash tables(DHT). The overall goal of this technique is to minimize the query execution time and the bandwidth consumption by reducing the cardinality of intermediate results. The DHT-based optimizer make use of three greedy optimization algorithms for best plan selection.

Ludwig and Tran[18] proposed a mixed query engine; that assumes some incomplete knowledge about the sources to select and discover new sources at run time. A Symmetric Hash Join is implemented to incrementally produce answers. This approach was extended to query both remote and local linked data[19].

Maribel et al.[21] presented ANAPSID, an adaptive query engine that adapts query execution schedulers to SPARQL endpoints data availability and run-time conditions. ANAP-SID provides physical SPARQL operators that detect when a source becomes blocked or data traffic is bursty. The operators produce results as quickly as data arrives from the sources. Amol et al.[17] presents a survey of the adaptive query processing techniques, common issues, and the settings in which each piece of work is most appropriate. Avalanche[24] gathers endpoints dataset statistics and bandwidth availability on-the-fly before the query federation. The advantage of this technique is the live query execution over the Web of Data. However, on-the-fly statistics calculation can increase the overall query execution time. Olaf[25] proposed a heuristics-based technique to minimize query intermediate results.

Other noteable contributions LIMES[26], SILK[27], EA-GLE[28] provides time-efficient approaches for Link Discovery using various distance measure techniques. LIMES framework utilize the mathematical characteristics of metric spaces to compute similarity estimates between instances. It makes use of the triangle inequality to partition the metric space into different portions. Each of these portions of the space is then represented by an exemplar [33] that allows to compute an accurate distance approximation between different instances. SILK implements a multi-dimensional blocking approach for link discovery. The Levenshtein distance[34] is used for approximate for similarity estimation between different instances. EAGLE presents a novel active learning approach to learning link specifications based on genetic programming. EAGLE generates highly accurate link specifications while reducing the annotation burden for the user.

None (to the best of our knowledge) of the above techniques has taken into account the possibility of mutual overlap between different RDF datasets before query execution. Like DARQ engine, we use service descriptions for query federation. In addition, we store MIPs vector statistics for each of the indexed data source. Query federator makes use of this special statistic information to estimate the possible overlap between the results of different sub-queries.

2.2 Dataset Overlap Estimation

Avoiding the retrieval of duplicate records is a crucial issue in large scale distributed datasets. As the SPARQL endpoints are autonomous, there is always the possibility of high mutual overlap between different datasets residing in different endpoints. It is thus likely that an optimized query execution plan contains a sub-query that retrieves records which are already retrieved by another sub-query from a different SPARQL endpoint. In this case it is useless to execute the former sub-query, as it would not only result in duplicate records being retrieved but also increase the total processing time. The overlap between two datasets can be estimated using min-wise independent permutations[9], bloom filters[10], or compressed bloom filters[11]. Bender et al. [12] also proposed a technique for predicting the query-specific mutual overlap between peers. However, MIPs have proved to be the more accurate overlap estimation with small bandwidth consumption[9].

3. NOTATION

In this section, we present the core of the notation that will be used throughout this paper. We denote data sources with S and the total number of data sources with n. The set of all data sets is dubbed S. The set of all possible result sets is denoted R while the set of all possible SPARQL queries is labeled with Q. A data source ranking function $rank: S \times Q \rightarrow \{1 \dots n\}$ is a function that assigns a ranking to each data source given a particular query $q \in Q$. Note that for any source ranking function rank, we assume that $\forall S, S' \forall q \in Q : S \neq S' \iff rank(S,q) \neq rank(S',q)$. A result set estimation function $est: S \times Q \rightarrow \mathbb{N}$ aims at approximating the size of the result set that will be returned by a given query. Note that this function plays a crucial role in the processing of federated queries as it is most commonly used to decide upon the ranking of data sources for a given query. The aim of a federated query system such as the one described in this work is thus to optimize its estimation

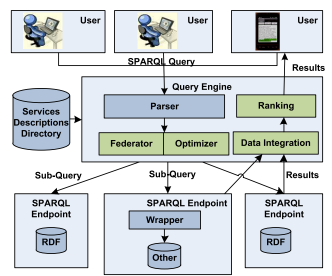


Figure 2: Federated SPARQL query processing architecture

function *est* so as to ensure a ranking of the source close to the optimal ranking.

