

Discovery of Positive and Negative Rules in Knowledge-Bases

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ABSTRACT

We present RuDiK, a system for the discovery of declarative rules over knowledge-bases (KBs). RuDiK output is not limited to rules that rely on “positive” relationships between entities, such as “if two persons have the same parent, they are siblings”, as in traditional constraint mining for KBs. On the contrary, it discovers also negative rules, i.e., patterns that lead to contradictions in the data, such as “if two persons are married, one cannot be the child of the other”. While the former class is fundamental to infer new relationships in the KB, the latter class is crucial for other tasks, such as error detection in data cleaning, or the creation of negative examples to bootstrap learning algorithms. The algorithm to discover positive and negative rules is designed with three main requirements: (i) enlarge the *expressive power* of the rule language to obtain complex rules and wide coverage of the facts in the KB, (ii) allow the discovery of *approximate* rules to be robust to errors and incompleteness in the KB, (iii) use disk-based algorithms, effectively enabling the mining of large KBs in commodity machines. We have conducted extensive experiments using real-world KBs to show that RuDiK outperform previous proposals in terms of efficiency and that it discovers more effective rules for the application at hand.

1. INTRODUCTION

Building large RDF knowledge-bases (KBs) is a popular trend in information extraction. KBs store information in the form of triples, where a *predicate*, or relation, expresses a binary relation between a *subject* and a *object*. KB triples, often referred as facts, store information about real-world entities and their relationships, such as “Michelle Obama is married to Barack Obama”, or “Larry Page is the founder of Google”. Significant effort has been put in the research community on KBs creation in the last 10 years (DBPedia [8], FreeBase [9], Wikidata [37], DeepDive [33],

Yago [35], NELL [10], TextRunner [6]) as well as in the industry (e.g., Facebook¹, Google [17], Wal-Mart [16]).

Unfortunately, due to their creation process, KBs are usually erroneous and incomplete. KBs are bootstrapped by extracting information from sources, oftentimes from the Web, with minimal or no human intervention. This leads to two main problems. First, errors are propagated from the sources, or introduced by the extractors, leading to false facts in the KB. Second, usually KBs do not limit the information of interest with a schema that clearly defines instance data. The set of predicates is unknown a-priori, and adding facts defined on new predicates is just a matter of inserting new triples in the KB without any integrity check. Since *closed world assumption* (CWA) does not hold [17, 23], we cannot assume that a missing fact is false, but rather we should label it as *unknown* (*open world assumption*).

As a direct consequence, the amount of errors and incompleteness in KBs is significantly larger than classic databases [36]. Since KBs are large, e.g., WIKIDATA has more than 1B facts and 300M different entities, checking all triples to find errors or add new facts cannot be done manually and tools are needed to assist humans in the KBs curation. A natural approach to assist curators of KBs is to discover *declarative rules* that can be executed over the KBs to improve the quality of the data [2, 11, 23]. However, these approaches so far have focused on the discovery of rules to derive new facts only, while for the first time we target the discovery of two different types of rules: (i) *positive rules*, used to enrich the KB with new facts and thus increase its coverage of the reality; (ii) *negative rules*, used to spot logical inconsistencies and identify erroneous triples.

Example 1: Consider a KB with information about parent and child relationships. A positive rule can be:

$$\text{parent}(b, a) \Rightarrow \text{child}(a, b)$$

which states that if a person a is parent of person b , then b is child of a . If the KB has the fact “Clint Eastwood is the parent of Scott Eastwood”, we can infer the fact that “Scott Eastwood is the child of Clint Eastwood”. A negative rule has similar form, but very different semantics:

$$\text{birthDate}(a, v_0) \wedge \text{birthDate}(b, v_1) \wedge v_0 > v_1 \Rightarrow \neg \text{child}(a, b)$$

The above rule states that a person b cannot be child of a if a was born after b . By instantiating the above rule for the *child* facts in the KB, we may discover erroneous triples stating that a child is born before a parent.

¹<https://developers.facebook.com/docs/graph-api>

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While intuitive to humans, the rules above must be manually stated in a KB in order to be enforced, and there are thousands of rules in a large KB with hundreds of predicates [24]. Other than enriching and cleaning KBs, negative rules support other use cases. We will show how negative rules can improve Machine Learning tasks by providing meaningful training examples [31, 33].

However, three main challenges arise when discovering rules for KBs.

Errors. Traditional database techniques for rules discovery make the assumption that data is either clean or has a negligible amount of errors [4, 12, 26, 38]. We will show that KBs present a high percentage of errors and we need techniques that are noise tolerant.

Open Word Assumption. KBs contain only positive statements, and, without CWA, we cannot derive negative statements as counter examples – classic databases approaches rely on the presence of positive and negative examples [15, 30] or a fixed schema given as input.

Volume. Existing approaches for discovery on positive rules in KBs assume that data can fit into memory, something that it is not realistic given the large, and increasing, size of the KB [2, 23]. More recent approaches also rely on main memory algorithm and try to solve this problem with distributed architectures [11, 22].

We advocate that a rule discovery system should be designed to discover *approximate rules* since errors and incompleteness are in the nature of the KBs. Also, the main memory constraint not only reduces the applicability of the system, but also forces limitations on the *expressive power* of the language in order to reduce the search space. In fact, the larger is the number of patterns that can be expressed in the rules, the larger is the number of new facts and errors that can be identified, and with increasing precision. However, this comes with a computational cost, as the search space quickly becomes much larger. For this reason, existing approaches for mining positive rules prune aggressively the search space, and rely on a simple language [2, 11, 23].

We present RuDiK (Rules Discovery in Knowledge Bases), a novel system for the discovery of positive and negative rules over KBs that addresses the challenges above. RuDiK is the first system capable of discovering both approximate positive and negative rules without assuming that the KB fits into memory by exploiting the following contributions.

Problem Definition. We formally define the problem of rules discovery over erroneous and incomplete KBs. The input of the problem consists of a target predicate, from which we identify sets of positive and negative examples. In contrast with traditional ranking of rules based on a measure of support [15, 23, 29, 32], our problem definition aims at the identification of a subset of approximate rules. Given errors in the data and incompleteness, the ideal solution is a compact set of rules that cover the majority of the positive examples, and as few negative examples as possible. We map the problem to the well-known weighted set cover problem.

Search Algorithm. We give a $\log(k)$ approximation of the problem close in spirit to the greedy algorithm for set cover, where k is the maximum number of positive examples covered by a single rule. We discover the rules to feed the algorithm by judiciously using the memory. The algorithm incrementally materializes the KB as a directed graph, and discovers rules by navigating valid paths. To reduce the use

of resources, at each iteration we follow the most promising path in a A^* traversal fashion, allowing the pruning of unpromising paths. By materializing only the portion of the KB that is needed for the discovery, and generating nodes and edges *on demand*, the disk is accessed only whenever the navigation of a specific path is required. The significant reduction of the search space reduces the running time an order of magnitude in the best case scenario.

Sampling for Exploration. To further improve performance, we also introduce sampling techniques that are aware of the data challenges in KBs. Our sampling leads to the same improvements in execution times than traditional samplings approach, but, as it is design to mitigate the problems in KBs, it can lead to better rules, i.e., a larger number of correct rules in the output and a smaller number of false rules.

We experimentally verify the performance of rules discovery in RuDiK using real-world datasets (Section 5). The evaluation is carried on the 3 most popular and widely used KBs, namely DBPEDIA [8], YAGO [34], and WIKIDATA [37]. We show that our problem formulation leads to approximate rules that are better in quality than previous systems and that the search algorithm is general enough to deliver accurate rules both in the positive and negative settings, clearly outperforming state-of-the-art systems in running time. We also show the impact of the proposed optimization and give an example of application where discovered rules have a beneficial impact when providing representative examples for Machine Learning algorithms.

2. PRELIMINARIES AND DEFINITIONS

The work of this chapter focuses on discovering rules from RDF² KBs. An RDF KB is a database that represents common information through RDF triples $\langle s, p, o \rangle$, where a *subject* is connected to a *object* via a *predicate*. Triples are often called *facts*. As an example, the fact that Scott Eastwood is the child of Clint Eastwood could be represented with the triple $\langle \text{Clint_Eastwood}, \text{child}, \text{Scott_Eastwood} \rangle$. RDF KB triples respect the following constraints:

- triple subjects are always *entities*, i.e., concepts from the real world;
- triple objects can be either entities or *literals*. Literals represent primitive types – numbers, dates, and strings;
- triple predicates specifies real-world relationships between subjects and objects.

If we compare the KB model with object oriented programming languages, entities correspond to complex objects with properties and attributes, while literals correspond to primitive types such as integers, chars and strings.

Differently from, e.g., relational databases, KBs do not have a schema that instance data is forced to follow. The set of predicates is unknown a-priori, and adding new predicates is just a matter of inserting new triples having those predicates. Sometimes KBs include *T-Box* facts in order to define classes, domain/co-domain types for predicates, and relationships among classes. While the T-Box can be useful in order to check integrity and consistency within a KB, very often such information is incomplete or not present at

²<https://www.w3.org/RDF/>

all. Hence our focus in this thesis is only on the *A-Box*, the set of facts that describe instance data. If on the one hand this model allows great flexibility, on the other hand the likelihood of introducing errors and inconsistencies is much higher than traditional schema-guided databases.

2.1 Language

Inspired by Inductive Logic Programming [30], the goal of our work is to discover first-order logical formulas in KBs. More specifically, we target the discovery of *Horn Rules*. A Horn Rule is a disjunction of *atoms* with at most one unnegated atom. When written in the implication form, Horn Rules have one of the following format:

$$A_1 \wedge A_2 \wedge \dots \wedge A_n \Rightarrow B \quad A_1 \wedge A_2 \wedge \dots \wedge A_n \Rightarrow \neg B$$

where $A_1 \wedge A_2 \wedge \dots \wedge A_n$ consists of the *body* of the rule (a conjunction of atoms), while B is the *head* of the rule (a single atom). The head of the rule is either unnegated (left) or negated (right). We call the former *definite clause* or more simply *positive rule*, while the latter *goal clause* or *negative rule*. In the context of KBs, an atom is a predicate connecting two variables, two entities, or an entity and a variable. For simplicity, we represent an atom with the notation $\mathbf{rel}(a, b)$, where \mathbf{rel} is the name of the predicate, and a and b are either variables or entities. In the context of Horn Rules, all variables appearing in a rule are implicitly universally quantified. Given a Horn Rule r , we define r_{body} as the body of the rule, and r_{head} as the head of the rule. We define the variables appearing in the head of the rule as the *target variables*. The first rule of Example 1 shows a plausible positive rule, while the second rule is a potential negative rule. a and b are the target variables in both cases.

Furthermore we extend the language in order to include *literals comparison*. A literal comparison is a special atom $\mathbf{rel}(a, b)$, where $\mathbf{rel} \in \{<, \leq, \neq, >, \geq\}$, and a and b can only be assigned to literal values except if \mathbf{rel} is equal to \neq . In such a case a and b can be also assigned to entities. We will explain later on in the thesis why this exception is important.

Given a KB kb and an atom $A = \mathbf{rel}(a, b)$ where a and b are two entities, we say that A *holds over* kb iff $\langle a, \mathbf{rel}, b \rangle \in kb$. Given a KB kb and an atom $A = \mathbf{rel}(a, b)$ with at least one variable, we say that A can be *instantiated over* kb if there exists at least one entity from kb for each variable in A such that if we substitute variables with entities in A , A holds over kb . Transitivity, given a body of a rule r_{body} and a KB kb , we say that r_{body} can be instantiated over kb if every atom in r_{body} can be instantiated.

