

POLITECNICO DI TORINO

DEPARTMENT OF CONTROL AND COMPUTER ENGINEERING

Master of Science in Computer Engineering

Master Degree Thesis

# Deep Learning on Academic Knowledge Graphs

Predicting new facts in a novel semantic graph built on top of the Politecnico di Torino scholarly data



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ACADEMIC YEAR 2018-2019



*To Monia*

*To my Grandfather*



# Abstract

The publication and sharing of new research results is one of the main goal of an academic institution. In recent years, many efforts have been made to collect and organize the scientific knowledge through new comprehensive data repositories. Knowledge graphs are a particular class of graphs that are used to semantically describe the human knowledge in a specific domain by linking semantic entities through labeled and directed edges. In this work we are going to present a novel semantic graph built on top of the scholarly data produced by the researchers of the Politecnico di Torino.

Such graph, built by leveraging Semantic Web technologies, connects publications, researchers, fields of study and scientific journals in order to build a knowledge base that describes the Politecnico di Torino academic community. Once the knowledge base is built, advanced statistical relational learning techniques specifically developed to learn from graph data can be used to predict new facts. The prediction of non-existent links between entities is one of the most challenging tasks in this field, mainly because, in order to obtain meaningful predictions, the vector representations of the entities must embed their semantic characteristics.

To accomplish this goal, we decided to employ Deep Learning techniques derived from the image recognition field and specifically adapted to the task of representation learning for graph data. Such architectures allows to obtain representations that are directly learnt from the graph structure itself, without the use of any prior information. We then used such predicted facts to complete the knowledge base and build a recommendation system to empower researchers with useful insights. For instance, unexplored fields of study in which they might be interested in, or recommendations about other researchers with whom they have never worked before, but that may share the same research interests.

# Acknowledgements

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# Chapter 1

## Introduction

### 1.1 Motivation

Graphs are used to empower some of the most complex IT services available today, an example among all is the Google search engine <sup>1</sup>. Graphs can be used to represent almost any kind of information, and they are particularly capable of representing the structure of complex systems and describe the relationships between their elements.

Over the last decade, much effort has been put in trying to leverage the power of graphs to represent human knowledge and to build search tools capable of querying and understanding the semantic relations within them. RDF<sup>2</sup> graphs are a particular class of graphs that can be used to build knowledge bases. Ontologies are used to shape such knowledge bases, in order to have a semantically coherent representation of the domain knowledge. Given a domain and an ontology, RDF graphs allows to build a structured representation of the knowledge in such domain.

Modern machine learning techniques can be used to mine latent information from such graphs. One of the main challenges in this field is how to learn meaningful representations of entities and relations that embed the underlying knowledge. Such representations can then be used to evaluate new links within the graph or to classify unseen nodes. Deep learning

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<sup>1</sup><https://blog.google/products/search/introducing-knowledge-graph-things-not/>

<sup>2</sup>The Resource Description Framework will be introduced in section 2.1.2

techniques have proved to be first class citizens when dealing with representation learning tasks, being able to learn latent representations without any prior knowledge other than the graph structure.

## 1.2 Goal and contribution

Knowledge sharing is one of the main goals of research organizations. In particular, universities are among the most interested in making publicly available their research results. Today most universities have embraced the Open Science movement, making their scientific publications publicly available through web portals. An example is IRIS<sup>3</sup>, which stores all the scientific papers produced by the Politecnico di Torino, publicly sharing them through Open Access. IRIS allows to explore the published papers by searching for a field of study, matching it with the keywords inserted by the authors of the publications. This implementation has some limitations: being inserted by the authors, there could be more keywords that refers to the same scientific topic. Moreover, the author may have inserted acronyms or misspelled some words. The consequence is that the search engine of IRIS is unable to correctly retrieve all the publications about a specific research topic, because the system cannot match the searched field of study with an unambiguous *semantic entity*, but only with character strings that are not uniquely identified or semantically linked each other, and also prone to lexical errors.

Our goal is to overcome such limitations and enabling new possibilities for exploring and obtaining insights about the scientific community of the Politecnico di Torino. A new semantic-empowered search engine can be one of the possible solutions to obtain this results, allowing for coherent and precise results to be retrieved. At the foundations of this new semantic search engine there must be a data structure capable of representing semantic relations and concepts. Once such knowledge base of the scholarly data is obtained, it can be enhanced and completed by automatically extracting latent information through the use of advanced machine learning algorithms.

In the next chapters we are going to present a newly built structured and semantically coherent representation of the scholarly data produced

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<sup>3</sup><https://iris.polito.it/>

by the Politecnico di Torino, and how implicit facts can be automatically extracted from such knowledge repository by leveraging knowledge base completion techniques, implemented by means of an advanced deep learning algorithm.

## **1.3 Thesis structure**

### **1.3.1 Chapter 2**

### **1.3.2 Chapter 3**

### **1.3.3 Chapter 4**



# Chapter 2

## Background

### 2.1 Semantic Web

#### 2.1.1 From a Web of Contents to a Web of Data

The World Wide Web has been developed as a tool to easily access documents and to navigate through them by following hyperlinks. This simple description already resembles the structure of a graph: we can think of documents as nodes and hyperlinks as edges. The unstoppable growth of the *Web graph* led to the raise of new tools to explore such complexity. Search engines have been developed to easily navigate such a giant graph. First approaches were based on analytics evaluations, such as the number of times a document has been linked, as in the case of the PageRank [?] algorithm developed by Google.

The Web rapidly became one of the most innovative technology ever built, allowing to retrieve information quickly and easily as never before. The next evolutionary step has been to think about a Web not only exploitable by human beings but also by machines. In order to build such a comprehensive system, where information can be not only machine-readable, but machine-understandable, the World Wide Web had to move from a web of content, to a web of data.

The World Wide Web Consortium (W3C) introduced the Semantic Web as an extension to the prior standard of the WWW. Its primary goal has been to define a framework to describe and query semantic information

contained in the documents available on the Web, so as to allow machines to understand the semantic information contained in web pages. In the vision of Tim Berners-Lee, the father of WWW, this would bring to the transition from a World Wide Web to a Giant Global Graph <sup>1</sup>, where a web page contains metadata that provides to a machine the needed information to understand the concepts and meanings expressed in it.

### 2.1.2 The Semantic Web building blocks

The three key components of the Semantic Web standard are:

1. OWL: the Web Ontology Language [?]
2. RDF: the Resource Description Framework [?]
3. SPARQL: The SPARQL Protocol and RDF Query Language

OWL is a language used to define ontologies. In this context, an ontology is defined as a collection of concepts, relations and constraints between these concepts that describes an area of interest or a domain. OWL allows to classify things in terms of their meaning by describing their belonging to classes and subclasses defined by the ontology: if a thing is defined as member of a class, this means that it shares the same semantic meaning as all the other members of such class. The result of such classification is a taxonomy that defines a hierarchy of how things are semantically inter-related in the domain under analysis. The instances of OWL classes are called individuals, and can be related with other individuals or classes by means of properties. Each individual can be characterized with additional information using literals, that represent data values like strings, dates or integers.

The Resource Description Framework defines a standard model for the description, modelling and interchange of resources on the Web.

The first component of the framework is the *RDF Model and Syntax*, which defines a data model that describes how the RDF resources should be represented. The basic model consist of only three object types: resource, property, and statement. A resource is uniquely identified by an

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<sup>1</sup><https://web.archive.org/web/20160713021037/http://dig.csail.mit.edu/breadcrumbs/node/215>



Uniform Resource Identifier (URI). A property can be both a resource attribute or a relation between resources. A statement describes a resource property, and is defined as a triple between a subject (the resource), a predicate (the property) and an object (a literal or another resource).

The second component of the framework is the *RDF Schema* (RDFS), that defines a basic vocabulary for describing RDF resources and the relationships between them. Many of the vocabularies and ontologies available today are built on top of RDFS, such as the Friend of a Friend (FOAF) ontology [?], for describing social networks, or the one maintained by the Dublin Core Metadata Initiative [?], that defines common terms and relations used in the definition of metadata for digital resources.

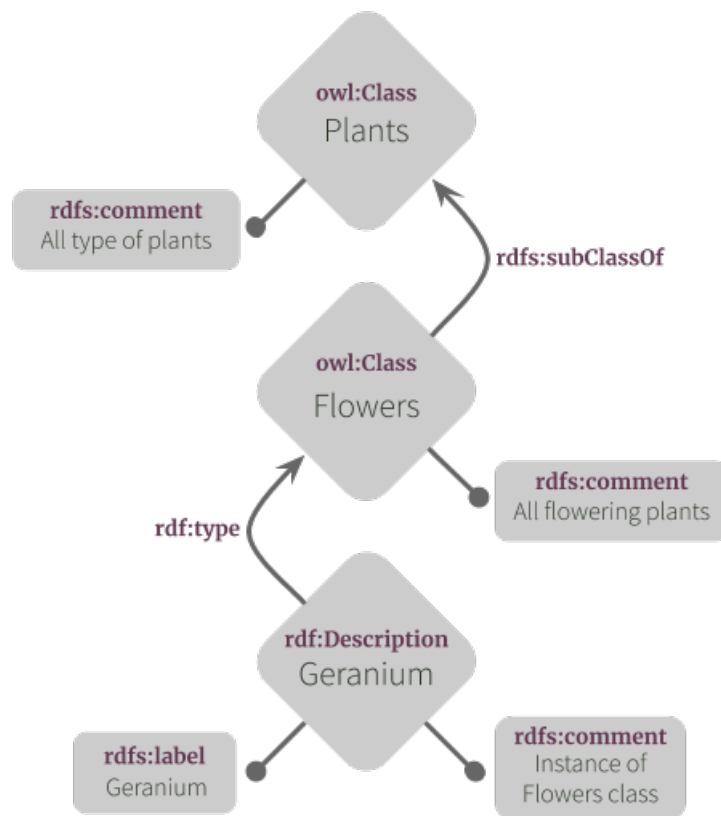


Figure 2.1: An example of ontology defined using OWL and RDF Schema.

SPARQL is a query language for triplestores, a class of Database Management Systems (DBMS) specialized in storing RDF databases. Such DBMS often expose endpoints that can be used to query the database

and obtain results. Given the complexity of the stored data, the query language has been designed to be as simple as possible, for instance by allowing the use of variables, whose definition is preceded by a question mark.

