A Large Scale Taxonomy Mapping Evaluation

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Abstract. Matching hierarchical structures, like taxonomies or web directories, is the premise for enabling interoperability among heterogenous data organizations. While the number of new matching solutions is increasing the evaluation issue is still open. This work addresses the problem of comparison for pairwise matching solutions. A methodology is proposed to overcome the issue of scalability. A large scale dataset is developed based on real world case study namely, the web directories of Google, Looksmart and Yahoo!. Finally, an empirical evaluation is performed which compares the most representative solutions for taxonomy matching. We argue that the proposed dataset can play a key role in supporting the empirical analysis for the research effort in the area of taxonomy matching.

1 Introduction

Taxonomic structures are commonly used in file systems, market place catalogs, and the directories of Web portals. They are now widespread as knowledge repositories (in this case they can be viewed as shallow ontologies [20]) and the problem of their integration and interoperability is acquiring a high relevance from a scientific and commercial perspective. A typical application of hierarchical classification interoperability occurs when a set of companies wants to exchange products without sharing a common product catalog. The typical solution to the interoperability problem amounts to performing matching between taxonomies. The Match operator takes two graph-like structures as input and produces a mapping between the nodes of the graphs that correspond semantically to each other.

Many diverse solutions to the matching problem have been proposed so far, for example [13, 14, 6, 17, 21, 3, 18, 16, 8], etc. Unfortunately nearly all of them suffer from the lack of evaluation. Until very recently there were no comparative evaluations and it was quite difficult to find two systems which were evaluated on the same dataset. At the same time the evaluation efforts were mostly concentrated either on datasets artificially synthesized under questionable assumptions or on the "toy" examples.

In this paper we introduce a large scale dataset for evaluating matching solutions. The dataset is constructed from the mappings extracted from real web directories and contains thousands of mappings. We have evaluated the dataset using the most representative state of the art solutions to the matching problem. The evaluation highlighted that the dataset has four key properties namely Complexity, Discrimination capability, Incrementality and Correctness. The first means that the dataset is "hard" for state of the art matching systems, the second that it discriminates among the various matching solutions, the third that it is effective in recognizing weaknesses in the state of the art matching systems and the fourth that it can be considered as a correct tool to support the improvement and research on the matching solutions. At the same time the current version of dataset contains only "true positive" mappings. This fact limits the evaluations on the dataset to measuring only Recall. This is a weakness of the dataset that we plan to improve. However, as highlighted in [15], the biggest problem in nowadays matching systems is recall, while completeness is much less of an issue.

The rest of the paper is organized as follows. Section 2 summarizes the definition of the matching problem and recalls the state of the art. Section 3 expands more on the notion of mapping evaluation problem. Section 4 illustrates how the large scale dataset has been arranged. Section 5 is devoted to a large scale empirical evaluation on two leading matching systems. Section 6 presents the results of our experiments and argues why the proposed dataset is of interest. Section 7 concludes the paper.

2 The Matching Problem

In order to motivate the matching problem and illustrate one of the possible situations which can arise in the data integration task let us use the two taxonomies A and B depicted on Figure 1. They are taken from Yahoo! and Standard business catalogues. Suppose that the task is to integrate these two taxonomies.

We assume that all the data and conceptual models (e.g., classifications, database schemas, taxonomies and ontologies) can be represented as graphs (see [9] for a detailed discussion). Therefore, the matching problem can be represented as extraction of graph-like structures from the data or conceptual models and matching the obtained graphs. This allows for the statement and solution of a generic matching problem, very much along the lines of what done in [14, 13].

The first step in the integration process is to identify candidates to be merged or to have relationships under an integrated taxonomy. For example, $Computer_Hardware_A$ can be assumed equivalent to $Computer_Hardware_B$ and more general than $Personal_Computers_B$. Hereafter the subscripts designate the schema (either A or B) from which the node is derived.

We think of a mapping element as a 4-tuple $\langle ID_{ij}, n1_i, n2_j, R \rangle$, $i = 1, ..., N_1$; $j = 1, ..., N_2$; where ID_{ij} is a unique identifier of the given mapping element; $n1_i$ is the i-th node of the first graph, N_1 is the number of nodes in the first graph; $n2_j$ is the j-th node of the second graph, N_2 is the number of nodes in



Fig. 1. Parts of Yahoo and Standard taxonomies

the second graph; and R specifies a similarity relation of the given nodes. A mapping is a set of mapping elements. We think of matching as the process of discovering mappings between two graph-like structures through the application of a matching algorithm.