4. APPROACH (MS)

4.1 Overview

Overall, our approach works as follows: The user begins by issuing a SPARQL query using any input form. The query is then forwarded to the query engine. Upon receiving, it is parsed to get all triples. The parser rewrites the original query in an execution-optimized form. The federator then makes use of the services descriptions directory/index to decompose single query into multiple sub-queries. The query optimizer takes all sub-queries to generate an optimize execution plan. At last, all the optimized sub-queries are forwarded to the relevant SPARQL endpoints. The results of each sub-query execution are integrated and finally sent back to the user. This architecture for distributed query processing is shown in figure 2.

While this architecture is the same for all federated query systems, the federator of our system differ significantly from those of other approaches as it make use of the MIPs statistical information to generate duplicate-free query execution plan. The generation of MIPs statistical information is explained in the next section.

4.2 Service Description Directory

To find the set of relevant servers and generate multiple sub-queries, the query engine needs information about the different data sets available at different servers. We call this catalog the *service description directory*. Each service description provides a declarative description of the data available from a server along with statistical information.

Our service description directory stores server url, graph name, and for each distinct predicate p we record predicate name, MIPs vector, subject selectivity, object selectivity and total number of triples having predicate p. The MIPs vectors are used to estimate the dataset overlap among the capable datasets before the sub-query distribution. The subject and object selectivity are used for sub-query result set estima-

Listing 1: A Service Description Example

```
[] a sd:Service;
sd:url <a href="http://localhost:8890/sparql">sd:graph ''diseasome/DS1'';
sd:capability [
sd:predicate diseasome:name;
sd:MIPv ''-6908232 -7090543 -6892373 -7064247 ...'';
sd:subjectSelectivity ''0.0068'';
sd:objectSelectivity ''0.0069'';
sd:triples 147;
];
sd:capability [
sd:predicate diseasome:chromosomalLocation;
sd:MIPv ''-7056448 -7056410 -6845713 -6966021 ...'';
sd:subjectSelectivity ''0.0062'';
sd:subjectSelectivity ''0.0062'';
sd:objectSelectivity ''0.0072'';
sd:triples 160;
];
```

tion when *subject*, *object* are bound respectively. A sample service description is given in the listing 1.

It is very crucial to keep the index size enough to minimize pre-query federation time. However, it must contain sufficient information to enable sub-query generation, optimization and pre-processing dataset overlap estimation. Our index size is mostly depended upon the size of the MIPs vectors which can be adjusted to any length. The generation of the MIPs vectors is explained in the next section.

4.3 MIPs Operation

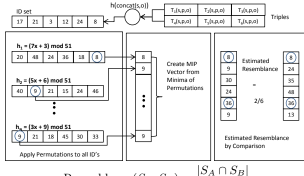
Min-Wise Independent Permutations (MIPs) assumes that the set elements can be ordered and computes N random permutations of the elements. Each permutation uses a linear hash function of the form $h_i(x) := a_i * x + b_i \mod U$ where U is a big prime number and a_i , b_i are fixed random numbers. By ordering the resulting hash values, we obtain a random permutation. For each of the N permutations, the MIPs technique determines the minimum hash value, and stores it in an N-dimensional vector, thus capturing the minimum set element under each of these random permutations. The technique is illustrated with an example in figure 3. Its fundamental rationale is that each element has the same probability of becoming the minimum element under a random permutation. By using sufficiently many different permutations, we can approximate the set cardinality.

An unbiased estimate of the pair-wise resemblance of sets using their N-dimensional MIPs vectors is obtained by counting the number of positions in which the two vectors have the same number and dividing this by the number of permutations N. Essentially, this holds as the matched numbers are guaranteed to belong to the intersection of the sets.

For a predicate $p \in S$, the set elements are all triples $T \in S$ having predicate p. We concatenate the *subject* and *object* of each triple in T and apply a hash function to get the ID set. Finally, we computes N random permutations of the ID set. The resulting MIP vector is stored against each capability of the service description. Note that we are not applying hash function to the complete triple because the predicate is already stored with each service capability and source selection is bound by query predicate.

4.4 Source Selection

A SPARQL query contains one or more basic graph patterns. The query planning step is performed separately for each basic graph pattern. The source selection algorithm used in DARQ for a query simply matches all triple patterns



Resemblance $(S_A, S_B) = \frac{|S_A \cap S_B|}{|S_A \cup S_B|}$

Figure 3: Min-Wise independent permutations

against the capabilities of the data sources. The matching compares the predicate in a triple pattern with the predicate defined for a capability. Let $S = \{(s_1, C_1), ..., (s_n, C_n)\}$ be a set of all indexed data sources $s_1..s_n$ and their capabilities $C_1..C_n$, where $C_i = \{(p_i, mipsv_i, sbjsel_i, objsel_i, totTrpl_i)\}$.