The syntax of SPARQL is heavily derived from SQL, with some minor adaptations to be more suited for querying graph data. The following is an example of query which selects all the labels (human-readable description of a resource) of all the entities that match the given resource type.

```
PREFIX plants:<http://example.org/plants/>

SELECT ?name
WHERE {
    ?subject rdf:type plants:flowers .
    ?subject rdfs:label ?name .
}
```

### 2.1.3 Knowledge Bases as knowledge repositories

Even though the raise of the Semantic Web has suffered a slowdown in its growth due to the complexity of its vision, many new projects were born from its enabling technologies. Efforts have been put by profit and non-profit organizations in trying to build complex knowledge repositories starting from the knowledge already available in the Web. An example among all is the DBpedia<sup>2</sup> project, which developed a structured knowledge base from the semi-structured data available on Wikipedia. Another example is the *Google Knowledge Graph*<sup>3</sup>, which is used to enhance the Google search engine and virtual assistant capabilities, allowing to retrieve punctual information about everything that has been classified in its ontology and described in its knowledge base, or the *Open Academic Graph*<sup>4</sup>, a Scientific Knowledge Graph that collects more than three hundred million academic papers. A comparison between some of the biggest

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<sup>2</sup><https://wiki.dbpedia.org/>

<sup>3</sup><https://blog.google/products/search/introducing-knowledge-graph-things-not/>

<sup>4</sup><https://www.openacademic.ai/oag/>

Table 2.1: Comparison of some of the biggest industry-scale knowledge graphs developed to this date. Table adapted from [?].

	Data model	Graph size	Development stage
<b>Microsoft</b>	Entities, relations and attributes defined in an ontology.	2 billion entities and 55 billion facts.	Actively used in products.
<b>Google</b>	Strongly typed entities, relations with domain and range inference.	1 billion entities, 70 billion assertions.	Actively used in products.
<b>Facebook</b>	All of the attributes and relations are structured and strongly typed.	50 million primary entities, 500 million assertions.	Actively used in products.
<b>eBay</b>	Entities and relation, well-structured and strongly typed.	100 million products, more than 1 billion triples.	Early stages of development.
<b>IBM</b>	Entities and relations with evidence information associated with them.	Various sizes. Proven on scales documents >100 million, relationships >5 billion, entities >100 million.	Actively used in products and by clients.

knowledge graphs developed to this date is available in Table 2.1.

From an implementation perspective, knowledge bases can be created to describe a specific domain by defining an ontology and a vocabulary for such domain using OWL and RDF Schema, and then by describing the concepts of such domain using the RDF Model and Syntax. The RDF document obtained can then be stored in a triplestore and queried using SPARQL. The main effort in building knowledge bases is to have a correct understanding and prior knowledge of the domain of interest, to avoid the risk of mischaracterizing and misrepresenting concepts.

If all the requirements and cautions are met, a well formed knowledge base may prove to be a critical resource for an organization. It permits to build new services upon it, and also to improve the existing knowledge inside the organization by performing reasoning upon the available knowledge, in order to derive implicit facts starting from the existing entities and relationships. Another field of applications is the development of Expert

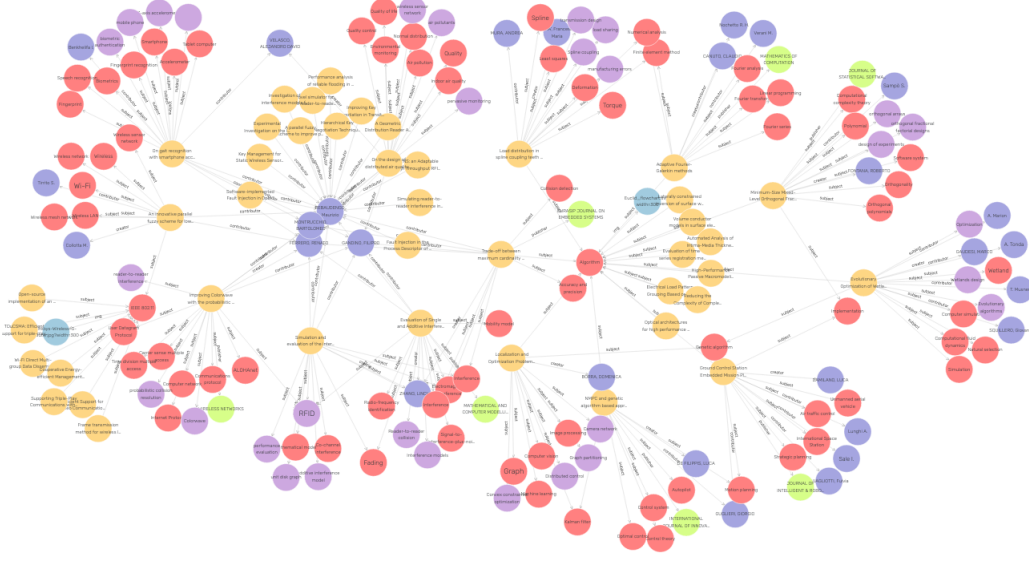


Figure 2.2: An extract of the Polito Knowledge Graph whose details will be described in section 5.1.

Systems<sup>5</sup>, AI software that emulates the behavior of a human decision-making process by navigating the knowledge base and taking decisions like in a rule-based system.

Today’s knowledge bases are commonly composed of tens of thousands nodes and by hundreds of thousands of edges. Considering such dimensions, storing and querying giant graphs requires the adoption of specialized DBMS that are capable of efficiently store and query the RDF input representation. Moreover, performing analysis and gathering statistics from such giant graphs requires the adoption of highly efficient algorithms in order to retrieve the desired output in an acceptable time.

The availability of such a complex and informative data structure leads to the opening of interesting scenarios, especially when thinking about the latent information that can be extracted from it. In fact, a knowledge base is a structured representation of the human knowledge in a specific field, thus its comprehensiveness is restricted by the human understanding.

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<sup>5</sup>[https://en.wikipedia.org/wiki/Expert\\_system](https://en.wikipedia.org/wiki/Expert_system)

## 2.2 Learning on Graphs

### 2.2.1 Representation learning

Machine learning (ML) algorithms are used to learn models from the available data, with the final goal to obtain a set of parameters that are fine-tuned to identify seen characteristics in the data used for training. The obtained models can be used to recognize unseen inputs by leveraging the knowledge embedded in such parameters. ML algorithms require the input data to be available in a machine-understandable vector representation. An important task in the ML field is the learning of such representations, task known as representation learning.

Natural Language Processing (NLP) is one of the research branches that in the past years has made a great use of machine learning algorithms both for language recognition and for embedding words *meaning* into words *vectors*. One of the most successful algorithms when dealing with representation learning of words is Word2Vec [?], where the model obtained is trained to learn a vector representation for each word in a vocabulary. In Word2Vec, the concept of meaning of a word is related to the context in which such word is frequently used, so two words are recognized as similar if they're used in similar contexts, thus in the vector space of the learnt representations words that have similar meaning have higher cosine similarity<sup>6</sup> with respect to dissimilar ones. For instance, the cosine similarity between the word vectors of "Man" and "King" is roughly the same as the one between the words "Woman" and "Queen", since such words are used in similar contexts. This has opened up new scenarios for language recognition and processing, since it allowed to perform vector operations on such words which brought interesting results, as can be seen in Figure 2.3.

This idea of words characterized by the context in which they're used can be generalized and applied to other fields of research, such as the field of representation learning on graphs.

Graphs are composed of nodes and edges, and are used to describe

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<sup>6</sup> Cosine similarity is a heuristic method to measure the similarity between two vectors by computing the cosine of the angle between them:  $similarity(A, B) = \cos(\theta) = \frac{A \cdot B}{\|A\| \|B\|}$

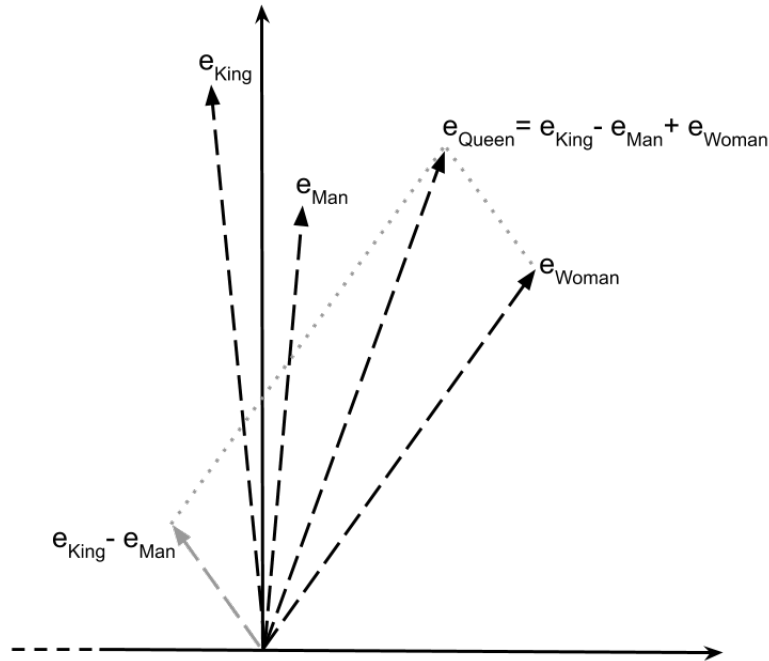


Figure 2.3: Word vectors allows to perform vector operations, the results obtained reflect the fact that Word2Vec is capable of embed the meaning of such words.

complex systems, such as social networks or the interactions in a molecular biology system. To apply machine learning algorithms to such data structures vector representations of nodes and edges are needed in order to be able to learn from the available data and predict new facts. Such vector representations are often referred to as *embeddings* because they should embed the characteristics of the graph nodes, so that similar nodes have similar embeddings. In example, in a scholarly knowledge base publications with same authors and similar subjects should have similar embeddings.

Early approaches required these representations to be learned from feature vectors that where handcrafted, task that required not only a relevant amount of effort, but also a deep understanding of the domain of interest. This has long been one of the main obstacles when dealing with representation learning tasks, since who has knowledge of the domain and who has to engineer the features were unlikely the same individual.

### 2.2.2 Deep Learning on graphs

In the latest years a big shift towards deep architectures has been made in machine learning, mainly thanks to the development of highly parallelized architectures that are able to efficiently compute at the hardware level vector and matrix multiplications, operations that are at the basis of any machine learning task. Deep Learning (DL) algorithms are able to extract relevant features from raw data by applying simple mathematical operations, such as convolution, to the input data. An example of one of the most successful applications of DL is in image recognition, where matrix representations of images are convolved with self-trained filters that are able to extract the relevant features needed to recognize patterns present in the input images.

Deep learning techniques have proven to perform well also in the field of representation learning for graph data. As can be seen in figure 2.4, a digital image is composed of pixels which can be thought of as nodes in a graph, where each pixel is connected by an edge to its immediate neighbors. This suggests that the techniques used when dealing with images can be adapted, with some major changes, to the field of representation learning on graphs, but also in other fields of research, such as learning on manifolds.

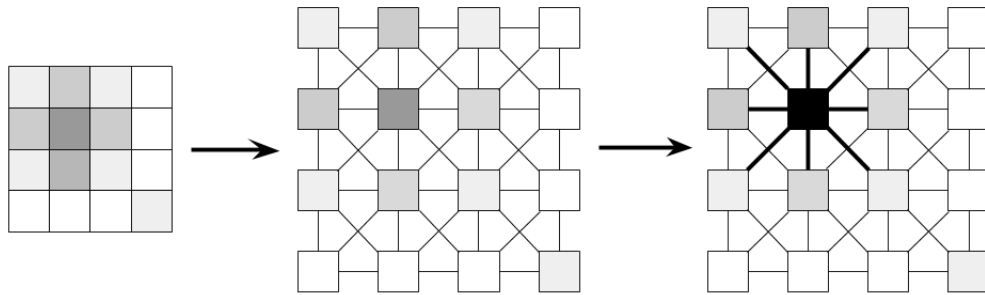


Figure 2.4: A digital image can be thought of as a graph.