Eventually, we introduce two language biases in order to avoid the explosion of the search space. Following the biases introduced by other approaches for rules discovery in KBs [23] (TODO: cite Sigmod Paper Ontological Path...), we introduce such restrictions in order to speed up the discovery process. Language biases introduce a compromise between the expressiveness of target rules and the complexity of the discovery problem. Therefore we define a rule *valid* iff it follows the following constraints.

Connectivity. We say that in a rule an atom A_1 can be reached by an atom A_2 iff A_1 and A_2 share at least one variable or one entity. The connectivity constraint requires that every atom in a rule must be *transitively* reached by any other atom in the rule.

Repetition. Every variable in a rule must appear at least twice. Since target variables already appear once in the head of the rule, the repetition constraint limited to the body of a rule requires that each variable that is not a target variable must appear at least twice, while target variables at least once.

Language restrictions obviously limit the output of the discovery algorithm to a subset of all plausible rules. We will show in Section 3 how these restrictions can significantly speed up the discovery process through a fast disk-based approach that leverages on these limitations.

2.2 Rules Coverage

Given a pair of entities (x, y) from a KB kb and a Horn Rule r , we say that r_{body} *covers* (x, y) if $(x, y) \models r_{body}$. In other words, given a Horn Rule $r = r_{body} \Rightarrow \mathbf{r}(a, b)$, r_{body} covers a pair of entities $(x, y) \in kb$ iff we can substitute a with x , b with y , and the rest of the body can be instantiated over kb . Given a set of pair of entities $E = \{(x_1, y_1), (x_2, y_2), \dots, (x_n, y_n)\}$ and a rule r , we denote by $C_r(E)$ the *coverage* of r_{body} over E as the set of elements in E covered by r_{body} : $C_r(E) = \{(x, y) \in E \mid (x, y) \models r_{body}\}$.

Given the body r_{body} of a Horn Rule r , we denote by r_{body}^* the *unbounded body* of r . The unbounded body of a rule is obtained by keeping only atoms that contain a target variable and substituting such atoms with new atoms where the target variable is connected to a new unique variable. As an example, given $r_{body} = \mathbf{rel}_1(a, v_0) \wedge \mathbf{rel}_2(v_0, b)$ where a and b are the target variables, $r_{body}^* = \mathbf{rel}_1(a, v_1) \wedge \mathbf{rel}_2(v_2, b)$. While in r_{body} the target variables are bounded to be connected to the same variable v_0 , in r_{body}^* the target variables are not bounded to share the same variable. Given a set of pair of entities $E = \{(x_1, y_1), (x_2, y_2), \dots, (x_n, y_n)\}$ and a rule r , we denote by $U_r(E)$ the *unbounded coverage* of r_{body}^* over E as the set of elements in E covered by r_{body}^* : $U_r(E) = \{(x, y) \in E \mid (x, y) \models r_{body}^*\}$. Note that, given a set E , $C_r(E) \subseteq U_r(E)$. In other words, the unbounded coverage of a rule over E always contains all the elements of the coverage over E .

Example 2: Given the negative rule r of Example 1 and a KB kb , we denote with E the set of all possible pairs of entities in kb . The coverage of r over E ($C_r(E)$) is the set of all pairs of entities $(x, y) \in kb$ such that both x and y have the **birthDate** information and x is born after y , while the unbounded coverage of r over E ($U_r(E)$) is the set of all pairs of entities (x, y) such that both x and y have the **birthDate** information, no matter what the relation between the two birth dates is.

The unbounded coverage is essential to distinguish between missing and inconsistent information: if for a pair of entities (x, y) the **birthDate** information is missing for either x or y (or both), we cannot say whether x was born before or after y , therefore we cannot state whether the negative rule of Example 1 covers (x, y) or not. On the other hand, if both x and y have the **birthDate** information and x is born before y , we can affirm that the negative rule of Example 1 does not cover (x, y) . Given that modern KBs are largely incomplete [17,28], discriminating between missing and conflicting information becomes of paramount importance.

We can now define the coverage and the unbounded coverage for a set of rules $R = \{r_1, r_2, \dots, r_n\}$ as the union of individual coverages:

$$C_R(E) = \bigcup_{r \in R} C_r(E) \quad U_R(E) = \bigcup_{r \in R} U_r(E)$$

Our problem tackles the discovery of positive (negative) rules for an input given predicate (target predicate). We uniquely identify a predicate with two different sets of pairs of entities. *G* – *generation set*: *G* contains good examples for the target predicate (*G* contains examples of parents and children if we are discovering positive rules for a child predicate). *V* – *validation set*: *V* contains counter examples for the target predicate (pairs of people that are not in a child relation). We will explain in Section 3.1 how to generate these two sets for a given predicate. Note that our approach is not less generic than those for mining rules for an entire KB (e.g., [2, 23]): it is true that we require a target predicate as input, however we can generically apply our setting for every predicate in the KB and compute rules for each of them (see Section 5.2).

We can now formalise the *exact discovery problem*.

Definition 1: Given a KB *kb*, two sets of pairs of entities *G* and *V* from *kb* such that $G \cap V = \emptyset$, and a universe of Horn Rules *R*, a solution for the *exact discovery problem* is a subset *R'* of *R* such that:

$$R_{opt} = \underset{|R'|}{\operatorname{argmin}} (C_{R'}(G) = G) \wedge (C_{R'}(V) \cap V = \emptyset)$$

□ The ideal solution

is a set of rules that covers all examples in *G*, and none of the examples in *V*. Note that given a pair of entities (x, y) , we can always generate a Horn Rule whose body covers only (x, y) by assigning target variables to *x* and *y*.

Unfortunately, since the solution is not allowed to cover any element in *V*, in the worst case the exact solution may be a set of rules such that each rule covers only one example in *G*, making such a set of rules difficult to use and understand.

2.3 Weight Function

In order to allow flexibility and errors in both *G* and *V*, we drop the strict requirement of not covering any element of *V*. However, since covering elements in *V* is an indication of potential errors, we want to limit the coverage over *V* to the minimum possible. We therefore define a *weight* to be associated with a rule.

Definition 2: Given a KB *kb*, two sets of pair of entities *G* and *V* from *kb* such that $G \cap V = \emptyset$, and a Horn Rule *r*, the weight of *r* is defined as follow:

$$w(r) = \alpha \cdot \left(1 - \frac{|C_r(G)|}{|G|}\right) + \beta \cdot \left(\frac{|C_r(V)|}{|U_r(V)|}\right) \quad (1)$$

with $\alpha, \beta \in [0, 1]$ and $\alpha + \beta = 1$. □ The weight is a value between 0 and 1 that captures the *goodness* of a rule w.r.t. *G* and *V*: the better the rule, the lower the weight – perfect rules would have a weight of 0. The weight is made of two components normalised by the two parameters α and β . 1) The first component captures the coverage over the generation set *G* – the ratio between the coverage of *r* over *G* and *G* itself. Note that if *r* covers all elements in *G*, then this component is 0 because of the subtraction from 1. 2) The second component aims at quantifying potential errors of *r*, or rather the coverage over *V*. The coverage over *V* is not divided by total elements in *V*, because for those elements

in *V* that do not have predicates stated in *r*_{body} we cannot be sure whether such elements are not covered by *r*. Thus we divide the coverage over *V* by the unbounded coverage of *r* over *V*. Ideally this number is close to 0. The parameters α and β are used to give relevance to each component. We would set a high β if we want to discover rules with high precision that identify few mistakes, or we would set a high α if we are more interested in recall and the discovered rules should identify as many examples as possible.

Example 3: W.r.t. the negative rule *r* of Example 1, given two sets of pair of entities *G* and *V* from a KB *kb*, the two components of *w_r* are computed as follow: 1) the first component is computed as 1 minus number of pairs (x, y) in *G* where *x* is born after *y* divided by the total number of elements in *G*; 2) the second component is the ratio between number of pairs (x, y) in *V* where *x* is born after *y* and number of pairs (x, y) in *V* where the date of birth (for both *x* and *y*) is available in *kb*.

Definition 3: Given a set of rules *R*, the weight for *R* is defined as:

$$w(R) = \alpha \cdot \left(1 - \frac{|C_R(G)|}{|G|}\right) + \beta \cdot \left(\frac{|C_R(V)|}{|U_R(V)|}\right)$$

□

Assigning a weight to one or multiple rules allows us to take into consideration an important aspect of modern KBs: the presence of errors. We will show in the experimental evaluation that very often universally correct rules have a significant coverage over *V*, which corresponds to errors in the KB. The exact discovery problem implies the absence of errors in the input KB, however such an assumption is too strong for modern KBs that are usually automatically built from external web sources [17, 33, 34] and thus inconsistencies are common [36].

2.4 Problem Definition

We can now state the approximate version of the problem.

Definition 4: Given a KB *kb*, two sets of pair of entities *G* and *V* from *kb* where $G \cap V = \emptyset$, a universe of rules *R*, and a *w* weight function for *R*, a solution for the *approximate discovery problem* is a subset *R'* of *R* such that:

$$R_{opt} = \underset{w(R')}{\operatorname{argmin}} (R'(G) = G)$$

□

We can map this problem to the well-known weighted set cover problem, which is proven to be a **NP-complete** problem [14]. The universe corresponds to *G* and the input sets are all the possible rules defined in *R*.

Since we want to minimise the total weight of the output rules, the approximate version of the discovery problem aims to cover all elements in *G*, and as few as possible elements in *V*. Since for each element $g \in G$ there always exists a rule that covers exactly only *g* (single-instance rule), an optimal output is always guaranteed to exist. We expect the output to be made of some rules that cover multiple examples in *G*, while remaining examples in *G* to be covered by single-instance rules. In the best-case scenario a single rule covers all elements in *G* and none of the elements in *V*.

Section 4 will describe a greedy polynomial algorithm to find a good solution for the approximate discovery problem.

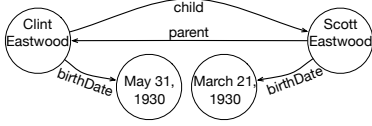


Figure 1: Graph Portion of DBpedia

3. RULES GENERATION

We introduce RuDiK, a disk-based approach to discover first-order Horn Rules in RDF KBs. The first task RuDiK needs to address is the generation of the universe of all possible rules R . Each rule in R must cover one or more examples of the generation set G .

In this setting, the universe of all possible rules can be generated by just inspecting elements of G . In fact, a rule that does not cover any element of G will never be part of the optimal solution, since it will not give any contribution to the set cover problem. This is a key part of our generation approach: we do not inspect the entire KB, but just entities belonging to G . The smaller the size of G is, the smaller is the search space for rules generation. On the contrary, recent KB rule mining approaches tackle the problem by searching for valid rules over the entirety of the KB. This is typically done by initially looking at all facts that share a common predicate, and then expanding these facts with other predicates that share common entities [23] (TODO : CITE Ontological path finding). Alternatively other approaches leverage on well established relational database techniques such as functional dependencies discovery (TODO: cite SIGMOD DEMO Data Lakes).

The generation problem is simple but presents significant scalability issues. The number of all possible rules increases exponentially with the size of the KB and enumerating all of them means exploring a huge search space. This is the main reason why previous approaches solve the problem by loading the entire KB into main memory and by aggressively indexing triples on subject, object, and predicate. Unfortunately, modern KBs can easily exceed the size of hundreds of GBs, making a memory-based solution unfeasible on common machines. We introduce in RuDiK a novel rules generation technique that inspects only a portion of the KB. On the one hand, we load into main memory only a small fraction of the database. On the other hand, this model allows the extension of the language to be more expressive than previous works. One could argue that reducing the search space may lead to missing some meaningful rules. We will show in Section 5.2 that our generation technique for G creates a representative (reduced) view of the entire search space.