Let BGP be a set of triple patterns in a filtered basic graph pattern. The result of the source selection is a set of data sources D_j for each triple pattern $t_j = (s_j, p_j, o_j) \in BGP$ with

$$D_i = \{ s | (s,C) \in S \land \exists p_i \in C \}$$

We extend the DARQ source selection algorithm by comparing MIP Vectors for similarity. For a triple pattern, we compare the MIP vectors of all the relevant data sources for possible similarity. A specific source s is only added to the set of capable sources D_j , if it is able to produce new results above a specific threshold value defined for similarity. For a triple $t \in BGP$ and dataset $s \in S$, if the value of the new record generated is less than the threshold, s is ignored from set D_j . For 100% recall the threshold value should be 1. For sufficient (not complete) record retrieval the threshold can be fixed to any number greater than 1. For all of the 79 queries in this work, our threshold value is 1.

4.5 Source Ranking

4.5.1 Triple Pattern Source Ranking

Algorithm 1 shows the single triple pattern source ranking. For a triple $t(s,p,o) \in T$ the set of capable data sources D is obtained using the source selection algorithm explain in previous section. A source $d \in D$ which has the highest tot-Trpl value in the matched capability C(p,mipsv,sbjsel,objsel,totTrpl) is selected as first rank. The union and intersection of MIPs Vectors is explained in section 4.3. In order to get the accurate result set estimate, we multiply sbjsel, $objsel \in C$ to the orignal size of the MIPs Vectors if subject or object of the triple pattern is bound respectively. The same is true for rank 1 dataset selection. However, the multiplication is performed with totTrpl of mached capability instead of orignal size.

It is important to note that the number of elements in the $MIPs\ Vector$ may be different from its $orignal\ size$. The $orignal\ size$ is the number of elements in the $ID\ set$ over which we applied N random permutations to get the $MIPs\ Vector$ as explained in MIPs operation. In line 24 we set the threshold value of 1 is chosen to get 100% recall. For each

Algorithm 1 triple pattern source ranking

```
Require: t(s,p,o) \in T //triple pattern
 1: D = getCapableSources(t) //all sources capable of an-
    swering triple t
 2: rank_1 Source = getMaxSizeSource(D,t)
3: rnkNo = 1
4: unionMipv = rank_1Source.getMIPv(t)
 5: D_s[\text{rnkNo}] = \text{selectedSource}
 6: D = D - {selectedSource}
 7: rnkNo = rnkNo + 1
 8: while D \neq \emptyset do
9:
      selectedSource = null
10:
      \max DistinctRecords = 0
11:
      for each d_i \in D do
        mipv = d_i.getMipv(t)
12:
13:
        if s is bound in t then
           mipv.orignalSize = mipv.orignalSize *d_i.getSbjSel(t)
14:
15:
        else if o is bound in t then
           mipv.orignalSize = mipv.orignalSize *d_i.getObjSel(t)
16:
17:
        overlap = unionMipv.intersectionSize(mipv)
18:
19:
        distinctRecords = mipv.getOrignalSize()-overlap
20:
        if distinctRecords > maxDistinctRecords then
21.
           selectedSource = d_i
22:
           maxDistinctRecords = distinctRecords
23:
        end if
24:
      end for
25:
      if maxDistinctRecords \geq 1 then
26:
         D_s[\text{rnkNo}] = \text{selectedSource}
        selectedMipv = selectedSource.getMipv(t)
27:
28:
        unionMipv = unionMipv.union(selectedMipv)
29:
        rnkNo = rnkNo + 1
30:
      end if
      D = D - \{selectedSource\}
31:
32: end while
33: return D_s //rank wise selected sources
```

triple pattern $t_j = (s_j, p_j, o_j)$ in query $q \in Q$, we apply the algorithm 1 to get a set of selected capable data sources D_{s_j} which is used in sub-query generating algorithm explained in the section 4.6.