One of the issues when working with graph data is that commonly graphs are built to describe complex systems, such as the knowledge of a domain or field for knowledge graphs, and thus are composed of a fairly high amount of nodes and edges. The matrices used to store the graph structure can thus explode in dimensionality, becoming impractical as input data. Moreover, graphs are not regular structures with a given

shape and size, such a matrix of pixels for images, but they live in an irregular domain which led to highly irregular structures. The first issue can be solved by randomly sampling the graph at each training epoch, the immediate drawback being that more than one epoch is required to train over all graph nodes. The second issue can instead be solved by adapting known algorithms to work on irregular domains. One of the possible approaches, which has proven to work well, is the one based on convolutions.

Graph Convolutional Networks (GCNs) [?] are a class of semi-supervised deep learning algorithms for graphs which are based on the same convolution and backpropagation operations as the well known Convolutional Neural Networks [?] (CNNs) used for feature learning on images. The main difference between CNNs and GCNs is in how the convolution is performed, instead the backpropagation phase is the same as the one used to update the parameters of CNNs, with a task-specific loss function. In a CNN the input matrix of each network layer, which is the pixel matrix of the input image for the first layer, is convolved with a convolutional filter, whose parameters are then updated during the backpropagation phase.

GCNs works similarly by convolving at the  $l$ -th layer of the network the feature vector of each node with the feature vectors of its  $l$ -nearest neighbors by means of a convolutional filter. This operation is done by applying the following transformation:

$$H^{(l+1)} = \sigma(\tilde{D}^{-1/2} \tilde{A} \tilde{D}^{-1/2} H^{(l)} W^{(l)}) \quad (2.1)$$

Where  $H^{(l)} \in \mathbb{R}^{N \times d^{(l)}}$  is the nodes hidden representation matrix, which is the output of the previous layer, with  $N$  being the number of nodes in the graph and  $d^{(l)}$  being the dimensionality of the current layer. For the first layer,  $H^0$  is equal to the input feature matrix  $X$ , where each row can be initialized with the respective node feature or with a one-hot encoded vector. For the last layer,  $H^{(l+1)}$  is the embeddings matrix.  $\tilde{A}$  is an adjacency matrix of the graph that includes self loops.  $\tilde{D}$  is the node degree matrix of  $\tilde{A}$  and is used to normalize it.  $W^l \in \mathbb{R}^{d^{(l)} \times d^{(l+1)}}$  is the convolutional filter that is shared among all nodes and is unique for each layer, just like the convolutional filter of a CNN layer. The shape of this filter will directly impact the dimensionality of the embeddings obtained, in fact  $H^{l+1}$  is shaped as a  $N \times d^{(l+1)}$  matrix. Finally,  $\sigma$  is a non linear activation function, for example *ReLU*.



Looking at the update rule of a single node embedding will make more clear how the convolution is actually performed. The forward rule to update the embedding of a single node at the  $l$ -th layer of the network is the following:

$$h_i^{(l+1)} = \sigma\left(\sum_{j \in \eta_i} \frac{1}{c_{i,j}} h_j^{(l)} W^{(l)}\right) \quad (2.2)$$

Where  $\eta_i$  is the set of neighbors of node  $i$ , which contains the node itself because of the added self loop, and  $c_{ij}$  is a normalization constant obtained from the multiplication between the adjacency and degree matrices. The updated feature vector  $h_i^{(l+1)}$  of the node  $i$  is obtained by performing the following operations:

1. The feature vectors  $h_j^{(l)}$  of its neighbors are transformed by the matrix multiplication with the layer filter  $W^{(l)}$ .
2. The resulting  $1 \times d^{(l+1)}$  shaped features vectors are multiplied with the respective normalization constants and summed together.
3. The non-linear function is applied to the result of the summation, obtaining the updated feature vector of the node  $i$ .

As a consequence of applying such transformation to all nodes, at the  $k$ -th layer a node is represented by its transformed feature vector, which embeds the structural information within the node's  $k$ -hop neighborhood. So the amount of layers of the network is an hyperparameter that controls how much information from furthest nodes has to be collected in each node embedding. This represents a fundamental architectural difference between CNNs and GCNs: while the former are commonly built by stacking a lot of layers, the latter relies on architectures that are more wider, due to the dimensionality of the graphs involved, and that consist of a fairly low amount of layers, in order to characterize the nodes only by their immediate surroundings.

As a result of the propagation step each node embedding will be characterized by its context, just like it happens in Word2Vec, but in a non-Euclidean domain. So for example, in a social graph where each person is characterized by its friends, interests, places visited and so on, two people will have similar embeddings if they are both linked to the nodes that represent such characteristics.

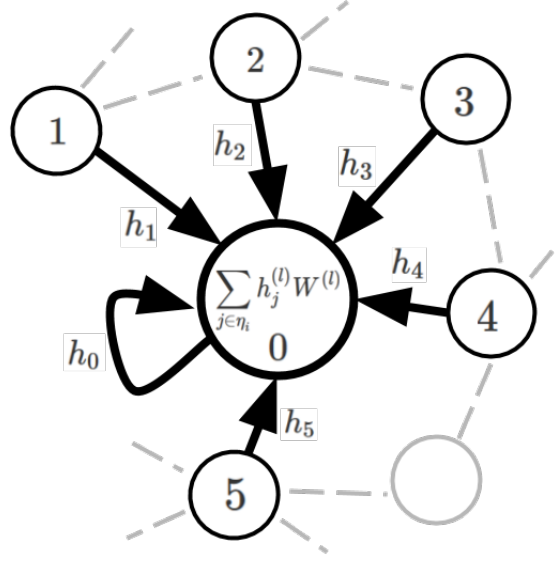


Figure 2.5: Representation of a GCN updating the feature vector of the node  $0$  by summing the convolved features of its adjacent nodes.

The node embeddings obtained by applying a GCN or one of its variants can then be used to perform some learning task on the graph, two examples are the classification of unseen nodes or the link prediction of non-existent edges. The latter is one of the most interesting tasks, since it allows to complete the information contained in the graph by predicting unseen facts.

### 2.2.3 Link prediction on knowledge graphs

Predicting new facts is one of the most common task in the field of knowledge graph completion. The goal of such task is to predict new, unseen triples that correspond to missing facts in the knowledge base, and that can be later added to the graph. Deep learning techniques for link prediction are based on the following two main steps:

1. Train an *encoder* model that is able to embed the node features and produce meaningful embeddings.
2. Apply a factorization model that act as a *decoder*, which is used to score the unseen triples under evaluation.

Deep learning techniques such as GCN can be exploited to obtain meaningful node embeddings, but fall short when dealing with graphs where nodes are connected by different relations (multi-relational graphs). In fact, if we use a single layer GCN to obtain the embeddings of two nodes that share the same neighborhood, but are connected to the neighbors via different relations, we'll obtain almost the same embeddings, even if it's clear that they do not share the same characteristics. For example in a producer-consumer framework one node could be the producer while the other the consumer, having both a common neighborhood of nodes which are produced by the former and consumed by the latter. If a GCN is used, the embeddings obtained would be very similar, even if the two nodes clearly don't have the same role and don't belong to the same class.

To overcome this limitation, changes to the basic GCN architecture have been proposed, so to obtain models that works well when applied to multi-relational graphs.

The Relational Graph Convolutional Network [?] (R-GCN) is an extension of GCNs which is focused on modeling multi-relational graphs composed by labeled and directed edges, and thus is particularly capable of embedding both nodes and relations of a knowledge graph. R-GCN can be used for both link prediction and node classification tasks.

At an high level the R-GCN architecture can be seen as a special case of the *message passing framework* [?], which groups together under a common scheme most of the existing neural models for graph data. The framework defines two major phases: a per-edge message computation and a per-node message reduction. In the first phase a function or linear transformation is applied to each edge to obtain an edge-specific message. Then, in the reduce phase, the embedding of each node is updated by aggregating together all the messages of the incoming edges.

This two phases can be grouped together through by the following equation:

$$h_i^{(l+1)} = \sigma \left( \sum_{m \in \mathcal{M}_i} g_m(h_i^{(l)}, h_j^{(l)}) \right) \quad (2.3)$$

Where  $\mathcal{M}_i$  is the set of incoming messages for node  $i$  and  $g_m$  is a message specific transformation.

The idea behind R-GCN is to have different set of parameters for different relations. At each step inside the network, the feature vector of a node is updated by convolving its first neighbors features with a convolutional filter that is different based on the kind of relation that connects the nodes. The forward rule to update the embedding of a node at the  $l$ -th layer is the following:

$$h_i^{(l+1)} = \sigma \left( W_0^{(l)} h_i^{(l)} + \sum_{r \in \mathcal{R}} \sum_{j \in \mathcal{N}_i^r} \frac{1}{c_{i,r}} W_r^{(l)} h_j^{(l)} \right) \quad (2.4)$$

Where  $h_i^{(l)}$  is the embedding (or, for the first layer, the input feature vector) of the node  $i$ ,  $W_0^{(l)}$  is the learnt kernel for the self loop relation,  $\mathcal{N}_i^r$  is the set of indices of the neighbors of node  $i$  under the relation  $r \in \mathcal{R}$ , with  $\mathcal{R}$  being the set of all the relations present in the graph.  $W_r^{(l)} \in \mathbb{R}^{d^{(l+1)} \times d^{(l)}}$  is the learnt filter for the relation  $r$ . As for the GCN architecture,  $\sigma$  is a non linear activation function and  $c_{i,r}$  is a normalization constant, commonly initialized to  $|\mathcal{N}_i^r|$ .

From a message passing framework perspective, the message function (per-edge transformation) is equal to the linear transformation  $W_r h_j$ , and the reduce function (per-node transformation) is just the sum of all the messages computed for the edges connected to each node.

As can be seen the update rule looks similar to the one for GCNs (2.2), with the major difference that in the case of a R-GCN the filters used to convolve the feature vectors of neighboring nodes are relation specific, so the number of filters at each layer will be equal to the number of relations inside the graph. As a consequence, the kind of relation that connect the node to its neighbors has an important role in determining the transformed node embedding.