Matching approaches can be classified into syntactic and semantic depending on how mapping elements are computed and on the kind of similarity relation R used (see [10] for in depth discussion):

- In syntactic matching the key intuition is to find the syntactic (very often string based) similarity between the labels of nodes. Similarity relation R in this case is typically represented as a [0, 1] coefficient, which is often considered as equivalence relation with certain level of plausibility or confidence (see [13, 7] for example). Similarity coefficients usually measure the closeness between two elements linguistically and structurally. For example, the similarity between $Computer_Storage_Devices_A$ and $Data_Storage_Devices_B$ based on linguistical and structural analysis could be 0,63.
- Semantic matching is an approach where semantic relations are computed between concepts (not between labels) at nodes. The possible semantic relations (R) are: equivalence (=); more general or generalization (\supseteq) ; less general or specification (\subseteq) ; mismatch (\bot) ; overlapping (\cap) . They are ordered according to decreasing binding strength, e.g., from the strongest (=) to the weakest (\cap) . For example, as from Figure 1 Computer_Hardware_A is more general than Large_Scale_Com-puters_B

In this paper we are focused on taxonomy matching. We think about taxonomy as a $\langle N, A, F_l \rangle$, where N is a set of nodes, A is a set of arcs, such that $\langle N, A \rangle$ is a rooted tree. F_l is a function from N to set of labels L (i.e., words in natural language). An example of taxonomy is presented on Figure 1. Notice that the distinguishing feature of taxonomies is the lack of formal encoding semantics.

3 The evaluation problem

Nearly all state of the art matching systems suffer from the lack of evaluation. Till very recently there was no comparative evaluation and it was quite difficult to find two systems evaluated on the same dataset. Often authors artificially synthesize datasets for empirical evaluation but rarely they explain their premises and assumptions. The last efforts [19] on matching evaluation concentrate rather on artificially produced and quite simple examples than real world matching tasks. Most of the current evaluation efforts were devoted to the schemas with tenth of nodes and only some recent works (see [6] for example) present the evaluation results for the graphs with hundreds of nodes. At the same time industrial size schemas contain up to tenth thousands of nodes.

The evaluation problem can be summarized as the problem of acquiring the reference relationship that holds between two nodes. Given such a reference relationship it would be straightforward to evaluate the result of a matching solution. Up to now the acquisition of the reference mappings that hold among the nodes of two taxonomies is performed manually. Similarly to the annotated corpora for information retrieval or information extraction, we need to annotate a corpus of pairwise relationships. Of course such an approach prevents the opportunity of having large corpora. The number of mappings between two taxonomies are quadratic with respect to taxonomy size, what makes hardly possible the manual mapping of real world size taxonomies. It is worthwhile to remember that web directories, for example, have many tens thousands of nodes. Certain heuristics can help in reducing the search space but the human effort is still too demanding.

Our proposal is to build a reference interpretation for a node looking at its use. We argue that the semantics of nodes can be derived by their pragmatics, i.e. how they are used. In our context, the nodes of a taxonomy are used to classify documents. The set of documents classified under a given node implicitly defines the its meaning. This approach has been followed by other researchers. For example in [5] the interpretation of a node is approximated by a model computed through statistical learning. Of course the accuracy of the interpretation is affected by the error of the learning model. We follow a similar approach but without the statistical approximation. The working hypothesis is that the meaning of two nodes is equivalent if the sets of documents classified under those nodes have a meaningful overlap.

The basic idea is to compute the relationship hypotheses based on the cooccurrence of documents. This document-driven interpretation can be used as a reference value for the evaluation of competing matching solutions. A simple definition of equivalence relationship based on documents can be derived by the F1 measure of information retrieval.

The Figure 2 shows a simple example. In the graphical representation we have two taxonomies, for each of them we focus our attention on a reference node. Let be S and P two sets of documents classified under the reference nodes of the first and second taxonomies respectively. We will refer to A_S and A_P as the set of documents classified under the ancestor nodes of S and P. Conversely we will refer to T_S and T_P as the set of documents classified under the subtrees of S

and P. The goal is to define a relationship hypothesis based on the overlapping of the set of documents, i.e. the pragmatic use of the nodes.