4.5.2 Join Source Ranking (this section is not well written)

Same like DARQ, we perform individual triple based subquery federation. Consequently, triple pattern source ranking is most important for sub-query escaping. The advantage of this approach is that if a data source is not capable of answering the complete query, it may produce enough results for a specific sub-set of query triples. Just for the comparison we proposed complete query source ranking which works as follow; for a star shaped queries the over all result set is bound by the triple with lowest number of estimated records. Therefore, we perform source ranking for that triple (using algorithm 1) and use this as complete query ranking. The lowest records query triple can be estimated using the rank 1 source selection explained in previous section. However, this time we are looking for a source which has the lowest estimated value of all. For path shaped queries the total records are bound by the triple which has subject variable equal to object of the previous query triple. For example

Algorithm 2 sub-query generation

```
Require: T = \{ t_1, ..., t_n \}
    S = \{D_{s_1},...,D_{s_n}\} // sets of data sources obtained from
    algorithm 1
 1: sub-queries = \emptyset, seperateSub-queries = \emptyset
2: for each t_i \in T do
 3:
      if D_{s_i} = \{s\} then
         q = queries.getQuery(s)
 4:
 5:
         if q not null then
 6:
           q.T = q.T + t_i
 7:
         else
 8:
            sub-queries = sub-queries + (\{t_i\},s)
9:
         end if
10:
       else
11:
         for each s_j \in D_{s_j} do
12:
            seperateSub-queries = seperateSub-queries +
            (\{t_i\},s)
13:
         end for
14:
       end if
15: end for
16: return sub-queries \cup seperateSub-queries //Return all
    sub-queries
```

the result set in the query given in listing 2, is bound by the second triple in which the *object* variable in the first triple is used as *subject* in the second triple. Therefore, we perform the triple pattern ranking for second triple and use this as ranking for complete query.

Listing 2: A P-1 query example

4.6 Sub-query generation

The sub-query generation algorithm is same as DARQ. However, instead of using all capable sources, we use selected data sources (obtained from algorithm 1) for sub-query generation. Let (T,s) is a sub-query where T is a set of triple patterns, and s is the data source that can answer this sub-query. The sub-query generation is shown in algorithm 2. If a triple pattern t_j matches exactly one data source $(D_{s_j} = \{s\})$ he the triple will be added to the set of a sub-query for this data source. All triples in this set will be sent to the data source in one sub-query. If a triple matches multiple more than one data sources then the triple will be sent individually to all matching data sources in separate sub-queries.

5. EXPERIMENTS AND RESULTS

The goal of our experiments was to quantify how well our duplicate-aware solution approximates the result sets from each of the data source from which it can choose and how this approximation affects the way the data sources are ranked. For this purpose, we compared our approach with two other approaches: DARQ and the optimal duplicate-unaware solution to result set estimation and query ranking.

Dataset	BGP	S-1	S-2	P-1	P-2	P-3	Total
Diseasome	5	5	5	4	5	2	26
Geo	5	5	5	0	0	0	15
LinkedMDB	5	0	0	0	0	0	5
Publication	5	5	5	7	7	4	33
Total	20	15	15	11	12	6	79

Table 2: Distribution of query types across datasets

In the following, we describe our experimental setup in detail and contrast the results achieved by each of the approaches. All data used is either publicly available or can be found at the project webpage.⁴

5.1 Experimental Setup (AN)

5.1.1 Datasets

Within our experiments, we used the four datasets described in Table 1. We chose data sets of different data sizes to gain some insights on the scalability of our approach. The size of the MIPS vectors was set to achieve a compression ratio of approx. 98% greater than the previous work [5]. In addition, we emulated the difference between the number of relevant triples across datasets by introducing the discrepancy constant d. Given n data sources of sizes $|S_i|$

$$d = \max_{1 < i < n} |S_i| - \min_{1 < j < n} |S_j|. \tag{1}$$

Note that the discrepancy accounted for between ca, 1.6% and 2.8% of the triples and was to increasingly higher values for increasingly larger data sets. The number of slices was set to 10 in all experiments.