Some form of regularization is required in order to avoid overfitting on rare relations, thus to obtain a more generalized model, and also to avoid the rapid growth in the number of parameters of the network for highly multi-relational graphs. One of the solutions proposed by the original paper [?] is to decompose each relation filter  $W_r$  using basis decomposition:

$$W_r^{(l)} = \sum_{b=1}^B a_{r,b}^{(l)} V_b^{(l)} \quad (2.5)$$

This allows to store only the relation-specific coefficients and the basis,

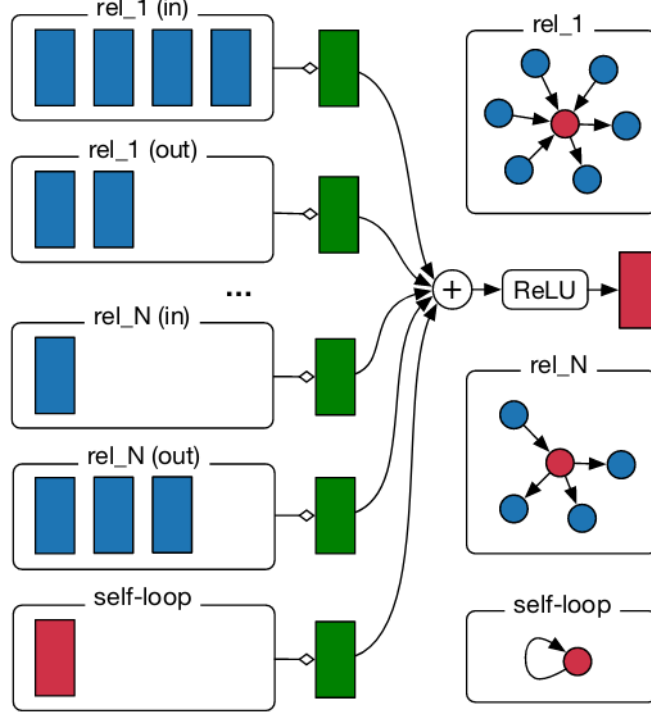


Figure 2.6: R-GCN encoder model. Image taken from [?].

which will be shared by all the relations.

The model obtained by training a R-GCN can then be used to build an encoder that given as input a graph, gives as output the embeddings of all nodes and the relations parameters. Then a factorization method can be used to evaluate unseen facts inside the graph, exploiting the resulting embeddings. Such methods are used as scoring functions in order to obtain, starting from the entities embeddings, a real value that can be used to score the unseen triples under evaluation.

DistMult [?] is one of the most common and simple factorization methods used to score unseen triples. Given a triple  $(s, r, o)$ , where  $s$  is the source node,  $o$  is the destination node, and  $r$  is the relation of the edge that links the source to the destination, DistMult allows to compute an associated real valued score as follows:

$$score_{(s,r,o)} = f(s, r, o) = e_s^T R_r e_o \quad (2.6)$$

Where  $e_s$  and  $e_o$  are the embeddings of the source and the destination node, obtained by means of an encoder model like R-GCN, and  $R_r \in \mathbb{R}^{d \times d}$  is obtained by transforming the embedding of the relation  $r$  into a diagonal matrix. Such relation embedding is not learnt by the R-GCN encoder, but is a randomly initialized vector that represent the relation.

The score obtained by applying the factorization method above can then be used to evaluate whether the triple  $(s, r, o)$  is a good candidate to be added to the graph: an high score has to be interpreted as a high confidence of the model in the fact that the triple should belong to the knowledge graph.

## Chapter 3

# Approach and methodology

This chapter introduces the architecture developed to build, enhance and visualize the Polito Knowledge Graph (PKG), an academic RDF graph built to organize in a structured and semantically coherent way the publications produced by the researchers of the Politecnico di Torino. The graph also includes publication-related entities, such as authors, journals, and fields of study.

The architecture is composed of three main modules, that at an high level can be seen as part of a producer-consumers architecture, where the former produces the graph data, and the latter consume the data produced.

The architecture, which is showed in Figure 3.1, is structured as a pipeline composed of the following three modules:

1. The *Builder*, which creates a first version of the RDF graph.
2. The *Enhancer*, which implements ML techniques to predict unseen facts. Such facts can then be added to the graph to improve its completeness.
3. The *Viewer*, a web application that allows to query and visualize the graph data.

The *Builder* act as producer taking as input the IRIS data and producing a set of RDF statements that together composes the Polito Knowledge Graph. The *Enhancer* act as both a producer and a consumer, given that

it takes as input the PKG and use it to predict unseen facts that can be later added as new RDF statements. The *Viewer* consumes the graph by storing it a triplestore and exposing a public web interface for querying and visualizing the data.

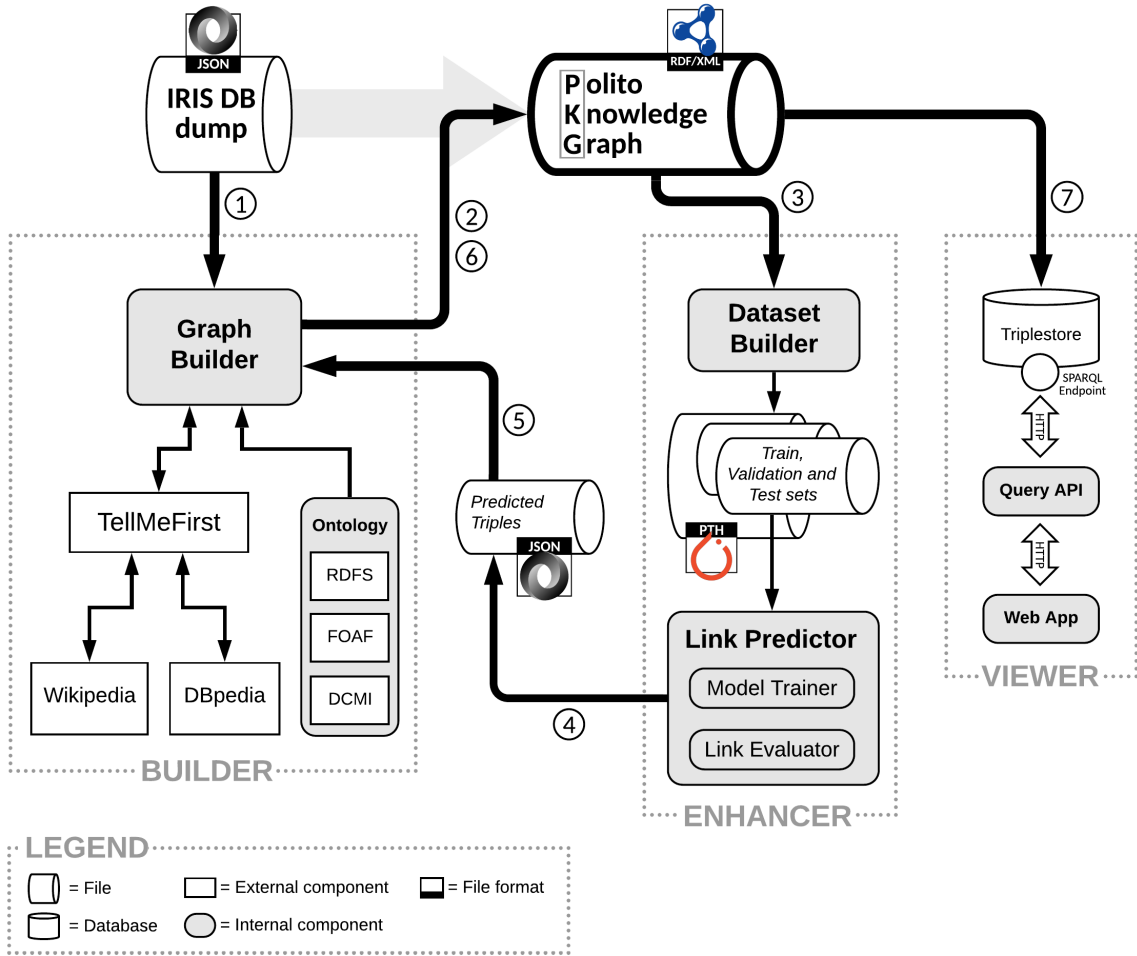


Figure 3.1: Schema of the pipelined software architecture developed to build, enhance and visualize the Polito Knowledge Graph.



## 3.1 Builder Module

The Builder input is a dump of the IRIS<sup>1</sup> database, which is the platform used by the Politecnico di Torino to store and share all the publications produced by its researchers. The dump is a JSON file that contains all the information available for the scientific paper published in a period of five years that goes from 2013 to 2017.

The Builder goal is to translate all the information contained in the dump in a set of semantically coherent RDF triples. To do so, an ontology that describes the domain of the IRIS scholarly data has been defined.

The builder uses as reference the defined ontology to analyze each record in the JSON dump, and builds facts as RDF triples by matching the information contained in the record with the concepts defined by the ontology. For example, given that a publication may have more than one contributor, the ontology that we defined differentiates the principal author from the contributors by using two different statements to tie the corresponding entities to the publication.

One of the fields of the publication record is the abstract. We wanted to link each publication to its research fields, to do so we used the abstract as input text for TellMeFirst [?], a tool for the automatic extraction of semantic keywords from texts. Such keywords, to which from now on we'll refer to as *topics*, are retrieved from the DBpedia taxonomy and are uniquely identified by their corresponding URI. Exploiting TellMeFirst we are able to automatically tie each publication to its relevant topics, by adding the corresponding RDF statements to the graph. Being the topics added as graph entities which are uniquely identified, all the publications that share a topic are linked to the same topic entity.

The result of this process is a set of RDF statements that constitutes the first version of the Polito Knowledge graph, a semantically coherent description of the publications produced by the Polito researchers, where each publication is uniquely identified and linked by means of semantic relations to the entities representing its authors, contributors and research topics.

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<sup>1</sup><https://iris.polito.it/>

## 3.2 Enhancer Module

Once the first version of the RDF graph has been built, it could be used as input for the Enhancer Module, whose main goal is to infer unseen facts to complete and enhance the information available inside the knowledge graph.

The Enhancer is composed of three main components:

1. The Dataset Builder.
2. The Model Trainer.
3. The Link Evaluator.

The Dataset Builder is in charge of translating the RDF graph into a usable dataset for the Model Trainer, given that the RDF statements cannot be used as input data for the link prediction algorithm. The dataset will then be splitted into three disjoint<sup>2</sup> sets: one for training, one for validation and one for testing.

The Model Trainer uses this three sets to train a link predictor. The training set is used to train the model at each epoch, while the validation set is used to evaluate which model parameters to keep, identifying the best epoch. The test set is used to evaluate the accuracy of the model, loaded with the best epoch parameters, upon unseen triples. Once the model has been trained, it can be used to predict unseen facts.

The Link Evaluator loads the best model found during the training phase and uses it to evaluate unseen triples. The Evaluator creates an set of unseen RDF triples and produces a score for each of them. Only the triples that obtain an high score (and so an high probability to be true facts) and are correct in domain and range with respect to the ontology are kept. The predicted triples are then saved as RDF statements and could be added to the RDF graph, obtaining the enhanced version of the Polito Knowledge Graph. This new version will contain not only the information available in IRIS, but also the latent information inferred by the learnt model, like missing topics, similar authors or proposed journals, which can be used to empower a recommendation system for the Polito researchers.

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<sup>2</sup>Two sets are said to be disjoint if they have no element in common.

## 3.3 Viewer Module

The last module of the architecture is the Viewer, which is composed of a triplestore, an API layer, and a front end web interface.