The first step, the *equivalence* relationship, can be easily formulated as the F1 measure of information retrieval [2]. The similarity of two sets of documents is defined as the ratio between the marginal sets and the shared documents:

$$Equivalence = \frac{|M_P^S| + |M_S^P|}{|O_P^S|}$$

where the set of shared documents is defined as $O_P^S = P \cap S$ and $M_P^S = S \setminus O_P^S$ is the marginal set of documents classified by S and not classified by P (similarly $M_S^P = P \setminus O_P^S$). The following equivalence applies $O_P^S = O_S^P$. Notice that "O" stands for "overlapping" and "M" stands for "Marginal set".

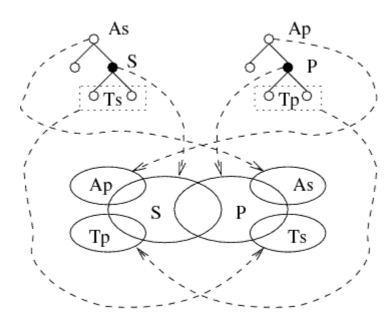


Fig. 2. The pairwise relationships between two taxonomies.

We do a step forward because we do not only compute the equivalence hypothesis based on the notion of F1 measure of information retrieval, but we extend such equation to define the formulation of generalization and specialization hypothesis. Generalization and specialization hypothesis can be formulated taking advantage of the contextual encoding of knowledge in terms of hierarchies of categories. The challenge is to formulate a generalization hypothesis (and conversely a specialization hypothesis) between two nodes looking at the overlapping of set of documents classified in the ancestor or subtree of the reference nodes [1].

The generalization relationship holds when the first node has to be considered more general of the second node. Intuitively, it happens when the documents classified under the first nodes occur in the ancestor of the second node, or the documents classified under the second node occur in the subtree of the first node. Following this intuition we can formalize the generalization hypothesis as

$$Generalization = \frac{|M_P^S| + |M_S^P|}{|O_P^S| + |O_{A_S}^P| + |O_{T_P}^S|}$$

where $O_{A_S}^P$ represents the set of documents resulting from the intersection between M_S^P and the set of documents classified under the concepts in the hierarchy above S (i.e. the ancestors); similarly $O_{T_P}^S$ represents the set of documents resulting from the intersection between M_P^S and the set of documents classified under the concepts in the hierarchy below P (i.e. the children).

In a similar way we can conceive the *specialization* relationship. The first node is more specific than second node when the meaning associated to the first node can be subsumed by the meaning of the second node. Intuitively, it happens when the documents classified under the first nodes occur in the subtree of the second node, or the documents classified under the second node occur in the ancestor of the first node.

$$Specialization = \frac{|M_P^S| + |M_S^P|}{|O_P^S| + |O_{T_S}^S| + |O_{A_P}^S|}$$

where $O_{T_S}^P$ represents the set of documents resulting from the intersection between M_S^P and the set of documents classified under the concepts in the hierarchy below S (i.e. the children); similarly $O_{A_P}^S$ represents the set of documents resulting from the intersection between M_P^S and the set of documents classified under the concepts in the hierarchy above P (i.e. the ancestors).

The three definitions above allow us to compute a relationship hypothesis between two nodes of two different taxonomies. Such an hypothesis relies on the assumption that if two nodes classify the same set of documents, the meaning associated to the nodes is reasonably the same. Of course this assumption is true for a virtually infinite set of documents. In a real world case study we face with finite set of documents, and therefore this way of proceed is prone to error. Nevertheless, our claim is that the approximation introduced by our assumption is balanced by the benefit of scaling with the annotation of large taxonomies.

4 Building a Large Scale Mapping Dataset

Let try to apply the notion of document-driven interpretation to a real world case study. We focus our attention to web directories for many reasons. Web directories are widely used and known; moreover they are homogeneous, that is they cover general topics. The meaning of a node in a web directory is not defined with formal semantics but by pragmatics. Furthermore the web directories

address the same space of documents, therefore the working hypothesis of cooccurence of documents can be sustainable. Of course different web directories don't cover the same portion of the web but the overlapping is meaningful.