5.1.2 Queries

We used three main types of queries. Simple basic graph pattern (BGP) queries consisted of exactly one triple pattern in the query. Star-shaped and path-shaped queries were defined as in [5]: star-shaped queries (denoted S-k) have one variable as subject and contained k triple patterns. Path-shaped queries, denoted P-k on the other hand were generated by using a random walk approach to generate a path of length k. Note that previous work has shown that these query shapes are the most common shapes found in real-world RDF queries [35]. Overall, our benchmark data consisted of 79 queries as shown in Table 2. Note that some query shapes could not be used on certain datasets due to the topology of the ontology underlying the datasets not being permissive for such queries. For example, S-2 queries could not be sent to LinkedMDB because

5.1.3 Metrics

We used two measures to compare our approach with the state of the art. First, we measured the mean squared error of each approach with respect to their result set evaluation as follows: For each query q_i , each source S_i contained a number $R_i(q_i)$ of non-duplicate results. Each of the approaches generated an approach $\tilde{R}_i(q_i)$ of $R_i(q_i)$. The mean squared error with respect to the result set size (short RSMSE) was then computed as follows:

$$RSMSE = \frac{1}{|Q|} \sum_{q_i \in Q} |R_i(q_i) - \tilde{R}_i(q_i)|^2.$$
 (2)

In addition to the MSERS, we also computed how well the approaches were able to rank the data sources with respect to the unseen triples they would return for a given query. Given the ideal ranking $rank(S_i, q_j)$ of each data source S_i for each query q_j , we computed the mean squared ranking error (short RankMSE) of an approach characterized by the the ranking function $\tilde{r}(S_i, q_j)$ by using the following formula:

$$RankMSE(q_j) = \frac{1}{|\mathfrak{S}|} \sum_{S_i} |rank(S_i, q_j) - \tilde{r}(S_i, q_j)|^2.$$
 (3)

$$RankMSE = \frac{1}{|Q|} \sum_{q_j \in Q} RankMSE(q_j). \tag{4}$$

For both measures, higher values are obviously worse.

5.2 Results

5.2.1 Result set estimation and ranking

The RSMSE results are shown in Figures 4 while the ranking results are shown in Figure 5. In each figure, we compared the results of the approaches w.r.t. both the whole input query and to each of the triples patterns contained in the input queries. Our results clearly show that we outperform the state of the art w.r.t to both result set estimation and ranking. Especially, our estimation of the result set sizes is approximately 4 orders of magnitude better than DARQ's or even DU's. This is due to our approach being able to capture the existence of duplicates in the data sets and thus to better approximate the number of new results that can be expected from an endpoint. As expected, the ranking error for datasets decreases when carrying the ranking of data sources at triple pattern level instead of query level (see Figures 4b and 4a). This is simply due to the overall error increasing when combining the several triple patterns to a query. The same overall behavior is also displayed when considering the result set estimation error (see Figures 5b and 5a). Interestingly, the approximation of our approach is slightly poorer on star-shaped queries (approx. 0.2 on S-2 queries). This is a result of our approach to generating the estimations for whole query, which is carried out by using the lower bound of the estimation of single query patterns in the query. Small errors in the estimation of the size of the result size can thus lead to minor absolute errors.

As stated before, we used a compression ratio of approx. 98% across all experiments. The main advantage of such a high compression ratio is the small size of the index. Yet, the higher the compression ratio, the larger the error margin of the ranking function. When given a query whose sub-queries lead to very similar result set estimations, this larger error margin can lead to substantial RankMSE and RSMSE rates. We thus studied the effect of decreasing the compression ratio on the RSMSE and RankMSE for such a query. Our results in Table 3 suggest that MIPS vectors can successfully detect even small differences in the sizes of the result sets when given a sufficiently small compression ratio

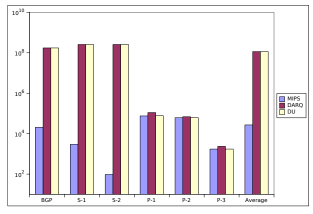
5.2.2 Recall and Runtime

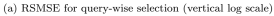
Table 4 shows the recall achieved by MIPS and DARQ in relation to the number of slices taken into consideration. Our results display the superiority of our ranking function, which is always able to achieve a recall of 1 by querying

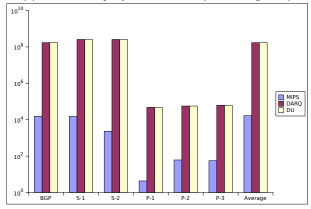
⁴http://???