The data structure RuDiK leverages on is a straightforward translation of a KB kb into a directed graph: entities and literals are nodes of the graph, while there is a direct edge from node a to node b for each triple $\langle a, rel, b \rangle \in kb$. Edges are labelled, where the label is the relation rel that connects subject to object in the triple. Figure 1 shows a portion of DBPEDIA [8] that connects two person in a child and parent relationship, along with their dates of birth. The graph represents information of four KB triples (two birth dates, one parent and one child relation).

The body of a Horn Rule can be seen as a path in the graph. W.r.t. Figure 1, the body $child(a, b) \wedge parent(b, a)$ corresponds to the path *Clint Eastwood* \rightarrow *Scott Eastwood*

\rightarrow *Clint Eastwood*. In Section 2.1 we defined valid rules. A valid body of a rule contains target variables a and b at least once, and every other variable at least twice. In a valid body also each atom is transitively connected to every other atom. If we allow navigation of edges in any direction (no matter what the direction of the edge is on the graph), we can easily translate bodies of valid rules to valid paths on the graph. Given a pair of entities (x, y) , a valid body corresponds to a valid path p on the graph that meets the following criteria:

1. p starts at the node x ;
2. p touches y at least once;
3. p ends in x , in y , or in a different node that has been touched before.

In other words, given the body of a rule r_{body} , r_{body} covers a pair of entities (x, y) iff there exists a valid path on the graph that corresponds to r_{body} . Therefore, given a pair of entities (x, y) , we can generate bodies of all possible valid rules by simply computing all valid paths from x with a standard BFS. Note how the transitively connection between atoms is always guaranteed by the construction property of a path p : for each node n in p there always exists a subpath that connects n to every other node in p . The key point is the ability of navigating each edge in any direction, which basically means turning the original directed graph into an undirected one.

Despite the navigation over an undirected graph, we still need to keep track of the original direction of the edges. This is essential when translating paths to Horn Rule bodies. In fact, if we navigate an edge rel between a and b , an original edge direction from a to b produces the atom $rel(a, b)$, while a direction from b to a produces the atom $rel(b, a)$. These two atoms are different, where the position of variables is determined by the original direction of the edge.

One should notice that for two entities x and y , there might exist infinite valid paths starting from x since every node can be traversed multiple times. Thus we introduce the $maxPathLen$ parameter that determines the maximum number of edges in the path. When translating paths to Horn Rules, $maxPathLen$ determines the maximum number of atoms that we can have in the body of the rule. This parameter is essential to avoid the discovery of rules with infinite body length.

The main advantage of inspecting just the generation set G is the capability of loading only a small portion of the graph that is currently needed. Given a pair of entities (x, y) , we retrieve from the KB all those nodes at distance $maxPathLen - 1$ or less from x or y , along with their edges. Retrieving such nodes and edges can be done recursively: we maintain a queue of entities, and for each entity in the queue we fire a SPARQL query against the KB to get all entities (and edges) at distance 1 from the current entity – we call these queries *single hop queries*. We then add the new found entities to the queue iff they are at distance less than $maxPathLen - 1$ from either x or y and they have not been visited before. The queue is initialised with x and y . By doing so we retrieve a small portion of the entire KB, the only one needed to discover rules that cover (x, y) . We will show in the experimental section that SPARQL engines are very fast at executing single hop queries.

The generation of the universe of all possible rules for G is then straightforward: for each element $(x, y) \in G$, we

construct the portion of the graph as described above and compute all valid paths starting from x . Computing paths for every example in G implies also computing the coverage over G for each rule. The coverage of a rule r is simply the number of elements in G where there exists a path that corresponds to r_{body} . Our discovery technique will also generate single-instance rules: rules that cover only one example $(x, y) \in G$ by instantiating target variables a and b in the rule with x and y . Once the universe of all possible rules has been generated (along with coverages over G), computing coverage and unbounded coverage over V is just a matter of executing two SPARQL queries against the KB for each rule in the universe. We will show in Section 4 how some queries can be avoided, as well as there is no need of enumerating all possible rules in the universe as some of them will never be part of the final solution.

Clearly, the size of G has a direct impact on the search space and hence on the running time. Since we generate all valid rules for each example in G , the search space grows roughly linearly with the size of G . If we could know a-priori the minimum subset of examples that lead to the generation of all valid rules, then we could use only those few examples. A future direction we are working on exploits exactly this point: how to select a small number of representative examples in order to reduce the size of G , without significantly affecting the quality of the output.

3.1 Input Examples Generation

A crucial role in our approach is played by the two input sets G and V , used to generate and validate rules. We will show that discovering negative rules is the dual problem of discovering positive rules, therefore we assume to be in the setting of generating positive rules. We will explain how to switch to the negative setting at the end of this section.

Given an input KB kb and a predicate $rel \in kb$, we automatically build a generation set G and a validation set V as follows. G consists of positive examples for the target predicate rel . It is generated as all pairs of entities (x, y) such that $\langle x, rel, y \rangle \in kb$ – if the target predicate is **child**, it consists of all pairs of entities in a child relation. V instead consists of counter (negative) examples for the target predicate and it is slightly more complicated to generate, since the closed world assumption does not longer hold in KBs. Differently from classic database scenarios, we cannot assume that what is not stated in a KB is false (closed world assumption). Because of large incompleteness, everything that is not stated in a KB is *unknown* rather false. In order to generate negative examples, we make use of a popular technique for KBs: *Local-Closed World Assumption (LCWA)* [17, 23]. LCWA states that if a KB contains one or more object values for a given subject and predicate, then it contains all possible values (if a KB contains one or more children of Clint Eastwood, then it contains all possible children). This is definitely true for *functional* predicates (predicates such as **capital** where the subject can have at most one object value), while it might not hold for non-functional predicates (e.g., **child**). KBs contain many non-functional predicates, we therefore extend the definition of LCWA by considering the dual aspect: if a KB contains one or more subject values for a given object and predicate, then it contains all possible values.

To generate negative examples we then take the union of the two LCWA aspects: for a given predicate rel , a negative

example is a pair (x, y) where either x is the subject of one or more triples $\langle x, rel, y' \rangle$ with $y \neq y'$, or y is the object of one or more triples $\langle x', rel, y \rangle$ with $x \neq x'$. As an example, if $rel = \text{child}$, a negative example is a pair (x, y) such that x has some children in the KB that are not y , or y is the child of someone that is not x . By considering the union of the two LCWA aspects we do not restrict predicates to be functional (such as in [23]).

The number of negative examples generated with the above technique could easily explode (for a target **child** predicate it is nearly the cartesian product of all the people having a child with all the people having a parent). In order to apply the same approach for positive and negative rules discovery, we require G and V to be of comparable sizes. We therefore introduce a further constraint, that significantly shrinks the size of negative examples: given a pair of entities (x, y) , x must be connected to y via a predicate that is different from the target predicate. In other words, given a KB kb and a target predicate rel , (x, y) is a negative example if $\langle x, rel', y \rangle \in kb$, with $rel' \neq rel$. This intuition exploits the LCWA for predicates rather than for entities. If a KB contains a relation between two entities x and y , then it contains all possible relations between x and y . If x and y are in a relation that is not the target predicate, then most likely x and y are not connected by the target predicate in the real world. This further restriction has multiple advantages: on the one hand it makes the size of V of the same order of magnitude of G (see Section 5), on the other hand it guarantees the existence of a path between x and y , for every $(x, y) \in V$. For positive examples, the existence of a path was already guaranteed since pairs in G are always connected by at least one predicate, the target one.

Eventually V is generated with the intersection of the two constraints defined above. A negative example (x, y) for the target predicate **child** has the following characteristics: (i) x and y are not connected by a **child** predicate; (ii) either x has one or more children (different from y) or y has one or more parents (different from x); (iii) x and y are connected by a predicate that is different from **child**.

Often KBs use same predicates to connect different semantic categories. We may find that a **child** predicate is used not only to connect two entities of type person, but also two companies to denote an ownership relation. In order to enhance the quality of the input examples and avoid cases of mixed unrelated semantic categories, we introduce the *type* restriction when generating G and V . The type restriction requires that for every example pair (x, y) belonging to either G or V , x is always of the same type and y is always of the same type. All modern KBs include entity types (often through the **rdf:type** statement), therefore we make use of this information. For example, G and V for a target **child** predicate are two sets of pairs, where each pair consists in two entities of type person. The type information can be manually provided or, as we will see in Section 5.2, we can automatically compute it from the KB.

In the dual problem of negative rules discovery our approach remains unchanged, we just switch the role of G and V . The generation set becomes V (negative examples), while the validation set becomes G (positive examples). Now it should be more clear why we introduced the second constraint of LCWA on predicates. Since V becomes the generation set in the negative rules setting, V must be small in size and it must guarantee the existence of a path between

pairs of entities in each example. Furthermore, we stress that our approach is independent on how G and V are generated: they could also be manually crafted by some domain experts, which would require additional manual effort.

To the best of our knowledge, RuDiK is the first approach that is capable of discovering both positive and negative rules in KBs. Previous works have put their focus either on the positive setting [2, 23] (TODO: cite Ontological Pathfinding), or on the negative one (TODO: cite Data Lakes Sigmod). However, none of these techniques can be easily adapted to work in the dual scenario.

3.2 Literals and Constants

We defined our target language in Section 2.1 which, other than normal predicate atoms, includes literals comparison. The scope of literals comparison is to enrich the language with smarter comparisons among literal values other than equalities, such as greater than or less than. In order to discover such kind of atoms, the KB graph must contain edges that connect literal values with one (or more) symbol from $\{<, \leq, \neq, >, \geq\}$. As an example, Figure 1 should contain an edge ‘<’ from node “*March 31, 1930*” to node “*March 21, 1986*”. Unfortunately, the original KB does not contain this kind of information, and creating such comparisons among all literals in the KB is unfeasible.

Once again, the use of a generation set G is the key point to introduce literals comparison. Since we discover paths for a pair of entities from G in isolation, thus the size of a graph for a pair of entities is relatively small, we can afford to compare all literal values within a single example graph. This implies the creation of a quadratic number of edges w.r.t. the number of literals in the graph. We will show in the experimental section that within a single example graph the number of literals is usually relatively small, thus the quadratic comparison affordable. Modern KBs include three types of literals: numbers, dates, and strings. Besides equality comparisons, we add ‘>’, ‘≥’, ‘<’, ‘≤’ relationships between numbers and dates, and \neq between all literals. These new relationships are treated as normal atoms: $x \geq y$ is equivalent to $\text{rel}(x, y)$, where rel is equal to \geq . Once we added artificial edges for every input example graph, the discovery of literal comparisons is equivalent to discover normal predicate atoms.

Furthermore, we noticed that ‘ \neq ’ relation could be useful for entities as well other than literals. Think about the following negative rule:

$$\text{bornIn}(a, x) \wedge x \neq b \Rightarrow \neg \text{president}(a, b)$$

The rule states that if a person a is born in a country that is different from b , then a cannot be president of b . The rule holds for most of the countries in the world. To consider inequalities among entities, we could add artificial edges among all pairs of entities in the graph. This strategy however, despite being inefficient for the high number of edges to add, would lead to many meaningless rules. We noticed that it is reasonable to compare two entities only when they are of the same type, e.g., they belong to the same conceptual category. As with the generation of G and V , we make use of **rdf:type** triples. We add an artificial inequality edge in the input example graph only between those pairs of entities of the same type. In the above rule it is reasonable to compare x and b because they are both countries.