The triplestore is a specialized DBMS for storing and retrieving RDF statements. It stores an online copy of the Polito Knowledge Graph and expose a SPARQL endpoint that allows to query the graph itself.

The API layer exposes a REST API that allows to retrieve the information contained in the graph without the need of SPARQL queries. It accept HTTP requests with URL-encoded parameters and proceeds to query the SPARQL endpoint of the triplestore by matching the parameters upon some predefined queries. Then it translates the response obtained from the triplestore in a JSON file, which is sended back to the requesting client, which is typically the front end.

The front end is a responsive web application that act as the entry point for the user, mimicking the functionalities of a modern search engine. It allows to query and visualize the data contained in the graph by performing HTTP requests to the API layer and displaying the results received in a structured and comprehensible way.



# Chapter 4

## Related work

Currently there is a growing interest in scientific knowledge graphs, both by academic institutions and by private organizations. An example is the Open Academic Graph<sup>1</sup> (OAC), an academic knowledge graph built by unifying two of the largest scientific KG available, the Microsoft Academic Graph<sup>2</sup> and AMiner<sup>3</sup>. It has been publicly released to allow the study of citation networks, papers content and more. The first version of the graph, released in 2017, has been built by merging together the aforementioned graphs and by linking the matching publications, obtaining a graph that is composed of more than three hundred million publications. In the first version the only type of entity in the graph was the publication: authors, journals, and all the other publications information were added as attributes, and not as entities.

In January 2019 the second version of the OAC has been released, adding even more publications to the graph. However, the biggest change of this new version is the addition of authors and venues as graph entities, instead of being simple publications attributes.

However, the OAC does not contain the publications topics as graph entities, but as author keywords, thus being prone to the same limitations of IRIS: the keywords are the ones chosen by the authors and are not referencing to semantic concepts, being simple character strings.

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<sup>1</sup><https://www.openacademic.ai/oag/>

<sup>2</sup><https://academic.microsoft.com/>

<sup>3</sup><https://www.aminer.cn/>

Regarding the development of tools similar to the Polito Knowledge Graph by other academic institutions, an example of employment of knowledge graphs and natural language processing techniques to build new academic search engines is Wiser [?], an expertise search tool developed by the University of Pisa and publicly released at the beginning of 2019. The KG of Wiser is composed of approximately 1'500 authors, 65'000 publications and 35'000 topics<sup>4</sup>. The main function of Wiser is to allow the search of expertise in a given research field. The system has proven to be particularly effective, representing a strategic tool and being actively used by the university technological transfer office.

As saw in the previous chapter, one of the main components of the PKG pipeline is TellMeFirst, tool used to automatically extract the topics of interest from the publications abstracts. By automatically extracting the topics, TMF allows to add them as entities in the Polito Knowledge Graph, so that each publication is directly linked to its main topics, and each topic is linked to all the publications of which it is a subject (reverse relation).

Other tools that are able to extract the subjects of a publication exists, an example is the CSO Classifier [?]. This tool is able to automatically classify a research paper according to the Computer Science Ontology<sup>5</sup> (CSO), an automatically generated ontology of research topics in computer science. The fact that the CSO Classifier relies on a predefined ontology has some disadvantages with respect to TellMeFirst, the biggest being the fact that the Computer Science Ontology is restricted to the computer science field only, while TMF, using DBpedia as its source of knowledge, is able to extract topics (and so classify a research paper) regarding every field of research. However, the approach of CSO Classifier has also its advantages: being the ontology more restricted, the classification could be more accurate, and the structure of the ontology itself may be tailored for such classification task.

Regarding the learning task on graph data, many architectures that implements the Message Passing Framework [?] and that are derived from

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<sup>4</sup>Numerical statistics about the Polito Knowledge Graph are available in the following chapter.

<sup>5</sup><https://cso.kmi.open.ac.uk/home>

the classical Convolutional Neural Network are available, some example being [?] and [?]. We decided to implement the R-GCN architecture for our prediction task being specifically designed to work with relational graph data. Moreover, due to the different approach followed by the authors of R-GCN, which relies on the convolutional architecture only as an encoder, in future works we could try different factorization methods while keeping the same encoder architecture based on R-GCN to learn the node embeddings. Some possible factorization methods that we would like to implement as decoders are [?] and [?].

To the best of our knowledge we are the first to employ a Relational-GCN for the completion task of a scholarly knowledge graph.





## Chapter 5

# Development and implementation

In this chapter we will describe how we developed and implemented the architecture behind the Polito Knowledge Graph, particularly focusing on the components in charge of the graph creation and enhancement, whose architecture has been introduced in chapter 3.

We will firstly introduce a detailed description of the input data used to create the graph. We will also describe the ontology used to shape the knowledge of the scholarly domain and the graph schema obtained by structuring the information made available by IRIS with such ontology. Then, we will discuss how we implemented the Builder Module, which is responsible for the actual creation of the Polito Knowledge Graph starting from the IRIS publications metadata.

In the second part of the chapter we will focus on the implementation of the link predictor module, which is used to predict new facts inside the knowledge base. We will describe how starting from the Polito Knowledge Graph we created a usable dataset for the training of an encoder model, which architecture and hyperparameters we choose, how we trained such model and how it has been validated during the training phase in order to choose the best parameters. We will also discuss how we tested the model and which metrics we choose to evaluate the accuracy of the predictions on the test set. Finally, we will present how we employed the trained model to obtain predictions about new facts, and how the predictions obtained can be used to build a recommendation system for the Polito Knowledge Graph.

## 5.1 Building the Polito Knowledge Graph

As already mentioned when introducing the pipeline, the input data used to build the PKG is a dump of the IRIS database which contains the information about all the scientific papers published by the researchers of the Politecnico di Torino from the beginning of 2013 to the end of 2017. The dump is a JSON file composed of 23'268 records, where each record contains the metadata about a single scientific publication. Many of the metadata present are generated by the IRIS platform itself in order to manage its internal processes of acceptance and update of the publication status. We have discarded such information, since they are not significant for the characterization of the publication.

We selected the following metadata as the ones that truly characterize a scientific paper:

1. The publication identifier.
2. The title.
3. The abstract.
4. The author name, surname and identifier.
5. The contributors and coauthors names, surnames, and identifiers (if present).
6. The date of publication.
7. The journal title and ISSN (if present).
8. The keywords entered by the authors to tag the publication.

The publication identifier is a unique numeric code associated by IRIS to each scientific paper. Also the authors, contributors and coauthors should be uniquely identified by an alphanumeric identifier, however, only the researchers of the Politecnico di Torino have such identifier assigned. External researchers that may be contributors or coauthors of the publication have only their name and surname listed. The author is instead always a Polito researcher.

If the publication has been published in a journal, and its information are available, then the journal ISSN is used as identifier. If the publications is a conference paper, no information about the conference is present other than the title, thus we cannot uniquely identify the conference, at least in

this first version of the Graph Builder. The publication title, the abstract, the names, surnames and the keywords are simple character strings, while the date of publication is formatted as *dd-mm-yyyy*.

As already discussed in previous chapters, the keywords entered by the authors cannot be treated as semantic concepts. We wanted to solve this issue, in order to have all the publications that refer to a specific topic linked to a same semantically unambiguous graph entity that uniquely represents such topic. To do so, we employed TellMeFirst (TMF), a tool for the automatic extraction of semantic keywords that relies on the DBpedia ontology and knowledge graph as its source of knowledge.

In the following sections we will describe how we developed the Graph Builder and how, leveraging TMF, we have been able to extract the relevant topics of a publication from its abstract, so to add such semantic concepts as graph entities in the Polito Knowledge Graph.

### 5.1.1 PKG ontology and schema

Starting from the metadata discussed above, we defined the PKG ontology as composed of five different classes:

1. The *Publication*.
2. The *Author*.
3. The *Journal*.
4. The *AuthorKeyword*.
5. The *TMFResource*.

In order to build a knowledge graph, the instances of such classes must be linked together by means of semantic relations, called predicates. To do so, we employed some predicates already defined by the FOAF [?] and the DCMI [?] ontologies, together with some terms defined by the RDFS [?] standard. The graph structure obtained is represented by the schema in Figure 5.1, where we show the classes, the attributes of such classes and the predicates that links them together.

As can be saw from the schema showed in Figure 5.1, the *Publication* class is linked to the *Author* class by means of two relations both defined

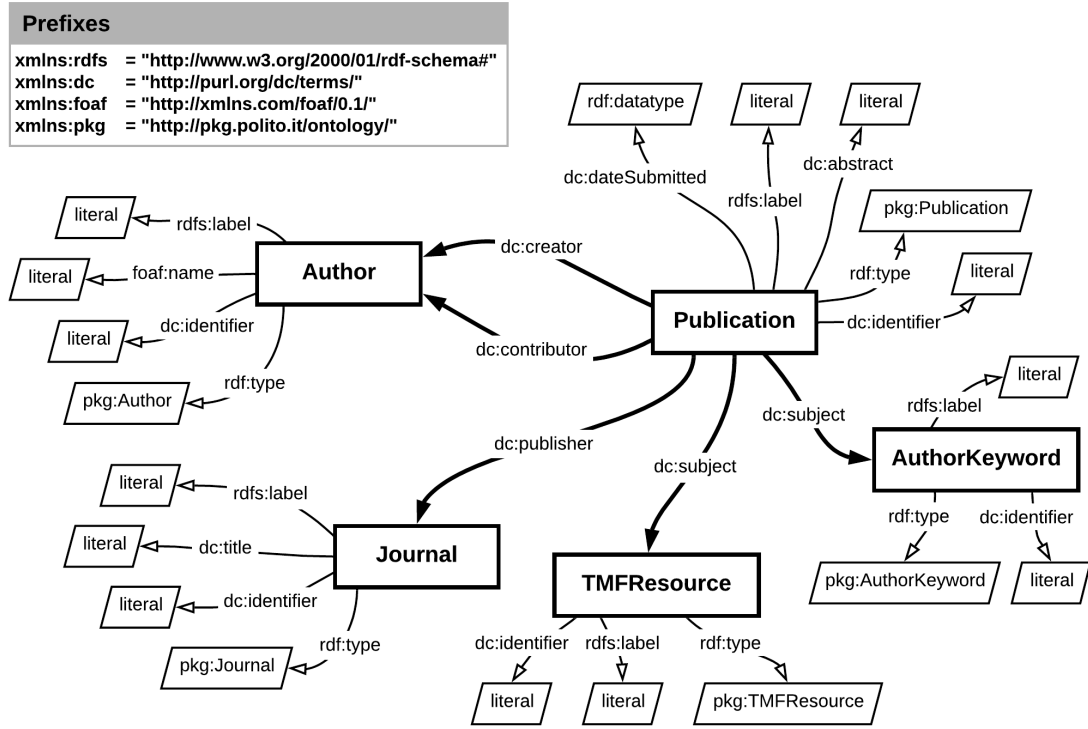


Figure 5.1: Schema of the Polito Knowledge Graph. The ontologies used to define the structure are showed in the prefixes table.

by the DCMI ontology: `dc:creator`<sup>1</sup> and `dc:contributor`<sup>1</sup>.