Dataset	Size	Summary	CR	Slices	Discrepancy	Duplicate	Duplicate	Total	Sum. Gen.
	(MB)	Size (MB)	(%)			Slices	Datasets	Triples	Time (sec)
Diseasome	18.6	0.64	96	10	1,500	1	10	91,122	9
Publication	39.0	1.15	97	10	2,500	1	10	234,405	16
Geo Coordinates	274.1	4.66	98	10	50,000	2	5,8	1,900,006	1302
LinkedMDB	448.9	7.9	97	10	100,000	1	2	3,579,616	1837

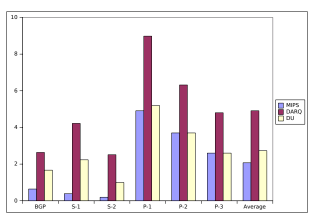
Table 1: Overview of datasets. CR stands for Compression Ratio.

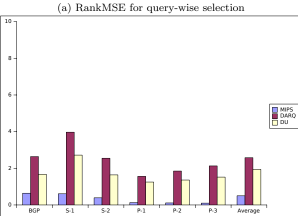






(b) RSMSE for subquery-wise selection (vertical log scale) Figure 4: Mean squared errors with respect to result set evaluation





(b) RankMSE for subquery-wise selection Figure 5: Mean squared errors with respect to ranking

Compression Ratio	RSMSE	RankMSE
97%	253.9×10^{6}	3.4
94%	33.3×10^{6}	0.6
91%	0	0

Table 3: Effect of compression ratio

Dataset	BGP	S-1	S-2	P-1	P-2	P-3	Total
Diseasome	7	21	21	8	20	12	89
Geo	10	15	23	0	0	0	48
LinkedMDB	5	0	0	0	0	0	5
Publication	1	2	5	0	0	0	8
Total	23	38	49	8	20	12	150

Table 5: Distribution of sub-query escaped by our approach

9 from 10 data sources. In contrast, DARQ always has to visit all data sources to achieve a comparable recall. We measured the number of sub-query executions that could be escaped by our approach without any loss in recall. The results shown in Table 5 clearly display that our approach can effectively reduce the total number of queries executed overall. Especially, on BGP queries, we were able to escape up to 10 subqueries. Our approach was most successful on star-shaped queries, where up to 21 subqueries could be escaped. The better ranking function also leads to a superior overall runtime as shown in Figure 6. In average, our approach requires solely between 79.85% (S-2 queries) and 93.46% (P-1 queries) of the runtime of DARQ to retrieve the complete results to each query.

6. DISCUSSION (AN)

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Rank	В	GP	S1		S2		P1		P2		P3	
	MIPS	DARQ	MIPS	DARQ	MIPS	DARQ	MIPS	DARQ	MIPS	DARQ	MIPS	DARQ
1	29,22	19,22	26,53	24,42	30,69	30,69	18,12	15,67	24,60	21,99	31,60	29,30
2	49,61	$39,\!50$	46,20	$39,\!67$	52,15	49,70	34,04	31,49	$38,\!84$	$38,\!56$	43,78	45,02
3	$67,\!48$	61,78	60,90	54,78	$66,\!48$	59,96	50,28	49,23	55,74	53,16	$61,\!86$	56,76
4	$78,\!44$	72,68	70,75	$63,\!86$	76,60	71,49	65,71	64,04	$72,\!25$	69,11	80,17	73,72
5	$85,\!65$	79,26	79,76	74,32	83,64	80,54	77,24	79,87	83,04	86,11	$90,\!45$	91,74
6	$90,\!54$	84,00	86,67	$79,\!59$	89,90	86,76	82,28	83,75	87,69	89,95	$94,\!48$	95,70
7	94,68	87,25	92,94	85,61	94,88	90,22	$85,\!56$	87,17	89,99	92,08	$95,\!59$	96,73
8	99,40	89,38	98,62	89,98	$99,\!35$	93,43	87,71	89,29	$91,\!47$	93,31	95,70	97,53
9	100,00	97,95	100,00	$96,\!52$	100,00	97,06	100,00	94,97	100,00	97,15	100,00	99,26
10	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00

Table 4: Comparison of recall on top-k data sources

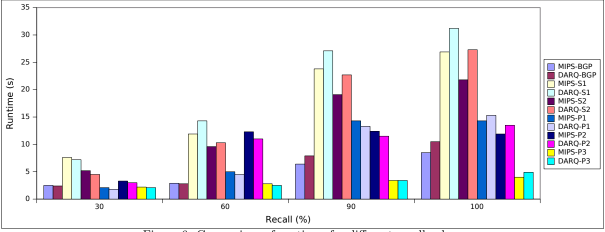


Figure 6: Comparison of runtimes for different recall values

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