Algorithm 1: Greedy Rules Selection.

```

input :  $G$  – generation set
input :  $V$  – validation set
input :  $R$  – universe of rules
output:  $R_{opt}$  – greedy set cover solution
1  $R_{opt} \leftarrow \emptyset$ ;
2  $r \leftarrow \underset{w_m(r)}{\text{argmin}}(r \in R)$ ;
3 repeat
4    $R_{opt} \leftarrow R_{opt} \cup \{r\}$ ;
5    $R \leftarrow R \setminus \{r\}$ ;
6    $r \leftarrow \underset{w_m(r)}{\text{argmin}}(r \in R)$ ;
7 until  $R = \emptyset \vee C_{R_{opt}}(G) = G \vee w_m(r) \geq 0$ ;
8 if  $C_{R_{opt}}(G) \neq G$  then
9    $R_{opt} \leftarrow R_{opt} \cup \text{singleInstanceRule}(G \setminus C_{R_{opt}}(G))$ ;
10 return  $R_{opt}$ 

```

As a last extension of our language, we also discover rules with constants (entities). For a given rule r , we promote a variable v in r to an entity e iff for every $(x, y) \in G$ covered by r , v is always instantiated with the same value e . Suppose that for the above negative rule for president, all examples in G are people born in the country “*U.S.A.*”. We can then promote variable b to “*U.S.A.*”, generating the rule:

$$\text{bornIn}(a, x) \wedge x \neq \text{U.S.A.} \Rightarrow \neg \text{president}(a, \text{U.S.A.})$$

4. A* GREEDY ALGORITHM

We introduce in RuDiK a greedy approach to solve the approximate version of the discovery problem (Section 2.4). The algorithm combines two phases: (i) it solves the set cover problem with a greedy strategy; (ii) it discovers new rules by navigating the graph in a A^* search fashion, allowing the pruning of unpromising paths.

4.1 Marginal Weight

In Section 2.3, we defined the weight associated with a set of rules R as follows:

$$w(R) = \alpha \cdot \left(1 - \frac{|C_R(G)|}{|G|}\right) + \beta \cdot \left(\frac{|C_R(V)|}{|U_R(V)|}\right)$$

Our goal is to discover a set of rules that covers as many elements as possible in G , and as few elements as possible in V . We follow the intuitions behind the greedy algorithm for weighted set cover by defining a *marginal weight* for rules that are not yet included in the solution [14].

Definition 1: Given a set of rules R and a rule r such that $r \notin R$, the marginal weight of r w.r.t. R is defined as:

$$w_m(r) = w(R \cup \{r\}) - w(R) = -\alpha \cdot \frac{|C_r(G) \setminus C_R(G)|}{|G|} + \beta \cdot \left(\frac{|C_{R \cup \{r\}}(V)|}{|U_{R \cup \{r\}}(V)|} - \frac{|C_R(V)|}{|U_R(V)|}\right)$$

□

The marginal weight quantifies the total weight increase that we would have by adding r to an existing set of rules R . In other words, the marginal weight indicates the contribution of r to R in terms of new elements covered in G and new elements uncovered in V . Due to the first negative part, $w_m(r) \in [-\alpha, +\beta]$. Since the set cover problem aims at minimising the total weight, we would never add a rule to

the solution if its marginal weight is greater than or equal to 0. Algorithm 1 shows the straightforward greedy rules selection procedure. The algorithm takes as input the generation set G , the validation set V , and the universe of all possible rules R . The set of output rules R_{opt} is first assigned to an empty set. At each iteration, the algorithm picks from R the rule r with the minimum marginal weight, and it adds r to R_{opt} . r is then removed from R . The algorithm stops when one of the following termination conditions are met: 1) R is empty – all the rules have been included in the solution; 2) R_{opt} covers all elements of G ; 3) the minimum marginal weight is greater than or equal to 0 – among the remaining rules in R , none of them has a negative marginal weight, hence the current solution is the one with the minimum weight. If the second termination condition is not met, there may exist examples in G that are not covered by R_{opt} . In such a case the algorithm will augment R_{opt} with single-instance rules (rules that cover only one example), one for each element in G not covered by R_{opt} .

Since the coverage of a rule is always contained in its unbounded coverage, the marginal weight is greater than or equal to 0 whenever the rule does not cover new elements in G and does not unbound cover new elements in V . More specifically, a rule r has a negative marginal weight iff the sum of its new elements covered in G with its new elements unbounded covered in V is strictly greater than its new elements covered in V multiplied by some γ , where γ depends on how we set α and β .

The greedy solution guarantees a $\log(k)$ approximation to the optimal solution [14], where k is the largest number of elements covered in G by a rule in R – k is at most $|G|$. If the output rules in the final solution cover disjoint sets of G , then the greedy solution coincides with the optimal one.

4.2 A^* Graph Traversal

Algorithm 1 assumes that the universe of rules R has been generated, so that we can iteratively pick the rule with minimum marginal weight. In order to generate the universe of all possible rules, we need to traverse all valid paths from a node x to a node y , for every pair (x, y) in the generation set G . In this section we will analyse the following aspect: is it essential to generate all possible paths for every example in G ?

Example 4: Consider the scenario where we are mining positive rules for the target predicate `spouse`. The generation set G includes two examples g_1 and g_2 , where the above figure shows the KB graph for the two examples. Assume for simplicity that all rules in the universe have the same coverage and unbounded coverage over the validation set V . One of the plausible rule is $r = \text{child}(x, v_0) \wedge \text{child}(y, v_0) \Rightarrow \text{spouse}(x, y)$, stating that entities x and y with a common child are married. Looking at the KB graph, r covers both g_1 and g_2 – in both g_1 and g_2 there exists a path that corresponds to r_{body} . Since all rules have the same coverage and unbounded coverage over V , if we include r in the solution before inspecting other potential rules, then there is no need to generate any other rule. In fact any other plausible rule will not cover new elements in G , therefore their marginal weights will be greater than or equal to 0. Hence any other rule will not be part of the solution. Thus the navigation (and creation) of edges `livesIn` in g_1 , `worksAt` in g_2 , and `partner` in g_2 becomes worthless.

Based on the above observation, we avoid the generation of the entire universe R , but rather we consider at each iteration the most promising path on the graph. The intuition is the same behind the A^* graph traversal algorithm [25]. Given an input weighted graph, A^* computes the smallest cost path from a starting node s to an ending node t . At each iteration, A^* maintains a queue of partial paths starting from s , and it expands one of these paths based on an *estimation* of the cost still to go to t . The path with the best estimation is expanded and added to the paths queue. The algorithm keeps iterating until one of the partial paths reaches t . RuDiK discovers rules with a similar technique. For each example $(x, y) \in G$, we start the navigation from x . We keep a queue of not valid rules (Section 2.1), and at each iteration we consider the rule with the minimum marginal weight, which corresponds to equivalent paths in the example graphs. We expand the rule by following edges on the graphs, and we add the new founded rules to the queue of not valid rules. Differently from A^* , we do not stop when a rule (path) reaches the node y . Whenever a rule becomes valid, we add the rule to the solution and we do not expand it any further. The algorithm keeps looking for plausible paths until one of the termination conditions of Algorithm 1 is met.

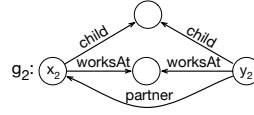
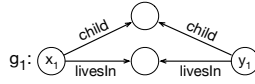
A crucial point in A^* is the definition of the estimation cost. In particular, if we want to guarantee the solution to be optimal, the estimation must be *admissible*, i.e., the estimation cost must be less than or equal to the actual cost. As an example, when searching for the shortest route on a map, an admissible estimation might be the straight-line distance to the goal for every node, since that is physically the smallest possible distance between any two points. In our setting, given a rule that is not yet valid and needs to be expanded, we need to define an admissible estimation of the marginal weight.

Definition 2: Given a rule $r = A_1 \wedge A_2 \cdots A_n \Rightarrow B$, we say that a rule r' is an *expansion* of r iff r' has the form $A_1 \wedge A_2 \cdots A_n \wedge A_{n+1} \Rightarrow B$. In other words, r' is generated by adding a new atom to the body of r . \square Given a

rule r , expanding r means adding a new atom to the body of r . From the graph traversal point of view, expanding r means traversing one further edge on the path defined by r_{body} . In order to guarantee the optimality condition, the estimated marginal weight for a rule r that is not valid must be less than or equal to the actual weight of any valid rule that is generated by expanding r . Given a rule and some expansions of it we can derive the following:

Lemma 4.1: *Given a rule r and a set of pair of entities E , then for each r' expansion of r , $C_{r'}(E) \subseteq C_r(E)$ and $U_{r'}(E) \subseteq U_r(E)$.* \square The above

Lemma states that the coverage and unbound coverage of a rule r' expansion of r are respectively contained in the coverage and unbound coverage of r . In fact expanding a rule means adding a new atom to the body of the rule, hence making the rule more selective. This is equivalent of adding an ‘AND’ condition to a SQL query q : the new result will obviously be a subset of the result obtained with q . Note that Lemma 1 is equivalent to the augmentation inference rule for functional dependencies [4].



We recall that the marginal weight of a rule r w.r.t. a solution R is defined as follow:

$$w_m(r) = -\alpha \cdot \frac{|C_r(G) \setminus C_R(G)|}{|G|} + \beta \cdot \left(\frac{|C_{R \cup \{r\}}(V)|}{|U_{R \cup \{r\}}(V)|} - \frac{|C_R(V)|}{|U_R(V)|} \right)$$

The only positive contribution to marginal weights is given by $|C_{R \cup \{r\}}(V)|$. $|C_{R \cup \{r\}}(V)|$ is equivalent to $|C_R(V)| + |C_r(V) \setminus C_R(V)|$, thus if we set $|C_r(V) \setminus C_R(V)| = 0$ for any r that is not valid, we guarantee an admissible estimation of the marginal weight. More simply, we estimate the coverage over the validation set to be 0 for any rule that can be further expanded, since expanding the rule may bring the coverage to 0.

Definition 3: Given a rule r and a set of rules R , we define the *estimated marginal weight* of r as:

$$w_m^*(r) = \begin{cases} -\alpha \cdot \frac{|C_r(G) \setminus C_R(G)|}{|G|} + \beta \cdot \left(\frac{|C_R(V)|}{|U_{R \cup \{r\}}(V)|} - \frac{|C_R(V)|}{|U_R(V)|} \right) & \text{if } r \text{ is not valid} \\ w_m(r) & \text{if } r \text{ is valid} \end{cases}$$

where a rule is valid iff it respects the language biases defined in Section 2.1. \square

The estimated marginal weight for a valid rule is equal to the actual marginal weight since a valid rule will not be considered for expansion – it corresponds to reaching the ending node in the A^* algorithm. Given Lemma 1, we can easily see that $w_m^*(r) \leq w_m^*(r')$, for any r' expansion of r . Thus our marginal weight estimation is admissible.

RuDiK uses the concept of *frontier nodes* for a rule r ($N_f(r)$). Given a rule r , $N_f(r)$ contains the last visited nodes in the paths that correspond to r_{body} from every example graphs covered by r . As an example, given $r_{body} = \text{child}(x, v_0)$, $N_f(r)$ contains all the entities v_0 that are children of x , for every $(x, y) \in G - v_0$ is the last visited node in the path. Expanding a rule r implies navigating a single edge from any frontier node. Algorithm 2 shows the modified set cover version that includes A^* -like rules generation. The set of frontier nodes is initialised with starting nodes x , for every $(x, y) \in G$ (Line 2). The algorithm maintains a queue of rules Q_r , from which it chooses at each iteration the rule with minimum approximated weight. The function **expandFrontiers** retrieves from the KB all nodes (along with edges) at distance 1 from frontier nodes and returns the set of all rules generated by this one hop expansion. Such expansions are computed with single-hop SPARQL queries. Q_r is therefore initialised with all rules of length 1 starting at x (Line 3). In the main loop, the algorithm checks if the current best rule r is valid or not. If r is valid, r is added to the output and it is not expanded (Line 8). If r is not valid, r is expanded iff the length of its body is less than $maxPathLen$ (Line 10). There is no point in expanding rules with body greater than or equal to $maxPathLen$ since such rules are not allowed in the output. The termination conditions and the last part of the algorithm are the same of Algorithm 1.