The case study of web directories meets two requirements of the matching problem: to have heterogeneous representations of the same topics and to have taxonomies of large dimensions.

We address three main web directories: Google, Yahoo! and Looksmart. Nodes have been considered as categories denoted by the lexical labels, the tree structures have been considered as hierarchical relations, and the URL classified under a given node as documents. The following table summarizes the total amount of processed data.

Web Directories	\mathbf{Google}	${f Looksmart}$	Yahoo!
number of nodes	335.902	884.406	321.585
number of urls	2.425.215	8.498.157	872.410

Let briefly describe the process by which we have arranged an annotated corpus of pairwise relations between web directories.

- **Step 1.** We crawled all three web directories, both the hierarchical structure and the web contents, then we computed the subset of URLs classified by all them.
- **Step 2.** We pruned the downloaded web directories removing all the URLs that were not referred by all the three web directories.
- **Step 3.** We performed an additional pruning removing all the nodes with a number of URLs under a given threshold. In our case study we fixed such a threshold at 10.
- **Step 4.** We manually recognized potential overlapping between two branches of two different web directories like

Google:/Top/Science/Biology

Looksmart:/Top/Science-and-Health/Biology

Yahoo:/Top/Computers-and-Internet/Internet

Looksmart:/Top/Computing/Internet

Google:/Top/Reference/Education

Yahoo:/Top/Education

We recognized 50 potential overlapping and for each of them we run an exhaustive assessment on all the possible pairs between the two related subtrees. Such an heuristic allowed us to reduce the quadratic explosion of cartesian product of two web directories. We focussed the analysis on smaller subtrees where the overlaps were more likely.

Step 5. We computed the three document-driven hypothesis for *equivalence*, generalization and specialization relationship as described above. The computation of equivalence, generalization and specialization hypothesis is estimated by a number in the range [0,1]. Since the cumulative hypothesis of

all three relationships for the same pair of nodes can not be higher than 1, we introduce a threshold to select the winning hypothesis. We fixed such a threshold to 0.5.

We discarded all the pairs where none of the three relationship hypotheses was detected. This process allowed us to obtain 2265 pairwise relationships defined using the document-driven interpretation. Half are equivalence relationships and half are generalization relationships (let notice that by definition generalization and specialization hypothesis are symmetric).

In the following we will refer to this dataset as TaxME, TAXonomy Mapping Evaluation.

5 The Empirical Evaluation

The evaluation was designed in order to assess the major dataset properties namely:

- Complexity, namely the fact that the dataset is "hard" for state of the art matching systems.
- Discrimination ability, namely the fact that the dataset can discriminate among various matching approaches.
- Incrementality, namely the fact that the dataset allows to incrementally discover the weaknesses of the tested systems.
- Correctness, namely the fact that the dataset can be a source of correct results.

We have evaluated two state of the art matching systems $COMA^3$ and S-Match and compared their results with baseline solution. Let us describe the matching systems in more detail.

The COMA system [13] is a generic syntactic schema matching tool. It exploits both element and structure level techniques and combines the results of their independent execution using several aggregation strategies. COMA provides an extensible library of matching algorithms and a framework for combining obtained results. Matching library contains 6 individual matchers, 5 hybrid matches and 1 reuse-oriented matcher. One of the distinct features of the COMA tool is the possibility of performing iterations in the matching process. In the evaluation we used default combination of matchers and aggregation strategy (NamePath+Leaves and Average respectively).

S-Match is a generic semantic matching tool. It takes two tree-like structures and produces a set of mappings between their nodes. S-Match implements semantic matching algorithm in 4 steps. On the first step the labels of nodes are linguistically preprocessed and their meanings are obtained from the Oracle (in the current version WordNet 2.0 is used as an Oracle). On the second step the

³ In the evaluation we use the version of COMA described in [13]. A newer version of the system COMA++ exists but we do not have it. However as from the evaluation results presented in [10, 11], COMA is still best among the other syntactic matchers.

meaning of the nodes is refined with respect to the tree structure. On the third step the semantic relations between the labels at nodes and their meanings are computed by the library of element level semantic matchers. On the fourth step the matching results are produced by reduction of the node matching problem into propositional validity problem, which is efficiently solved by SAT solver or ad hoc algorithm (see [10, 11] for more details).