We decided to only differentiate the first author (*creator*) from the others (*contributors*) due to the inability to discriminate which are the co-authors and which are the collaborators based on the metadata made available by IRIS.

The *AuthorKeyword* class represents the keywords entered by the authors. Even if such keywords do not represent any semantic information, we decided to add them to the knowledge base for the sake of completeness with respect to the metadata available.

Instead, the *TMFResource* class is used to instantiate the entities corresponding to the DBpedia resources extracted by TMF starting from the publications abstracts. Being uniquely identified by their DBpedia URI, the publications that shares a topic will be linked, through the `dc:subject`

<sup>1</sup>The *dc* keyword is used as prefix for the Dublin Core Metadata Initiative (DCMI) namespace: <http://purl.org/dc/terms/>

relation, to the same instance of the *TMFResource* class that represents such topic. An example is showed in Figure 5.2, where are showed all the publications that are linked to the entity instantiated for the *Knowledge Base* topic.

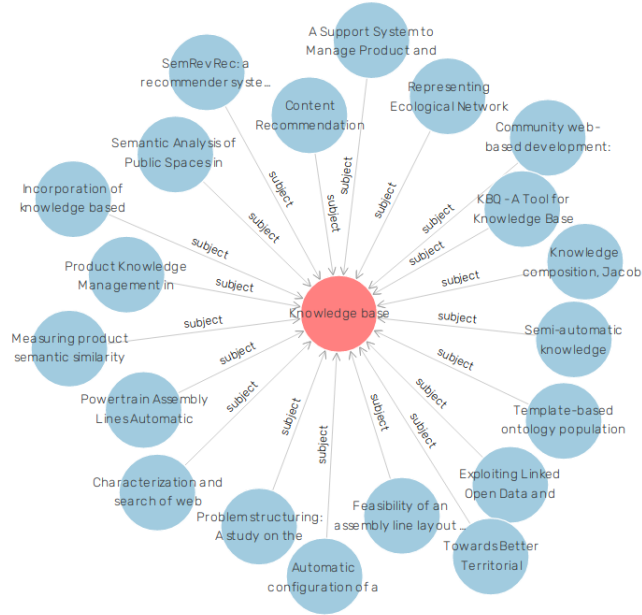


Figure 5.2: Visualization of the results obtained from the PKG when running a SPARQL query that returns all the publications that have *Knowledge Base* as topic. The RDF representation of this example will be listed in the following section.

### 5.1.2 The Graph Builder

We implemented the Graph Builder as a Python command-line script that uses the *rdflib*<sup>2</sup> library to create and manage an in-memory RDF representation of the graph. Such representation can be then serialized and saved as an XML file. The script takes as arguments the path to the JSON dump of IRIS, together with some options that allows to trigger specific functionalities of the script, such as:

1. The update of an already existing RDF graph with new statements, used for example to add the predicted triples.

<sup>2</sup><https://github.com/RDFLib/rdflib>

2. The number of topics that must be extracted from each abstract by TellMeFirst, the default value is seven.
3. The addition of the topics images, which are scraped from DBpedia and added as attributes of the *TMFResource* instances.

The Graph Builder firstly declares the namespaces and the ontology used to define the RDF representation of the entities and the relation inside the graph as written in the previous section, then parses the arguments and options received and execute the corresponding activities.

If the creation of a new graph is requested, the script reads the JSON dump in a Python list and instantiates a *ThreadPoolExecutor*, a Python abstraction that allows to execute a function asynchronously by spawning a predefined number of threads. Each thread asynchronously executes a function that process a single record of the dump. The concurrent access to the records list is not a problem, being the Python lists implemented as thread safe containers.

Each record is processed by:

1. Matching the record metadata with the ontology classes and instantiating the corresponding entities.
2. Requesting to TellMeFirst, via its API, the extraction of the topics from the publication abstract.

In the first step when a field that matches a class is found a corresponding RDF triple that instantiates a new object is created, so a new entity is added to the graph. If the entity is already present, an RDF triple that connects the publication to the already existing entity is added.

The topic extraction is requested to TMF by sending an HTTP POST request containing the publication abstract and the number of topics to be extracted to its REST API<sup>3</sup>. The response contains the list of the DBpedia resources that TMF extracted as main topics for the publication. Such topics are added to the graph by instantiating the corresponding *TMFResource* entities, and are linked to the publication by means of the *dc:subject* relation.

Even if the Python interpreter implementation poses some limitations in the actual advantage of executing multithreaded code in CPU bound

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<sup>3</sup>[http://tellmefirst.polito.it/TellMeFirst\\_API\\_Guide.pdf](http://tellmefirst.polito.it/TellMeFirst_API_Guide.pdf)

scenarios, in our case the use of multiple threads greatly improved the time required to build the graph, being the implementation I/O bound by the communication with the REST API of TellMeFirst.

The two steps described above allows to generate a full RDF description of a publication starting from its metadata, which is then enriched by the topics extracted by TellMeFirst. For instance, the listing that follows contains an example RDF description generated by the Graph Builder. As can be saw, not only a new publication entity is created, but also the author and the extracted topic are added to the graph and linked to the publication.

```

1 <?xml version="1.0" encoding="UTF-8"?>
2 <rdf:RDF
3   xmlns:pkg="http://pkg.polito.it/"
4   xmlns:dc="http://purl.org/dc/terms/"
5   xmlns:foaf="http://xmlns.com/foaf/0.1/"
6   xmlns:rdf="http://www.w3.org/1999/02/22-rdf-syntax-ns#"
7   xmlns:rdfs="http://www.w3.org/2000/01/rdf-schema#"
8   xmlns:dbpr="http://dbpedia.org/resource/"
9   xmlns:xm1s="http://www.w3.org/2001/XMLSchema#">
10  <rdf:Description rdf:about="pkg:publications/2670709">
11    <rdf:type rdf:resource="pkg:ontology/Publication"/>
12    <dc:identifier>2679709</ns1:identifier>
13    <rdfs:label>Title</rdfs:label>
14    <dc:abstract>Abstract of the publication</ns1:abstract>
15    <dc:subject rdf:resource="dbpr:Knowledge_Base"/>
16    <dc:creator rdf:resource="pkg:authors/rp00000"/>
17    <dc:dateSubmitted rdf:datatype="xm1s:date">
18      2017-01-01
19    </dc:dateSubmitted>
20  </rdf:Description>
21  <rdf:Description rdf:about="dbpr:Knowledge_Base">
22    <rdf:type rdf:resource="pkg:ontology/TMFResource"/>
23    <dc:identifier>Knowledge_Base</dc:identifier>
24    <rdfs:label>Knowledge Base</rdfs:label>
25  </rdf:Description>
26  <rdf:Description rdf:about="pkg:authors/rp00000">
27    <rdf:type rdf:resource="pkg:ontology/Author"/>
28    <foaf:name>Surname, Name</foaf:name>
29    <dc:identifier>rp00000</dc:identifier>
30    <rdfs:label>Surname, Name</rdfs:label>
31  </rdf:Description>
32 </rdf:RDF>

```

Table 5.1: Number of entities and edges in the Polito Knowledge Graph.

Polito Knowledge Graph				
Number of entities per-class				
Publication	Author	Journal	TMFResource	AuthorKeyword
23268	34886	3211	16988	41807
Number of edges per-relation				
Creator	Contributor	Publisher	Subject (TMF)	Subject (Keywords)
23268	80819	11243	107093	77492

By translating every record of the IRIS database dump into a set of RDF statements, we obtained an RDF graph composed of entities that are uniquely identified and linked together by meaningful relations. We decided to call such graph the *Polito Knowledge Graph*, being a comprehensive representation of the Politecnico di Torino academic community. The graph links together researchers, their fields of interest, the scientific papers produced by them and the journals in which such papers have been published.

In Table 5.1 we summarized some statistics about the PKG, showing the number of entities instantiated for each class, and the number of edges for each relation type.

After all the publications have been processed by the Graph Builder, the internal representation obtained can be serialized and exported as an XML file that contains the definition of all the RDF triples that forms the Polito Knowledge Graph. Such RDF representation will be then used as input for the Enhancer Module or loaded into a triplestore (for example Blazegraph<sup>4</sup>) in order to be queried by the viewer component.

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<sup>4</sup><https://blazegraph.com/>



## 5.2 Enhancing the Polito Knowledge Graph

Once the first version of the Polito Knowledge Graph has been built, the Enhancer Module can be used for the inference of new facts. In this section we will discuss how we created a usable dataset for the encoder starting from the RDF graph, how we employed a R-GCN architecture to encode the entity characteristics into meaningful embeddings and how we obtained and evaluated the new predicted triples.

### 5.2.1 The Dataset Builder

The Model Trainer cannot learn the nodes embeddings by directly taking as input the RDF representation of the graph. To solve this issue, another component, the Dataset Builder, is in charge of creating a useful dataset for the learning task.

As discussed in the previous section, some of the metadata available in the IRIS publications records are not uniquely identified, however, we have still added them to the graph for sake of completeness. As a consequence of this, the corresponding entities do not represent unambiguous semantic information, the effect being that such entities may only add noise to the dataset, thus leading to incorrect predictions.

To avoid this issue, we decided to include in the dataset only the entities which are unique and unambiguous in the whole graph:

1. The publications, identified by their IRIS identifier.
2. The authors, identified by their Politecnico di Torino internal identifier.
3. The journals, identified by their ISSN.
4. The topics extracted by TMF, identified by their DBpedia URI.

With respect to the full set of entities of the Polito Knowledge Graph, we discarded all the external contributors and co-authors, for which we only have as information the names and surnames, and all the authors keywords. The dataset is thus built from a reduced version of the initial graph.

In the following, we will explain how starting from such reduced version of the graph we built a usable dataset for the learning task.

Like the Graph Builder, also the Dataset Builder is implemented as a Python command-line script that process the RDF triples with *rdflib*, which offers a practical interface to work with RDF data. The Dataset Builder has to convert the RDF representation of the graph into an integer representation, where each class, entity and relation has a corresponding integer index assigned.

The following input data are needed by the Model Trainer to train the encoder:

1. The number of nodes, equal to the number of entities instantiated by the RDF statements.
2. The number of different relations that links together the graph entities.
3. The number of nodes labels, where each label is referred to a class of the PKG ontology.
4. A list of edges, where each edge is a Python tuple composed of three elements: the node index of the subject and the object, and the index of the predicate. How such indexes are generated is explained in the following.
5. a list of node-specific normalization constants.

To build such data structures starting from the RDF graph, the Dataset Builder leverages some look-up hash tables (implemented as Python dictionaries) that are created starting from the RDF statements and the PKG ontology. Such tables are accessed by URI, and allows to retrieve the corresponding entity or relation integer index.

1. The *nodes table* is populated by assigning to each entity a unique, increasing integer index. Such index will identify the entity when building the list of edges. Only the entities that are instances of the classes that we selected as part of the dataset are added to the table.
2. The *relations table* is built starting from the PKG ontology, assigning to each relation inside the graph a corresponding integer identifier.