Algorithm 2: RuDiK Rules Discovery.

```

input :  $G$  – generation set
input :  $V$  – validation set
input :  $maxPathLen$  – maximum rule body length
output:  $R_{opt}$  – greedy set cover solution

1  $R_{opt} \leftarrow \emptyset$ ;
2  $N_f \leftarrow \{x | (x, y) \in G\}$ ;
3  $Q_r \leftarrow \text{expandFrontiers}(N_f)$ ; // SPARQL queries
4  $r \leftarrow \text{argmin}(r \in Q_r)$ ;
    $w_m^*(r)$ 
5 repeat
6    $Q_r \leftarrow Q_r \setminus \{r\}$ ;
7   if isValid( $r$ ) then
8      $R_{opt} \leftarrow R_{opt} \cup \{r\}$ ;
9   else
10    // rules expansion
11    if  $\text{length}(r_{body}) < maxPathLen$  then
12       $N_f \leftarrow \text{frontiers}(r)$ ;
13       $Q_r \leftarrow Q_r \cup \text{expandFrontiers}(N_f)$ ; // SPARQL queries
14     $r \leftarrow \text{argmin}(r \in Q_r)$ ;
        $w_m^*(r)$ 
15 until  $Q_r = \emptyset \vee C_{R_{opt}}(G) = G \vee w_m^*(r) \geq 0$ ;
16 if  $C_{R_{opt}}(G) \neq G$  then
17    $R_{opt} \leftarrow R_{opt} \cup \text{singleInstanceRule}(G \setminus C_{R_{opt}}(G))$ ;
18 return  $R_{opt}$ 

```

The simultaneous rules generation and selection of Algorithm 2 brings multiple benefits. First of all, we do not generate the entire graph for every example in G . Nodes and edges are generated *on demand*, whenever the algorithm requires their navigation (Line 12). If the initial part of a path is not promising according to its estimated weight, the rest of the path will never be materialised since there is no need to navigate it. Materialising the graph is an expensive task, since we need to query the disk in order to retrieve target nodes and edges. Rather than materialising the entire graph and then traversing it, we propose a solution that gradually materialises parts of the graph whenever they are needed for navigation (Lines 3 and 12). Secondly, the weight estimation could lead to pruning unpromising rules. If a rule does not cover new elements in G and does not unbound cover new elements in V , then its estimated marginal weight is 0 (Definition 3). A rule with 0 marginal weight is never picked as best rule to be expanded, and if it is the algorithm terminates (due to one of the termination conditions). This implies that a rule with 0 estimated marginal weight is pruned away from the search space, as we will never generate rules from its expansion.

The partial materialisation of the graph and the pruning of the search space have a significant impact on the algorithm's resources. On the one hand, we noticed the running time is halved in the worst case and sometimes it is up to ten times faster. On the other hand, we can also significantly decrease the amount of memory needed, since we load sub-portions of the graph. Consider the extreme case where there exists a rule r that covers all elements in G , unbound

covers all elements in V and does not cover any element of V . In such a case, Algorithm 2 will output the optimal solution by just materialising paths on the example graphs that correspond to r_{body} with $\mathcal{O}(l \cdot |G|)$ SPARQL queries, where l is the length of r_{body} .

5. EXPERIMENTS

We carried out an extensive experimental evaluation of our rules discovery approach. We grouped the evaluation into 4 main sub-categories: (i) a first set of experiments aims at demonstrating the quality of our output, both for negative and positive rules; (ii) a second set of experiments compares our method with state-of-the-art systems; (iii) in the third set of experiments we outline the applicability of rules discovery by enhancing machine learning algorithms; (iv) in the last set of experiments we discuss internal system settings and some KB properties.

Settings. We evaluated our approach over several popular KBs. For each KB, we downloaded the most up-to-date core facts and loaded them into our SPARQL query engine. We experimented several SPARQL engines, including Jena ARQ³, OWLIM Lite⁴, and RDF-3x⁵. We also implemented a naïve relational database solution with PostgreSQL. Eventually we opted for OpenLink Virtuoso⁶, as it was the fastest among all the solutions. Virtuoso took on average 20 minutes to load a medium size KB (i.e., 10 GB) into its store, and around 100 milliseconds to execute a single hop query. All experiments are run on a iMac desktop with an Intel quad-core i5 at 2.80GHz with 16 GB RAM. We run Virtuoso server with its SPARQL query endpoint on the same machine, optimised for a 8 GB available RAM.

Our method needs the two input parameters α and β of Equation 1. α measures the importance of the coverage over the generation set, while β measures the coverage over the validation set. In other words, a high α privileges recall over precision, while a high β gives more importance to precision. We can afford a high α and a low β when the input KB is accurate and complete. We will show that KBs contain many errors and missing information, therefore we set $\alpha = 0.3$ and $\beta = 0.7$. Increasing α and decreasing β means a higher number of output rules with a lower accuracy.

We also set the *maxPathLen* parameter to 3. This number represents the maximum number of atoms that we can have in the body of a rule. We will show in Section 5.4 that increasing this parameter does not bring any benefits, as body rules longer than 3 atoms start to be very complicated and not insightful.

Evaluation Metrics. RuDiK discovers rules for a given target predicate. For each KB, we chose 5 representative predicates as follows: we first ordered predicates according to descending popularity (i.e., number of triples having that predicate), and then we picked the top 3 predicates for which we knew there existed at least one meaningful rule, and other 2 top predicates for which we did not know whether some meaningful rules existed. We repeated the procedure for each input KB, and for positive and negative rules. Despite working one predicate at time, RuDiK can also discover rules

for the entire KB by listing all predicates in the KB, and discover rules for each of them. We will show in Section 5.2 how this can be achieved.

Positive rules are very useful to enrich the KB by discovering new facts. Since we are generating new data, we cannot evaluate the output over the existing data. Inspired by [23], we proceed as follows: we run the algorithm over the KB, and for each output rule we generate all new predictions that are not already in the KB (we execute the body of the rule against the KB and remove all those pairs that are already connected by the target predicate in the KB). As an example, for the rule $\text{spouse}(b, a) \Rightarrow \text{spouse}(a, b)$, we retrieve all pairs (b, a) such that b is *spouse* with a but a is not *spouse* with b in the KB. If the rule is universally correct (like the previous example), we mark all the new predictions as true. If the rule is unknown, we randomly sampled 30 new predictions and manually checked them against the Web. The *precision* of the rule is then computed as the ratio of true predictions out of true and false predictions.

Negative rules are slightly more complicated to evaluate. In fact, despite KBs are usually incomplete, a large percentage of the data not stated in a KB is false – a very small fraction of the cartesian product of all the people in a KB will be actually married. Therefore negative rules will always discover many correct negative facts. However, negative rules are a great means to discover errors in the KB, and we leverage this aspect to evaluate them. For each discovered rule, we retrieve from the KB pairs of entities for which the body of the rule can be instantiated over the KB and that are also connected by the target predicate. As an example, for the rule $\text{child}(a, b) \Rightarrow \neg \text{spouse}(a, b)$, we retrieve all those pairs (a, b) such that b is *child* of a and a is *spouse* with b . We call these generated pairs of entities *potential errors*. Similarly to positive rules, whenever a rule is universally correct we mark all its potential errors as true, whereas if the rule is unknown we manually check 30 sampled potential errors. The final precision of a rule is computed as actual errors divided by potential errors. Furthermore, a potential error is evaluated as actual error whenever a single atom in the rule is incorrect, no matter if the atom is in the body or the head of the rule. For the negative rule $\text{child}(a, b) \Rightarrow \neg \text{spouse}(a, b)$, a specific instantiation (a, b) is an actual error if either b is not an actual *child* of a , or b is not an actual *spouse* of a . As we will explain further below, negative rules cannot point exactly where the error is, but they give a hint that something is wrong and needs to be further inspected.

The full suite of test results, together with all the KBs, induced rules, and annotated gold standard examples and rules is available online (TODO: add footnote online appendix).

5.1 Rules Discovery Accuracy

The first set of experiments aims at evaluating the accuracy of discovered rules over the 3 most popular and widely used KBs: DBPEDIA [8], YAGO [34], and WIKIDATA [37]. For each KB we downloaded the most recent version and selected core facts, facts about people, geolocations and transitive *rdf:type* facts. WIKIDATA provides only the entire dump, therefore we just eliminated from it no-english literal values. Table 1 shows the characteristics of the 3 KBs.

As the figure shows, the size of the KB is relevant. Loading the entire KB into main memory is not feasible unless

³<https://jena.apache.org/documentation/query/>

⁴<https://confluence.ontotext.com/display/OWLIMv54/OWLIM-Lite+Installation>

⁵<https://code.google.com/archive/p/rdf3x/>

⁶<http://virtuoso.openlinksw.com/>

Table 1: Dataset characteristics.

<i>KB</i>	<i>Version</i>	<i>Size</i>	<i>#Triples</i>	<i>#Predicates</i>
DBPEDIA	3.7	10.056GB	68,364,605	1,424
YAGO	3.0.2	7.82GB	88,360,244	74
WIKIDATA	20160229	12.32GB	272,129,814	4,108

we have high memory availability (TODO: cite the two SIGMOD16 papers), or we reduce the KB by eliminating facts such as `rdf:type` or literals [23]. We propose an approach that is disk-based where only a small portion of the KB is loaded into main memory, such that we can discover rules on any size KB with limited resources.

5.1.1 Positive Rules Discovery

We first evaluate the precision of positive rules for the top 5 predicates on the 3 KBs. The number of new induced facts varies significantly from rule to rule – a rule with literals comparison will produce a very high number of facts. In order to avoid the precision to be dominated by such rules, we first compute the precision for each rule as explained above, and then we average values over all induced rules. Table 2 reports precision values, along with predicates average running time.

Our first observation is that the more accurate is the KB, the better is the quality of induced rules. WIKIDATA contains very few errors, since it is manually curated and every triple is manually checked by different individuals before being inserted. DBPEDIA and YAGO instead are automatically generated by extracting information from the Web, hence their quality is significantly lower. Discovering perfect positive rules is a hard task, mostly because there is no guarantee of the existence of valid negative examples. A striking example in this direction is one of the rule induced for `founder` in DBPEDIA. Our approach discovers that if a person is born in the same place where a company is founded, then the person is the founder of the company. The rule is obviously wrong, as there are many people who are born in the same place of a company and have not founded the company. However this rule has a very high coverage over the generation set (many companies’ founders founded their company in their birth place), and a very low coverage over the validation set – indeed among the cartesian product of all the people and companies, a very small fraction includes people and companies born and founded in the same place. Despite such hard cases, our approach is always capable of producing correct rules for those predicates for which we knew there existed some valid rules. Cases like `academicAdvisor`, `child`, and `spouse` have a precision above 95% in all of the KBs, and final precision values are brought down by few predicates where meaningful rules probably do not exist at all.