We have compared the performance of these two systems with baseline solution. The pseudo code of baseline node matching algorithm is given in Algorithm 1. It is executed for each pair of nodes in two trees. The algorithm considers a simple string comparison among the labels placed on the path spanning from a node to the root of the tree. Equivalence, more general and less general relations are computed as the corresponding logical operations on the sets of the labels.

Algorithm 1 Baseline node matching algorithm

```
1: String nodeMatch(Node sourceNode, Node targetNode)
2: Set sourceSetOfLabels=getLabelsInPathToRoot(sourceNode)
3: Set targetSetOfLabels=getLabelsInPathToRoot(targetNode)
4: if sourceSetOfLabels \equiv targetSetOfLabels then
5:
      result = " \equiv "
6: else if sourceSetOfLabels \subseteq targetSetOfLabels then
7:
      result = "\subseteq"
8: else if sourceSetOfLabels \supseteq targetSetOfLabels then
9:
      result = "\supseteq"
10: else
      result="Idk"
11:
12: end if
13: return result
```

The systems have been evaluated on the dataset described in Section 4. We computed the number of matching tasks solved by each matching system. Notice that the matching task was considered to be solved in the case when the matching system produce specification, generalization or equivalence semantic relation for it. For example, TaxME suggests that specification relation holds in the following example:

Google:/Top/Sports/Basketball/Professional/NBDL Looksmart:/Top/Sports/Basketball

COMA produced for this matching task 0.58 similarity coefficient, which can be considered as equivalence relation with probability 0.58. In the evaluation we consider this case as true positive for COMA (i.e., the mapping was considered as found by the system).

Notice that at present TaxME contains only true positive mappings. This fact allows to obtain the correct results for Recall measure, which is defined as a ratio of reference mappings found by the system to the number of reference mappings. At the same time Precision, which is defined as ratio of reference

mappings found by the system to the number of mappings in the result, can not be correctly estimated by the dataset since, as from Section 4, TaxME guarantee only the correctness but not completeness of the mappings it contain.

6 Discussion of Results

Evaluation results are presented on Table 1. It contains the total number of mappings found by the systems and the partitioning of the mappings on semantic relations. Let us discuss the results through major dataset properties perspective.

Google vs. Looksmart Google vs. Yahoo Looksmart vs. Yahoo Total COMA 608 250 18 876 (38,68%) 608 250 18 876 \subseteq not applicable S-Match 584 83 669 (29,54%) 2 5 0 7 \subseteq 462 19 67 536 59 0 595 $130 \ \overline{(5,39\%)}$ Baseline 54 76 0 520 0 52 \subseteq 0 76 0 762 0 0 2

Table 1. Evaluation Results

6.1 Complexity

As from Table 1 the results of baseline are surprisingly low. It produced slightly more than 5% of mappings. This result is interesting since on the previously evaluated datasets (see [4] for example) the similar baseline algorithm performed quite well and found up to 70% of mappings. This lead us to conclusion that the dataset is not trivial (i.e., it is essentially hard for simple matching techniques).

As from Figure 3 S-Match found about 30% of the mappings in the biggest (Google-Yahoo) matching task. At the same time it produced slightly less than 30% of mappings in all the tasks. COMA found about 35% of mappings on Google-Looksmart and Yahoo-Looksmart matching tasks. At the same time it produced the best result on Google-Yahoo. COMA found slightly less than 40% of all the mappings. These results are interesting since as from [13, 10] previously reported recall values for both systems were in 70-80% range. This fact turn us to conclusion that the dataset is hard for state of the art syntactic and semantic matching systems.

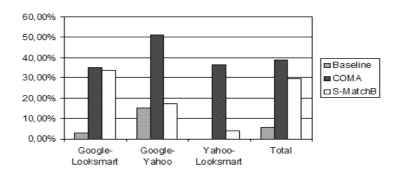


Fig. 3. Percentage of correctly determined mappings(Recall)

6.2 Discrimination ability

Consider Figure 4. It presents the partitioning of the mappings found by S-Match and COMA. As from the figure the sets of mappings produced by COMA and S-Match intersects only on 15% of the mappings. This fact turns us to important conclusion: the dataset is discriminating (i.e., it contains a number of features which are essentially hard for various classes of matching systems and allow to discriminate between the major qualities of the systems).