When building such tables, the Dataset Builder also keeps track of the total number of nodes and relations, and builds a labels list that stores in the  $i$ -th element the class of the  $i$ -th node.

Once the look-up tables and the other data structures are built, the Dataset Builder starts to process the RDF statements that composes the graph by looping through them.

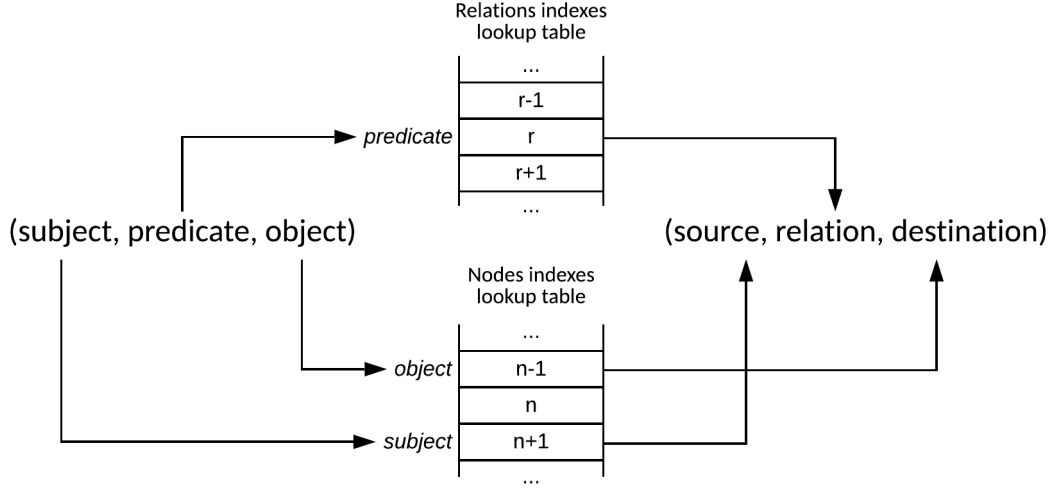


Figure 5.3: Process of translation from RDF statement composed of URIs to edge tuple composed of integer indexes.

For each statement, The Dataset Builder firstly checks whether the subject and object of the statement are present in the nodes table, if so, it builds the corresponding edge tuple by retrieving the indexes of the entities and relation through the look-up table. An example of this process is showed in Figure 5.3.

The obtained list of edges is then splitted into three separate and disjoint sets that will be used to train, evaluate and test the encoder model. We make sure to have in the training set at least one edge for every kind of relation and for each node, so to have representative nodes neighborhoods during training. This is mandatory to obtain meaningful embeddings, because as already explained in section 2.2.3, the R-GCN model learns the vector representations of the nodes by embeddings the nodes neighbors features. So is crucial to have a neighborhood structure in the training graph which is similar to the one in the full graph. Once at least one edge per relation and node have been picked, the remaining training edges are randomly taken.

The percentage of tuples used to create the three sets is an hyperparameter that can be chosen in advance. However, during our tests, we experienced that picking less than 70% of nodes for training does not allowed us to maintain a representative neighborhood for each node, resulting in an inaccurate link prediction model. As a consequence of this, we splitted the edges list in approximately 90% of the tuples for train, 5%

Table 5.2: Statistics of the dataset produced by the Dataset Builder.

Dataset			
Number of nodes	Number of classes/labels	Number of relations	Total number of edges
47996	4	4	170.593
Number of train, evaluation and test samples			
Train edges	Evaluation edges	Test edges	
153.531	8528	8534	

for the validation and 5% for testing.

Once the dataset have been created and the list of edges splitted into the three sets, it is serialized and saved, so that for future run of the Model Trainer is not required to create the dataset from scratch. Moreover, since a part of the edges are randomly picked, this allowed us to evaluate the different hyperparameters values upon the same set of triples.

Table 5.2 summarizes the size and characteristics of the dataset obtained by the Dataset Builder. The size of the dataset obtained, in particular the number of nodes and edges, can be compared to the size of the initial RDF graph, whose statistics are available in Table 5.1. As can be saw, more then half the nodes, the ones referred to the author keywords and the external authors, have been removed.

### 5.2.2 The Model Trainer

Modern machine learning algorithms are implemented as deep networks composed by many layers and parameters. Building such architectures from the ground-up would be almost unfeasible, especially when dealing with challenging kinds of data structures, such as multi-relational graphs. Many frameworks and libraries that relies on some basic abstractions have been introduced in the latest years, allowing the implementation of complex architectures without the need of coding them from scratch. Moreover, one of the key aspects when building machine learning models is to build highly efficient implementations, especially when working with lots of data, as in the case of knowledge graphs, thus leveraging highly optimized and tested software is crucial.

One of the most commonly used frameworks that allows to build and train ML models is PyTorch [?]. The goal of PyTorch is to provide some simple yet expressively powerful components that could be used to implement any sort of neural network, simplifying some of the most challenging aspects, like the implementation of the backpropagation phase. The two major abstractions introduced in the framework are the *tensor* and the *autograd*. Tensors are the basic data structure in PyTorch, they support both CPU and GPU execution and all the major operations for multidimensional matrices. The autograd package is the key novelty of PyTorch, allowing to perform automatic differentiation of all the operations performed on tensors by leveraging an internal representation in the form of a computational graph. However, even if the framework provides all the building blocks for developing complex neural networks, it is not suitable when dealing with graph data, due to the lack of support for the message passing paradigm by the tensor interface.

Deep Graph Library [?] (DGL) provides a more comprehensive solution specifically designed to work with graph data. DGL is built on top of existing ML frameworks (e.g. PyTorch, MXNet), and offers a simple interface that facilitate the implementation of the message passing framework. Supporting such framework, with DGL is possible to implement every deep learning algorithm whose forward pass can be divided into a per-edge message computation and a per-node message aggregation. Some of the architectures that can be implemented are depicted in [?], with R-GCN being one of them.

With DGL, a graph can be created by instantiating an object of the *DGLGraph* class, which offers a dictionary-like interface for adding to the graph both nodes and edges, together with their features or any other relevant data for the training, for instance the normalization constants. The message passing paradigm is implemented by means of the *send* and *recv* functions, which allows to define the two basic operations for the per-edge messages construction and the per-node aggregation.

As already mentioned before, we decided to use a R-GCN architecture to build an encoder model which is capable to produce the nodes embeddings starting from the graph structure. In the following we will look at how we trained such encoder model, how we implemented it with DGL, which learning problem we wanted our model to solve and how we evaluated such model, so to select the best parameters for the learning task.

To train and test the model we used a workstation whose hardware and software specifications are summarized in Table 5.3. The same workstation have been used to execute the Link Evaluator component. As we will discuss later, the training and evaluation speed have been bottlenecked by the system and video memory at our disposal.

Table 5.3: Hardware and software specifications of the workstation used to run the Model Trainer and the Link Evaluator.

Workstation hardware and software specifications	
<b>CPU</b>	x86-based with 4 cores and 8 threads clocked at 3.9GHz.
<b>System memory</b>	32GB of DDR3 1600MHz RAM.
<b>GPU</b>	8GB of VRAM and 2304 cores.
<b>Boot drive</b>	512GB SATA-III solid state drive.
<b>Swap memory</b>	A dedicated 64GB partition on the boot drive.
<b>OS</b>	Ubuntu Server 18.04 LTS with Linux Kernel 4.15.
<b>SW packages</b>	CUDA toolkit 10.1, PyTorch 1.2 and DGL 0.3.

As saw in previous section, the input dataset for the Model Trainer is composed by a list of edges, where 90% of them are used for the training, and the remaining are splitted in half and used for the validation and testing of the model. The training edges are used to build a *training graph*, which is the one that will be used to learn the node embeddings. Regarding the node features, we choose to follow a featureless approach, so to only leverage the graph structure to build the embeddings. Following this idea, we used as input features a one-hot encoded vector for each node, thus obtaining an  $N \times N$  feature matrix, where  $N$  is the number of nodes in the training graph.

Regarding the architecture, we employed a R-GCN composed of two hidden layers, with the first layer convolutional filters being  $500 \times N$ , and the second layer filters being  $500 \times 500$ . With such architecture the embeddings obtained as output of the forward pass are shaped as  $1 \times 500$  row vectors. For the per-node normalization constants we used the inverse of the nodes indegree.

Because of the size of the graph, the training is performed in batches. To do so, at each training epoch the Model Trainer randomly samples a

subset of the training edges. As a consequence of this, more than one epoch may be required to train over all the nodes in the training graph.

After the edges have been sampled, a predefined number of corrupted edges are generated starting from the sampled ones. This is done because the learning problem that we want to solve is a classification problem: we want our model to be able to give a high DistMult score to the true edges, and a low score to the corrupted ones. This will allow us to obtain a model that is able to recognize true facts, and thus can be used to predict the probability with which an unseen triple may belong to the Polito Knowledge Graph. The negative samples are associated to the label 0, while the positive samples (the true facts) are associated to the label 1. We have chosen a 1-to-10 ratio for the negative samples, having 10 corrupted edges generated starting from each true fact, as it is done for most models found in literature that leverages negative sampling.

The embeddings of the relations are randomly initialized, because as explained when introducing DistMult in section 2.2.3, they are not subject of the learning process.

At each epoch the training is performed in the following steps:

1. The forward pass of the R-GCN is executed, taking as input the *DGLGraph* generated starting from the positive samples (the sampled training edges), which contains both the nodes features and the per-edge normalization constants. The neural network gives as output the embeddings of the graph nodes.
2. Such embeddings are used to calculate the DistMult scores of both the positive and the negative samples. The sigmoid<sup>5</sup> function is used to cap the scores between zero and one. Such capped scores represents the probabilities with which the model considers the corresponding edges as true facts.
3. The batch loss is calculated and the filters parameters are updated through backpropagation using the *Adam* optimizer.

The loss function used is the *binary cross entropy loss*, which is commonly used for classification tasks with just two classes, as in our case:

---

<sup>5</sup>  $\sigma(x) = \frac{1}{1+\exp^{-x}}$

$$\mathcal{L}(x, y, f(x)) = -\frac{1}{E} \sum_{i=1}^E (y_i \cdot \log(f(x_i)) + (1 - y_i) \cdot \log(1 - f(x_i))) \quad (5.1)$$

$E$  is the number of edges for the current batch, taking into account both positives and negatives samples, and  $y_i$  represents their associated label.

In our case, the function  $f(x_i)$  computes the predicted probability for the  $i$ -th sample to be a true fact by applying the sigmoid function to its score, which is calculated with the DistMult factorization method:

$$f(x_i) = f(s_i, r_i, o_i) = \sigma(e_s^T R_r e_o) \quad (5.2)$$

Where  $e_s$  and  $e_o$  are the embeddings for the source node (subject) and destination node (object),  $R_r$  is obtained by transforming the embedding of the relation (predicate)  $e_r$  into a diagonal matrix and  $\sigma$  is the sigmoid function. The resulting capped score can be interpreted as the confidence of the model in the fact that the corresponding edge represents a true fact.