The running time is influenced by different factors. First of all the size of the KB has obviously a huge impact, as RuDiK is slower in WIKIDATA which is the biggest KB. Not only the number of triples is relevant, but also the differ-

ent number of predicates. In fact the more predicates we have in the KB, the more alternative paths we observe when traversing the graph, hence a bigger search space. The second relevant aspect is the target predicate involved. We noticed that some kind of entities have a huge number of outgoing and incoming edges (“*United States*” in WIKIDATA is connected to more than 600K entities). When the generation set includes such types of entities, the navigation of the graph is slower as we need to traverse a high number of edges. This is what happens in YAGO, where the most popular predicates are `isLeaderOf` and `exports`. Eventually the `maxPathLen` parameter also has a big say in the final running time. The longer the rule, the bigger is the search space. We will show in the next Section how we can be much faster if we set to 2 atoms the maximum length of the rule. In the next chapter we will discuss some future directions to significantly cut down the running time based on the above observations.

5.1.2 Negative Rules Discovery

Negative rules are very useful to discover inconsistencies in the KB. We evaluated negative rules as the percentage of correct errors discovered for the top 5 predicates in each KB. Table 3 shows, for each KB, the total number of potential erroneous triples discovered with negative rules, whereas the precision is computed as the percentage of actual errors among potential errors.

Negative rules generally have better accuracy than positive ones. This is mostly due to the completeness of the validation set: for negative rules the validation set is the universe of all possible counter examples stated in the KB, whereas for positive rules the validation set is just a small fraction of it. Therefore negative rules are usually better validated than positive ones. WIKIDATA shows lower numbers just because it does not contain as many errors as DBPEDIA and YAGO: even though discovered rules are almost correct, the percentage of actual errors identified is lower in WIKIDATA. For example, we identify the same rule that two people with same gender cannot be married both in YAGO and WIKIDATA. Such a rule retrieves errors in YAGO with 94% accuracy, while the accuracy in WIKIDATA for the same rule is 57%. YAGO is the KB with the highest number of errors. As an example, there are 9,057 cases in the online YAGO where a child is born before her parent. We cannot point exactly where the error is – there could be an error in one of the birth dates or an error in the parent relation – but we can affirm that at least one of these values is wrong and send them to human evaluators to spot the inconsistency.

Differently from positive rules, literals play a vital role in discovering negative rules. In fact in many cases correct negative rules rely on temporal aspects in which something cannot happen before/after something else. Temporal information are usually expressed through dates, years, or other primitive types that are represented as literal values in KBs.

Table 2: Positive Rules Accuracy.

<i>KB</i>	<i>Avg. Running Time</i>	<i>Precision</i>
DBPEDIA	34min, 56sec	63.99%
YAGO	59min, 25sec	62.86%
WIKIDATA	2h, 21min, 34sec	73.33%

Table 3: Negative Rules Accuracy.

<i>KB</i>	<i>Avg. Run Time</i>	<i># Potential Errors</i>	<i>Precision</i>
DBPEDIA	19min, 40sec	499	92.38%
YAGO	10min, 40sec	2,237	90.61%
WIKIDATA	1h, 5min, 38sec	1,776	73.99%

As a last observation, discovering negative rules is usually faster than discovering positive rules. This is mostly due to the time we spend executing validation coverage queries. Whenever we discover a rule that respect our language bias (Section 2.1), we execute the body of the rule against the KB with a SPARQL query to compute its coverage over the validation set. These queries are faster for negative rules since the validation set is just all the entities connected by the target predicate, whereas in the positive case the validation set corresponds to counter examples described in Section 3.1, which are usually more complex to evaluate for standard SPARQL engines.

Table 3 shows some interesting cases, for both correct and incorrect rules. The example rules show the full power of our language, including literals, smart literals comparisons, entities inequalities and constants. As previously mentioned, the extension including literals comparisons and entities inequalities gives a significant boost in accuracy for negative rules, while it is rarely used in discovery positive rules. The full set of discovered rules can be inspected online (TODO: footnote online appendix).

Rule
$\text{notableWork}(b, a) \Rightarrow \text{creator}(a, b)$
$\text{nationality}(a, v_0) \wedge \text{nationality}(b, v_0) \wedge \text{notableStudent}(b, a) \Rightarrow \text{academicAdvisor}(a, b)$
$\text{hasChild}(v_0, b) \wedge \text{isMarriedTo}(a, v_0) \Rightarrow \text{hasChild}(a, b)$
$\text{activeYearsEndDate}(a, v_0) \wedge \text{activeYearsStartDate}(b, v_0) \Rightarrow \text{successor}(a, b)$
$\text{inception}(a, v_0) \wedge \text{dateOfBirth}(b, v_1) \wedge v_1 > v_0 \Rightarrow \neg \text{founder}(a, b)$
$\text{country}(a, \text{England}) \wedge \text{country}(b, \text{United.Kingdom}) \wedge \text{England} \neq \text{United.Kingdom} \Rightarrow \neg \text{country}(a, b)$
$\text{actedIn}(a, b) \Rightarrow \neg \text{wroteMusicFor}(a, b)$
$\text{birthYear}(a, v_0) \wedge \text{birthDate}(b, v_1) \wedge v_0 < v_1 \Rightarrow \neg \text{academicAdvisor}(a, b)$

Table 4: Interesting Output Rules.

5.2 Comparative Evaluation

We compared the performance of our rules discovery method against AMIE [23], the state-of-the-art system in discovery Horn Rules from KBs.

AMIE is a rules discovery system designed to discover positive rules. It first loads the entire KB into memory, and then discovers positive rules for every predicate in the KB. AMIE lists all the predicates in the KB and inserts each of them as head of the rule. Once the head is filled, the system tries to expand the rule by pivoting on one of the variables of the current predicate and looks for predicates sharing the same variable with high coverage in the KB. The coverage of a rule is penalised with the partial closed world assumption, where the set of negative examples for a given pair (x, y) and a target predicate p is all those pairs where x is connected through p to an entity different from y . Differently from us, AMIE outputs all possible rules that exceeds a given threshold and ranks them according to their coverage function.

Given the in-memory implementation, AMIE cannot handle large KBs. We tried to run it on the KBs of Table 1, but the system goes quickly out of memory. Thus we downloaded and used the modified versions of YAGO and DBPEDIA used in their experiments⁷. These versions consist

⁷www.mpi-inf.mpg.de/departments/databases-and-information-systems/research/yago-naga/amie/



Figure 2: Predictions Accuracy on Yago

in the core facts of the KB, without literals and `rdf:type` facts. Table 5 summarises the characteristic of this dataset.

Table 5: AMIE Dataset characteristics.

KB	Size	#Triples	#Predicates	# <code>rdf:type</code>
DBPEDIA	551M	7M	10,342	22.2M
YAGO	48M	948.3K	38	77.9M

Removing literals and `rdf:type` triples drastically reduce the size of the KB (Table 1). Since our approach needs type information (both for the generation of G and V and for the discovery of entities inequality atoms), we run AMIE on its original dataset, while we run our algorithm on the same dataset plus `rdf:type` triples. The last column of Table 5 shows how many triples we added to the original AMIE dataset.

Positive Rules. We first compared RuDiK against AMIE on its natural setting: positive rules discovery. AMIE takes as input an entire KB, and discovers rules for every predicate in the KB. We adapted our system to simulate AMIE as follows: we first list all the predicates in the KB, and for each predicate that connects a *subject* to an *object* we computed the most common type for both subject and object. The most common type is computed as the most popular `rdf:type` that is not super class of any other most popular type.

After computing type domain and co-domain for each predicate, we run our approach sequentially on every predicate. Furthermore we set the `maxPathLen` parameter to 2, since this is the default setting for AMIE.

AMIE outputs a huge amount of rules along with their scores – 75 output rules in YAGO, and 6090 in DBPEDIA. We followed their experiments setting and picked the first 30 best rules according to their score. We then picked the rules produced by our approach on the head predicate of the 30 best rules output of AMIE. RuDiK is more conservative and produces much less rules than AMIE. We noticed that for every predicate AMIE always discovers more than one rule, while there are several predicates where the output of our algorithm is empty since none of the plausible rules has enough support. As a consequence, for instance, RuDiK outputs just 11 rules for 8 target predicates on the entire YAGO – for the remaining predicates RuDiK does not find any rule with enough support.

The precision of each rule is computed as described above with a minor modification: whenever a rule is unknown with its new predictions, we first check the existence of the new induced facts in a newer version of the KB. This is possible because the AMIE dataset does not contain the most up to date versions. If a new fact does not appear neither in a newer version nor on the Web, it is evaluated as false.

Figure 2 plots the total number of new unique predictions (x-axis) versus the aggregated precision (y-axis) on YAGO.

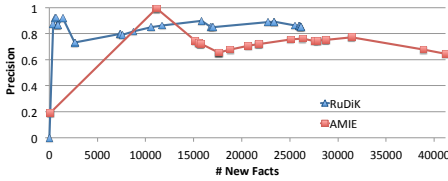


Figure 3: Predictions Accuracy on DBPedia

The n -th point from the left represents the total number of predictions and the total precision of these predictions, computed over the first n rules (sorted according to AMIE’s score). AMIE produces many more predictions (262K vs 102K), but with a significant lower accuracy. This is due to the high number of rules in output of AMIE, but also to the way these rules are ranked. In fact if we limit the output of AMIE to the best 11 rules (same output of our approach), the final accuracy is still 29% below our approach, with just 10K more predictions. AMIE outputs many good rules that are preceded by meaningless ones in the ranking, and it is not clear how to set a proper k in order to get the best top k rules. RuDiK instead understands that in some cases meaningful rules do not exist, and it outputs something only when it has a strong confidence. This results in a lower number of predictions with a very high accuracy – precision is above 85% before the last rule, with more than 80K predictions. Moreover, our approach can also simulate AMIE if we are more interested in recall. By modifying the α and β parameters, we can obtain a higher number of predictions in output, at the expense of accuracy.

Figure 3 shows the same evaluation on DBPEDIA. DBPEDIA has a richer set of relations, therefore also RuDiK is capable of producing 30 rules in output. Despite the same number of rules, once again our approach leads to a lower number of predictions (26K vs 41K) with a significant higher accuracy (85% vs 74%). The only point where AMIE outperforms our approach is when we consider the top 3 rules: the third rule discovered by AMIE is indeed a universally true rule that produces more than 11K correct predictions.

Negative Rules. As a second set of comparative experiments we used AMIE to discover negative rules. AMIE is not designed to work in this setting, and can discover rules only for predicates explicitly stated in the KB. Therefore we proceeded as follows: we sampled the top 5 most popular predicates in each KB and we created for each predicate a set of negative examples as explained in Section 3.1. For each negative example we added a new fact to the KB connecting the two entities with the *negation* of the predicate. For example, we added a `notSpouse` predicate connecting each pair of people who are not married according to our negative examples generation technique. We then run AMIE on these new created predicates. The evaluation of negative rules is then carried out as explained before: we generate potential errors in the KB, and we manually evaluated the precision of such errors. Table 6 shows the results on the two KBs. RuDiK outperforms AMIE in both cases of almost 20%. This is because for the negation of a predicate, we use the actual predicate as counter examples. AMIE instead is not aware that the actual predicate provides counter examples for the negation of it, hence it is much less precise. In fact the output of AMIE consists in a high number of rules for each predicate (often more than 30), and in many cases

Table 6: Negative Rules vs AMIE.

KB	AMIE		RuDiK	
	# Errors	Precision	# Errors	Precision
DBPEDIA	457	38.85%	148	57.76%
YAGO	633	48.81%	550	68.73%

Table 7: Run Time vs AMIE.

KB	#Triple	#Predicates	AMIE	RuDiK	Types Time
YAGO 2	948.3K	20	30s	18m,15s	12s
YAGO 2s	4.1M	26 (38)	> 8h	47m,10s	11s
DBPEDIA 2.0	7M	904 (10342)	> 10h	7h,12m	77s
DBPEDIA 3.8	11M	237 (649)	> 15h	8h,10m	37s
WIKIDATA	8.4M	118 (430)	> 25h	8h,2m	11s
YAGO 3*	88.3M	72	-	2h,35m	128s

AMIE produces same rules for both positive and negative scenarios. As an example, AMIE outputs that if a country a exports a good b , then a imports b and a does not import b .