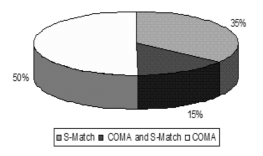


Fig. 4. Partitioning of the mappings found by COMA and S-Match

6.3 Incrementality

In order to evaluate incrementality we have chosen S-Match as a test system. In order to identify the shortcomings of S-Match we manually analyzed the mappings missed by S-Match. This analysis allowed us to clasterize the mismatches into several categories. In this paper we describe in detail one of the most important categories of mismatches namely $Meaningless\ labels$.

Consider the following example:

Looksmart:/Top/Science_&_Health/Social_Science/Archaeology/ By_Region/Andes_South_America/Nazca

In this matching task some labels are meaningful in the sense they define the context of the concept. In our example these are Social_Sciences, Archaeology, South_America, Nazca. The other labels do not have a great influence on the meaning of concept. At the same time they can prevent S-Match from producing the correct semantic relation. In our example S-Match can not find any semantic relation connecting Nazca_Lines and Nazca and the reason for this is By_Region label, which is meaningless in the sense it is defined only for readability and taxonomy partitioning purposes. An other example of this kind is

Here A and $Actors_A/Actors_Aa-Af$ does not influence on the meaning of the concept. At the same time they prevent S-Match to produce the correct semantic relation holding between the concepts.

An optimized version of S-Match (S-Match++) has a list of meaningless labels. At the moment the list contains only about 30 words but it is automatically enriched in preprocessing phase. A general rule for considering natural language label as meaningless is to check whither it is used for taxonomy partitioning purposes. For example, S-Match++ consider as meaningless the labels with the following structure by $\langle word \rangle$, where $\langle word \rangle$ stands for any word in natural language. However this method is not effective in the case of labels composed from alphabet letters (such as $Actors_Aa-Af$ from previous example). S-Match++deal with the latter case in the following way: the combination of letters are considered as meaningless if it is not recognized by WordNet, not in abbreviation or proper name list, and at the same time its length are less or equal to 3. The addition of these techniques allowed to improve significantly the S-Match matching capability. The number of mappings found by the system on TaxME dataset increased by 15%. This result gives us an evidence to incrementality of the dataset (i.e., the dataset allows to discover the weaknesses of the systems and gives the clues to the systems evolution).

6.4 Correctness

Analysis of S-Match results on TaxME allowed to identify 10 major bottlenecks in the system implementation. At the moment we are developing ad hoc techniques allowing to improve S-Match results in this cases. The current version of S-Match (S-Match++) contains the techniques allowing to solve 5 out of 10 major categories of mismatches. Consider Figure 5. It contains the results of comparative evaluation S-Match++ against the other systems. As from the figure S-Match++ significantly outperforms all the other systems. It found about

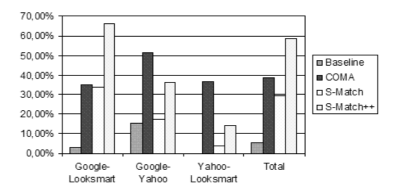


Fig. 5. Percentage of correctly determined mappings(Recall)

60% of mappings in all the matching tasks, what is twice better than *S-Match* result. This significant improvement would be impossible without comprehensive evaluation on TaxME. At the same time as from a preliminary manual analysis only 2-3% of mappings in TaxME is not correct. Taking into account the notion of idiosyncratic classification [12], such a mismatch can be considered as marginal.

7 Conclusions

In this paper we have presented a mapping dataset which carries the key important properties of Complexity, Incrementality, Discrementality and Correctness. We have evaluated the dataset on two state of the art matching systems representing different matching approaches. As from the evaluation, the dataset can be considered as a powerful tool to support the evaluation and research on the matching solutions.

The ultimate step which needs to be performed is to acquire the user mappings for TaxME dataset. We have already arranged such a kind of test and the results through preliminary are promising. Unfortunately at the moment more significant statistics needs to be collected in order to further improve TaxME .

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