Despite clearly outperforming AMIE, numbers look significantly lower than the ones showed in Section 5.1. This is because we are using the AMIE modified KBs which do not contain literals. As explained earlier, literals play a vital role when discovering negative rules, both in terms of total errors discovered and in terms of precision. Excluding literals is a big disadvantage, and we will emphasise this aspect in detail in Section 5.4. (TO DO: check if the internal evaluation actually exists)

Running Time. We report here the running time of AMIE compared to our approach. Note that numbers are different from [23], where AMIE was run on a 48 GB RAM server. AMIE could finish the computation only on YAGO 2, while for the other KBs it got stuck without outputting any more rules. When this happened, we stopped the computation after we did not see any new rule in output for more than 2 hours.

Table 7 reports the running time on different KBs. The first five KBs are the AMIE modified versions, while YAGO 3* is complete YAGO, including literals and `rdf:type`. The third column shows the number of predicates for which AMIE was able to produce at least one rule. In some cases AMIE got stuck without producing any rules for some predicates, hence we report the total number of predicates in brackets. For a fair comparison we run our algorithm only on those predicates for which AMIE could produce at least one rule. The fourth and fifth columns report the total running time of the two approaches. Despite being disk-based, RuDiK can successfully complete the task faster than AMIE in all cases, except for YAGO 2. This is because of the very small size of the KB, which can easily fits in main memory. However, when we deal with real KBs (YAGO 3*), AMIE is not even capable of loading the KB due to out of memory errors. Eventually the last column reports the total time needed to compute `rdf:type` information for each predicate in the KB. This time is negligible w.r.t. the total running time. The running time justifies our disk-based strategy: RuDiK can successfully discover rules for any size KB on any machine.

Other Systems. We found other available systems to discover rules in KBs. [2] discovers new facts at instance level, hence less generic than our approach. On AMIE YAGO 2 KB they can discover 2K new facts with a precision lower than 70%. The best rule we discover on YAGO 2 already produces more than 4K facts with a 100% precision (see on-

line appendix for detailed data (TODO: footnote online appendix)). (TODO: CITE Ontological Path Finding) implements AMIE algorithm with a focus on scalability. They do not introduce any novelty in the algorithmic part, but just a clever way of splitting the KB into multiple cluster nodes so that the computation can be run in parallel. The output is the same as AMIE. Eventually we did not compare with classic Inductive Logic Programming systems [15, 29], as these are already significantly outperformed by AMIE both in accuracy and running time.

5.3 Machine Learning Application

The main goal of this set of experiments is to prove the applicability of our approach in helping Machine Learning algorithms to provide meaningful training examples. We chose DeepDive [33], a Machine Learning approach to incrementally construct KBs. DeepDive extracts entities and relations from text articles via distant supervision. The key idea behind distant supervision is to use an external source of information (e.g., a KB) to provide training examples for a supervised algorithm. For example, the main showcase in DeepDive extracts mentions of married couples from text documents. In such a scenario DeepDive uses as a first step DBPEDIA in order to label some pairs of entities as *true* positive (those pairs of married couples that can be found in DBPEDIA). These labelled examples are then used to construct a factor graph, similar to Markov Logic, that will predict labels on the rest of candidates. Unfortunately, a KB can only provide positive examples. Hence in DeepDive the burden of creating negative examples is left to the user through manual rules definition.

In this set of experiments we will use our negative rules on DBPEDIA to generate negative examples, and we will compare the output of DeepDive trained with different set of negative examples. We used DeepDive showcase example⁸, where the goal is to extract mentions of married people from text articles. DeepDive already provides some negative rules to generate negative examples (e.g., if two people appear in a sentence connected by the words *brother* or *sister* then they are not married). We therefore compare the output of DeepDive using our generated negative examples and the ones generated with DeepDive rules.

Figure 4 shows DeepDive accuracy plot run on 1K input documents. The accuracy plot shows the ratio of correct positive predictions over positive and negative predications (y-axis), for each probability output value (x-axis). The dotted blue line represents the ideal situation, where the system finds high number of evidence positive predictions for higher probability output values – when the output probability is 0 there should not be positive predictions. The plot is computed over a test set, while the system is trained over a separated training set. The figure shows 4 lines other than the ideal ones. RuDiK is the output of DeepDive using our approach to generate negative examples. OnlyPos uses only positive examples from DBPEDIA, Manual uses positive examples from DBPEDIA and manually defined rules to generate negative examples, while ManualSampl uses only a sample of the manually generated negative examples in size equal to positive examples. The first observation is that OnlyPos and Manual do not provide valid training, as the former has only positive examples and labels everything as true, while the latter has many more negative examples than

⁸<http://deepdive.stanford.edu/>

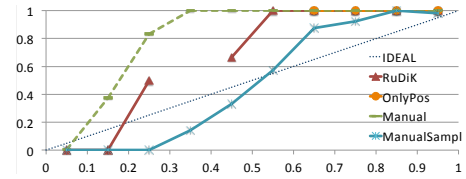


Figure 4: DeepDive Application 1K articles

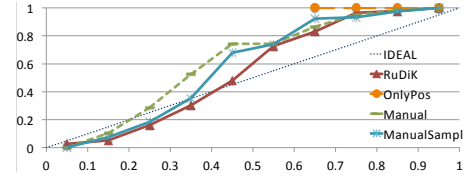


Figure 5: DeepDive Application 1M articles

positive and labels everything as false. ManualSampl is the clear winner, while our approach suffers mostly the absence of data: over the input 1K articles, we could find only 20 positive and 15 negative examples from DBPEDIA. The lack of enough evidence in the training data also explains the missing points for RuDiK in the chart, where there are not predictions in the probability range 25–45%.

If we extend the input to 1M articles, things change drastically (Figure 5). All the three approaches except OnlyPos can successfully drive DeepDive in the training, with the examples provided with RuDiK leading to a slightly better result. This is because of the quality of the negative examples: our negative rules generate representative examples that can help DeepDive in understanding discriminatory features between positive and negative labels. The output of ManualSampl and RuDiK are very similar, meaning that we can use our approach to simulate user behaviour and provide negative examples. With manually defined rules the number of generated examples is significantly higher (23K vs 5K generated with RuDiK), however the results are very similar since a small number of significant examples is enough to provide complete evidence for the training. This confirms the main finding of this experiment: as long as we have an external source of information with a decent coverage over the input articles, users do not need to worry about providing rules to generate negative examples.

5.4 Internal Evaluation

In this last set of experiments we measured the impact of RuDiK relevant features in order to quantify the benefit of three main aspects in rules discovery. We run RuDiK on DBPEDIA with different settings than the standard ones. We report results on the same top-5 predicates of Section 5.1 for both positive and negative rules.

Effect of Literals. Since previous KB rules discovery approaches exclude literals from the mining [2, 23] (TODO: cite Sigmod ontological), we wanted to quantify the impact of having literal rules. Thus we run RuDiK excluding all literal values. Table 8 reports the output precision with and without literals. Including literal values in the mining has a considerable impact on final accuracy, both for positive and negative rules. The effect is particularly evident for negative rules, where excluding literals involves finding less than half potential errors (numbers in brackets) with a lower precision.

founder is the most evident example: RuDiK discovers 79 potential errors with a 95% precision with literal rules, while there are not output errors with rules without literals.

Surprisingly, including literals reduces also the running time. This is due to the pruning effect of the A^* search: if we include literals the algorithm can find rather soon valid literal rules, which causes the pruning of several paths on the graph. If we exulted literals instead these paths cannot be pruned and needs to be inspected by the algorithm, which entails a bigger search space.

Rules Length Impact.

Negative Examples Generation. A key point in RuDiK is the generation of negative examples with a modified version of the LCWA (Section 3.1). We therefore evaluate two different strategies to generate negative examples. Given a target predicate p from a KB kb , we call t_a the most common type of entities that are subject of p , and t_b the most common type for the object. As an example, if $p = \text{founder}$, $t_a = \text{Company}$ and $t_b = \text{Person}$. We define k as the number of triples having p as predicate in kb – k is the cardinality of the positive examples set. The first alternative negative examples generation strategy is *Random*: we randomly select k pairs (x, y) from the cartesian product of all entities x of type t_a and all entities y of type t_b , such that the triple $\langle x, p, y \rangle \notin kb$. The second generation strategy, named *LCWA_Random*, leverages on the LCWA. In *LCWA_Random* we randomly pick k pairs (x, y) from the cartesian product of all entities x of type t_a and all entities y of type t_b , such that: (i) $\langle x, p, y' \rangle \in kb$, with $y' \neq y$; (ii) $\langle x', p, y \rangle \in kb$, with $x' \neq x$; (iii) $\langle x, p, y \rangle \notin kb$. *LCWA_Random* is equivalent to RuDiK generation strategy, minus the constriction that x and y must be connected by a predicate different from p . The constraint of having k examples aims at reducing the size of the cartesian product that can easily explode otherwise.

Table 10 reports the accuracy of discovering negative rules with the three different generation strategies (RuDiK is our modified LCWA strategy). *Random* and *LCWA_Random* strategies show very similar behaviours, with a slightly better precision than RuDiK. This is because whenever we randomly pick examples from the cartesian product of subject and object, the likelihood of picking entities from a different time period is very high, and negative rules pivoting on time constraints are usually correct. As an example, a correct rule for the target predicate **child** is $\text{birthDate}(a, v_0) \wedge \text{birthDate}(b, v_1) \wedge v_0 > v_1 \Rightarrow \neg \text{child}(a, b)$, stating that whenever a person a is born after a person b , then b cannot be child of a . The likelihood of randomly picking some pairs of persons (x, y) where x is born after y is very high, hence such kind of rules will often be discovered. However these are the *only* kind of rules that random generation strategies are able to find, as shown from the smaller number of errors that they can identify. Instead, the constriction of forcing x and y to be connected by a different predicate generates negative examples with heterogeneous properties that lead to differ-

Type	With Literals		Without Literals	
	Run Time	Precision	Run Time	Precision
Positive Rules	~35min	63.99%	~54min	60.49%
Negative Rules	~20min	92.38% (499)	~25min	84.85% (235)

Table 8: Rules Accuracy without Literals on DBPedia.

Table 9: maxPathLen Parameter Impact on DBPedia.

Type	$\text{MaxPathLen} = 2$		$\text{MaxPathLen} = 3$		$\text{MaxPathLen} = 4$
	Run Time	Precision	Run Time	Precision	Run Time
Positive	~3min	49.17%	~35min	63.99%	
Negative	~56sec	90% (131)	~20min	92.38% (499)	

Table 10: Effect of Negative Examples Generation Strategy on DBPedia.

Strategy	# Potential Errors	Precision
Random	247	95.95%
LCWA_Random	263	95.82%
RuDiK	499	92.38%

ent results. Rules such as $\text{parent}(a, b) \Rightarrow \neg \text{spouse}(a, b)$ are very unlikely to be generated with random strategies, since the likelihood of picking two people that are in a parent relation is very low. The generation strategy we adopt in RuDiK allows the discovery of more types of rules, and not only rules involving time constraints. This has multiple benefits: on the one hand, we discover more rules, which entails discovering a higher number of errors (Table 10); on the other hand, different rule types lead to high quality negative examples, that can be used in several applications such as training Machine Learning algorithms (see Section 5.3).

Effect of Weight Parameters. Our weight function is ruled by two different parameters: α and β , with $\alpha \in [0, 1]$ and $\beta = 1 - \alpha$ (Section 2.3). This experiment aims at establishing the optimal value to assign to α and β , and to quantify the impact of the two parameters on the overall performance. We recall that α quantifies the relevance of the coverage over the generation set, while β is the importance of the coverage over the validation set. In other words, a high α supports high recall over precision, while a high β supports a high precision over recall. Figure ?? shows the variation of F_1 -Score with different values of α for both positive and negative rules on DBPEDIA. For each different run, we manually label each output rule as true or false, where the total number of correct rules for each predicate is the union of all possible correct rules over all the runs. We then compute Precision, Recall, and F_1 -Score at rule level. On the one hand, setting $\alpha = 0$ produces an empty output, since we neglect the coverage over the generation set and we are just after rules that do not cover any element of the validation set. On the other hand, setting $\alpha = 1$ means chasing rules just based on the coverage over the generation set, no matter what the coverage over the validation set is. This strategy ends up in producing a huge amount of rules (high recall), with very few correct rules (low precision). Correct values lie somewhere in between. For positive rules, the best assignment is $\alpha = 0.3$ and $\beta = 0.7$. Since discovering correct positive rules is more challenging than negative ones, favoring precision over recall gives the best accuracy. For negative rules instead, the best assignment is $\alpha = 0.4$ and $\beta = 0.6$. Since negative rules are often correct, we can relax the constraint over precision and be slightly more recall oriented. In both cases, the variation in performance for $\alpha \in [0.1, 0.9]$ is anyway limited ($\leq 12\%$), showing the robustness of the set cover problem formulation.

A^* pruning impact. Another aspect we wanted to quantify is the benefit brought by the A^* algorithm on the

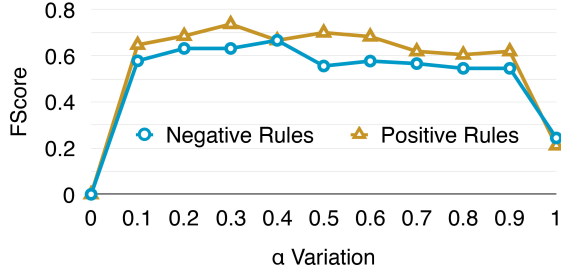


Figure 6: α Parameter Performance Impact

overall running time. Figure 7 shows the running time, for each target predicate, of the A^* algorithm (light-colored bars) against a modified version that first generates the universe of all possible rules, and then apply the greedy set cover algorithm on such a universe (dark-colored bars). For the last two predicates refer to the y-axis labels on the right hand side, as these predicates have a significant higher running time. In the figure (P) refers to positive rules, while (N) to negative rules. The A^* strategy allows the pruning of several unpromising paths and avoids the generations of such paths from the disk. This is directly reflected on running times, where we notice an average 50% improvement. When there exist rules that cover many examples from the generation set (e.g., *successor* (P), *founder* (P)), the algorithm is capable of identifying such rules rather early, thus pruning several unpromising paths that cover a subset examples from the generation set. In such cases the running time improvement is above 70%.

6. RELATED WORK

Our work is inspired by Inductive Logic Programming, but uses techniques from dependencies discovery in relational databases, KBs construction, and mining graph patterns. We review a few of the most relevant works below.

Relational Database Constraints. A significant body of work has addressed the problem of discovering constraints over relational database sources. Functional Dependencies (FDs), a formalism to express relations and dependencies among attributes, has been studied in constraints discovery literature for more than 20 years [4], with a recent focus on performance [3]. FDs can be grouped into two strands: the schema-level approaches [26] (similar to our positive rules discovery), and the instance-driven approaches [38]. More recently, Conditional FDs extend standard FDs by enforce-

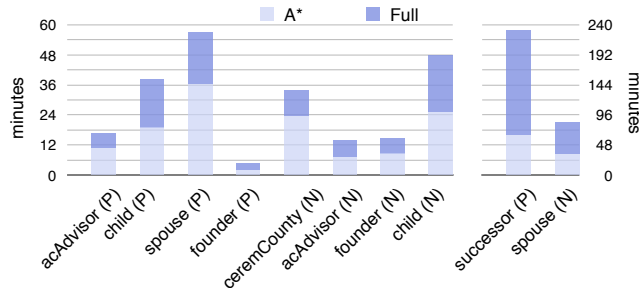


Figure 7: A^* Pruning Runtime Improvement

ing patterns of semantically related constants [21]. In the context of inconsistencies discovery for relational data, Denial Constraints (DCs) are the current state-of-the-art techniques [12]. DCs are a universally quantified first order logic formalism to express constraints over relational data, and they are directly related to our negative output rules. Efficient DCs algorithms have been proposed for data cleaning and consistent query answering [7, 13].

Despite being directly related to our output and expressing a richer language, FDs and DCs cannot be applied to RDF databases for three main reasons: (i) the schema-less nature of RDF data and the closed world assumption which no longer holds on RDF KBs; (ii) FDs and DCs techniques rely on the assumption that data is either clean or has a negligible amount of errors; (iii) scalability issues on large RDF dataset: applying relational database techniques on RDF KBs would imply to materialise all possible predicates combinations into relational tables. Some FDs approaches focus on dataset with erroneous data [1, 27], however they are still inapplicable to RDF data due to scalability problems.

KBs Rule Mining. Recently, the focus for constraints and rules discovery is moving towards RDF databases. The closest works to ours are AMIE [23] and OP algorithm (TODO: cite Ontological Path Finding), which discover positive Horn Rules from RDF KBs with same language biases. They both uses the same discovery algorithm: it first loads the entire KB into memory and then, working one predicate at time, tries to expand rules by connecting a predicate to others that share common variables. Rules are ranked according to a confidence measure that leverages on KBs partial closed world assumption. Our graph generation and navigation technique is similar to their approach, however our examples generation allows us to discover rules on just a small fraction of the KB. This is beneficial not only from a scalability point of view (see Section 5.2), but also from the language perspective: neither of the two approaches can afford smart literals comparisons. We have not tested our running time against (TODO: cite Ontological Path Finding) because we could not find an implementation of it, however it requires a powerful cluster of several machines to split the KB into multiple nodes. We showed in the experimental section how RuDiK outperforms AMIE both in final accuracy and running time.

[2] is an instance-level approach that discovers new facts for specific entities rather than generic variables. We showed in the experimental section that RuDiK is capable of generating more facts with a better precision with just a single rule. (TODO: cite Sigmod data lakes) is a modern system to discover Conditional Denial Constraints (CDCs) from RDF Data. Differently from other systems, it includes literals in its language. CDCs can be directly mapped to our negative rules, however there is not a general correlation between CDCs and positive rules. Another major difference of our setting is the hardware: our disk-based approach is designed to handle large KBs with limited memory resources, while (TODO: cite Sigmod Lakes) works on a distributed environment with a total of 832 GB RAM memory.

To the best of our knowledge, RuDiK is the first approach that is generic enough to use the same algorithm to discover both positive and negative rules in RDF KBs.

Inductive Logic Programming. Inductive Logic Programming (ILP) is a sub-field of Machine Learning and Logic Programming which investigates the inductive con-

struction of first-order Horn Rules from examples and background knowledge, usually expressed through logic formalisms [30]. RuDiK can be seen as an ILP system where the KB is the background knowledge, and the generation and validation sets correspond to positive and negative training examples.

WARMR is an ILP system that discovers frequent patterns (expressed through DATALOG queries) that succeed with respect to a sufficient number of examples [15]. When translated to databases, such patterns correspond to conjunctive queries. ALEPH⁹ is an available ILP system that is based on Prolog Inverse Entailment [29]. ALEPH works iteratively, by selecting examples from the background knowledge. It first constructs the most specific clause that entails the example selected (*bottom clause*), and then searches for some subset of the literals in the bottom clause that has the *best score* in order to define more general rules. The system allows the user to choose among several scoring functions. ILP systems such as WARMR and ALEPH are designed to work with a closed world assumption, and always require the definition of positive and negative examples. AMIE clearly outperforms these two systems [23], showing evident scalability issues when dealing medium-size KBs. Moreover, one of the main limitations of classic ILP systems is the assumption of having high-quality, errors-free training examples. We showed how this assumption does not longer hold on incomplete and erroneous KBs.

Sherlock [32] is an interesting ILP system that learns first-order Horn Rules from Web text. One of the key advantage of Sherlock is being unsupervised: it does not require negative training examples. It uses statistical significance and statistical relevance in order to discover rules that exceeds a given threshold. Differently from our setting, Sherlock is specifically designed to learn Horn Rules from open domain facts that are extracted from the Web. An interesting future direction is to adapt RuDiK to discover rules from relations extracted from free text rather than on a well-defined KB.

ILP systems take inspiration from Association Rule Mining [5], where given a database of costumer transactions the goal is to discover all frequent itemsets, i.e., all combinations of items that are found together with some minimum confidence. A well known example in a supermarket database could state that 90% of transactions that purchase bread and butter also purchase milk. As previously mentioned, adapting such a relational database setting to KBs would require the materialisation of all possible predicates combinations into relational tables.

Eventually, most of the ILP and KBs rules discovery systems rank rules according to a support value, and output only those rules that exceed a given threshold. We showed in Section 5.2 that properly setting such thresholds is not trivial, as often good rules are preceded in rank by meaningless ones. RuDiK does not use any threshold and outputs rules only when coverages over generation and validation sets are considered acceptable.

Relation to other areas. Our examples generation strategy leverages on the Local Closed World Assumption (LCWA). When dealing with incomplete KBs, the LCWA is a popular technique that replaces the canonical Closed World Assumption of standard relational databases. The

LCWA has been used in Google Knowledge Vault to estimate the quality of extracted triples [17, 18], AMIE [23] uses the LCWA to penalise discovered rules, and sometimes the LCWA is used to evaluate the quality of a target KB [19]. We see our examples generation strategy as complementary to our approach. It is possible to run RuDiK with any input examples, no matter how such examples have been generated.

Our graph-based rules discovery approach is close in spirit to mining graph patterns [20, 40], where given a (big) input graph the goal is to discover the most frequent patterns (subgraphs) according to some scoring functions. Our setting presents a key difference: we primarily look at edge labels and we are not interested in node labels, since nodes are mapped to variables when translating subgraphs to Horn Rules. Moreover, given the portion of the graph between two entities x and y , we are interested in discovery *all* possible subgraphs between x and y , which makes the problem easily solvable with BFS-like techniques.

Another interesting setting is the one of SpiderMine [39]. SpiderMine looks for the top- K largest patterns in a graph, where each node in the pattern is at most at distance r from a head vertex u . In our setting we do not have a single head vertex, but rather many vertexes (starting nodes) from which we begin the path computation. Furthermore our goal is not to find graph patterns (subgraphs), but we are simply interested in all possible paths between a pair of vertexes.

Fan et. al. (TODO: cite "Wenfei Fan. Functional Dependencies for Graphs") laid the theoretical foundations of Functional Dependencies on Graphs (GFDs). They also propose parallel algorithms for GFDs computation and evaluate the accuracy on YAGO and DBPEDIA. Despite existing a natural correlation between FDs and Horn Rules, the language they propose covers only a portion of our negative rules to detect inconsistencies in graph databases. Moreover their language does not include smart literals comparisons, shown to be vital when detecting errors in KBs.

7. CONCLUSION

Extend rules with temporal and local information.

Combine positive and negative rules: if a positive and negative rule identifies same triples, then one of them must be wrong.

Introduce functions and smarter literals comparison: if two people are not born in the same century, then they cannot be married.

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⁹<http://www.cs.ox.ac.uk/activities/machinelearning/Aleph/aleph>

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