The loss function used sets as training goal the correct classification of negative and positive samples, so to obtain a score as low as possible for the negative ones, while instead identify the true triples sampled from the PKG as true facts, thus giving them an high score.

As a consequence of this, the obtained model should be able to correctly predict if an unseen triple can be considered as a good candidate for the inclusion in the Polito Knowledge Graph.

To avoid overfitting, the following  $L2$  regularization is added to the training loss:

$$L2_{regularization} = \lambda \cdot \left( \frac{1}{N} \sum_{i=0}^N e_i^2 + \frac{1}{R} \sum_{r=0}^R w_r^2 \right) \quad (5.3)$$

Where  $\lambda$  is an hyperparameter used to scale the regularization,  $N$  is the number of nodes,  $R$  is the number of relations,  $e_i$  is the embedding of the  $i$ -th node and  $w_r$  is a randomly initialized vector that represents the embedding of the relation  $r$ .

In addition to the regularization, dropout with a probability of 20% is used to randomly update only some of the network weights at each training epoch.



The training can be performed both on CPU, which in respect to the GPU has access to four times the memory, but has less execution units, or on the GPU, which is memory-limited but can leverage its highly parallelized architecture. The usage of CPU or GPU and the number of CPU threads to be used are hyperparameters that can be changed before starting the training.

We were able to perform the training on GPU with a sampling size of 20'000 edges, that adding the negative samples produces a per-epoch training set composed of a total of 220'000 edges. While having to train over more epochs due to video memory constraints, in comparison to the execution on CPU the training time has been reduced by several order of magnitudes: from approximately 30 seconds per epoch for the forward plus backward pass, to less than three hundred milliseconds.

The biggest bottleneck while training remains the sampling phase, which is implemented as single threaded code using NumPy [?], and requires approximately 5 seconds to be performed with the hardware at our disposal, which has fairly limited single-threaded performances. We leave the implementation of more efficient sampling methods for future releases.

In order to identify the best parameters, we evaluate the model on a separate validation set during training. The validation is not performed at every epoch due to computation time constraints, and the frequency with which it is executed is an hyperparameter decided in advance. How the evaluation of the model is performed will be presented in the next chapter. If during the validation the model trained at the current epoch is identified as the best so far, its parameters are serialized and saved on local storage. Once the training is finished, the previously exported best epoch model is loaded from memory and its accuracy is evaluated on the test set.

After the training, validation and test phases are completed, the obtained encoder model should be able to give as output meaningful embeddings of the PKG nodes, which can be used to evaluate new facts using a factorization method like DistMult as decoder.

### 5.2.3 The Link Evaluator

Once the model has been trained, and the testing phase demonstrated that it has embedded the characteristics needed to build semantically meaningful nodes embeddings, the Link Evaluator can be used to leverage the

trained model for the prediction of new facts.

As already saw, a *fact* in the knowledge based is composed of a source and a destination nodes, the subject and the object, and a relation, called predicate, that connects the two nodes establishing a connection between them.

We are interested in the prediction of facts that are not already present in the knowledge base, and that are coherent with the defined ontology, both in domain and range.

The Link Evaluator loads in memory the model produced by the Model Trainer and performs the following tasks:

1. It builds the graph of existing facts by merging together the train, test and evaluation edges.
2. Then it gets the embeddings of the nodes of such graph by feeding it to the trained encoder.
3. It creates the set of every possible fact, and scores the triples in such set, only taking the ones with the highest score.
4. Finally, the obtained predicted triples are saved in a JSON file and imported by the Graph Builder, that will add them to the Polito Knowledge Graph as new RDF statements.

The node embeddings are obtained by feeding the graph of existing facts to the R-GCN encoder in evaluation mode, so to disable the gradients tracking in the PyTorch computational graph, which greatly improves the performances.

To create the set of new facts, the Link Evaluator firstly generates the set of all the graph nodes, and then creates the set of possible facts by generating, for every node, all the possible links to every other node in the graph. This is done by taking for each source node (subject) all the possible combinations of relations (predicates) and destination nodes (objects).

However, a majority of this automatically generated triples will be semantically incorrect. For example, among the permutations there will be triples that connect together two publications using as predicate the relation *dc:subject*, which is clearly meaningless, because it is saying that the major topic of a publication is another publication. For this reason, the triples that are not correct with respect to the PKG ontology are immediately discarded.

Once the semantically incorrect facts are removed, the remaining edges could be scored by applying the DistMult factorization. Then, the scored triples are grouped by subject and relation and sorted by their score in descending order. The number of scored triple to save for each couple subject-relation is an hyperparameter decided in advance, we used 30 as default value.

The resulting new facts are then exported as a JSON file that can be used by the graph builder to build the corresponding RDF triples. Such triples could then be added the Polito Knowledge Graph, completing its knowledge with new facts.

Moreover, the predicted triples could be used to obtain insights and recommendations that can be showed by the Viewer module.

For instance, among all the predictions there will be triples that links a researcher to a publication of which he is not an author: such predicted authors of a scientific paper can be thought as researchers that shares the same research interests of the actual publication authors.

Leveraging such predictions, the recommendation system could suggest to a researcher unexplored research topics, the profile of other researchers which share the same research interest but with which he has never worked before, or scientific journals that publish papers in his same research field.



## Chapter 6

# Evaluation

In this chapter we will describe the methods used to evaluate the trained model. The evaluation is performed both during training and testing. For the training task, we evaluate the model over the validation set, which is used to check whether the latest updated parameters led to a more accurate model. After the training is finished, the evaluation is performed over a test set whose data has never been seen by the model during training, so to test whether it is able to correctly score unseen true facts.

We evaluated the model using the *entity ranking* approach, which is commonly used for the evaluation of knowledge base embedding models. Such approach evaluates the ability of a model to identify the triples that belongs to the knowledge base with respect to the set of all possible facts.

To do so, it is firstly required to generate two sets of corrupted facts for each triple inside the knowledge base. This is actually performed by corrupting the source and destination node of every edge in the evaluation set: starting from the  $i$ -th evaluation edge  $(s_i, p_i, o_i)$  we obtain two sets of corrupted edges  $(a, p_i, o_i)$  and  $(s_i, p_i, b)$ , with  $a, b \in [0, N]$ , where  $N$  is the number of nodes of the whole graph.

Then, for every triple in the evaluation set, the triple itself and its corresponding corrupted ones are scored using the same factorization method used for the training, in our case DistMult. The triples are then sorted in descending score, and the position of the evaluation triples inside such sorted list represent their *rank*.

Ideally, the best possible model should be able to always rank all the triples in the evaluation set on first position, thus giving them the highest rank, which is equal to 1. Such model would be always able to identify

the correct facts, and therefore we could be confident in the truthfulness of its predictions. However, obtaining such model is unfeasible in practice.

To obtain a measure of the accuracy with which the trained model is able to identify the correct triples, we used the following metrics:

1. The Reciprocal Rank (RR), which can be computed for a single evaluated edge  $(s_i, p_i, o_i)$  as  $1/\text{rank}_{(s_i, p_i, o_i)}$ , where the rank is obtained as explained above.
2. The Mean Reciprocal Rank (MRR), which is the average of the RR of all the evaluation edges. The value obtained is representative of the accuracy of the entire model.
3. The Hits-at-N (HITS@N), which is the number of triples for whom the computed rank is between 0 and  $N$ .

So for example, a triple ranked in the tenth position of the sorted list of scores will have a RR of 0.1. If the triples in the evaluation set are 100, with half of them ranked in first position and the other half in the tenth, the obtained MRR will be  $\frac{1}{100} \sum_{i=1}^{100} \frac{1}{\text{rank}_{(s_i, r_i, o_i)}} = 0.55$ , while the HITS@1 will be equal to 50, and the HITS@10 will be equal to 100.

We performed the evaluation on the same workstation used for the training, whose specifications can be found in Table 5.3. However, in contrast to the training, which is performed on GPU to leverage its parallelized architecture, the evaluation had to be made on CPU due to memory constraints.

In fact, as a consequence of the dimensionality of the graph, and therefore that of the set of corrupted edges, the matrices used to perform the evaluation cannot be stored in the GPU memory. Moreover, also the system memory at our disposal is not sufficient to perform such task in parallel for all the edges, thus requiring the evaluation to be performed in batches. With the memory at our disposal we have been able to perform the evaluation in batches of 80 triples. Given that we were not able to run the computation on the highly parallel architecture of the GPU, and that we were limited by the small amount of memory and cores of our CPU, the evaluation proved to be the slowest task of the whole pipeline, taking approximately twenty minutes per run. This is why we have been not able to perform the evaluation of the model at every training epoch, however,

we found that evaluating every 500 epochs still allowed us to obtain useful metrics for the choice of the best hyperparameters.

Table 6.1: Cross-validation of learning rate and regularization parameters using as benchmark the Mean Reciprocal Rank (MRR) value. The first table shows the MRR of the best model found during the training phase. The second table shows the MRR obtained by the best model when tested over the test set.

(a) MRR of the best model found during training.

		Learning Rate				
Regularization	MRR	0.05	0.01	0.005	0.001	0.0005
	1.0	0.0	0.0623	0.0504	0.0823	0
	0.5	0.0	0.0677	0.0718	0.0891	0.0865
	0.1	0.0004	0.0703	0.852	0.0764	0
	0.05	0.0	0.0707	0.0763	0.0673	0
	0.01	0.0	0.062	0.0725	0.0	0

(b) MRR obtained when evaluating the best model over the test set.

		Learning Rate				
Regularization	MRR	0.05	0.01	0.005	0.001	0.0005
	1.0	0.00	0.0611	0.0497	0.081	0
	0.5	0.00	0.0662	0.0723	0.0852	0.0818
	0.1	0.0002	0.0693	0.0827	0.0726	0
	0.05	0.00	0.0687	0.0726	0.0654	0
	0.01	0.00	0.0589	0.07	0.00	0

In Table 6.1 we show the result of the evaluation of the model over some possible combinations of hyperparameters, in particular, we computed the

MRR for possible combinations of the learning rate and of the regularization parameter. As can be seen from the table, we found the best values to be 0.001 for the former and 0.5 for the latter.

The fairly high value of the regularization parameter (0.5) is particularly interesting: we motivate this result as a consequence of the fact that our graph is composed of a very low number of different relations, which is particularly true if compared to other graphs, such as the one used for benchmarking. Given how the R-GCN encoder works, less relations means less convolutional filters, so less characteristics that are used to generate the embeddings. In this regard, the regularization parameter helps the model to differentiate the embeddings obtained, even if there are few relations, and allows to not overfit on common features.

Regarding the MRR value obtained for the best model, it can be interpreted as the fact that, in average, the evaluated triples are ranked in the *eleventh* position. The authors of R-GCN obtained an MRR value of 0.125 over the FB15k-237 dataset, which is commonly used as benchmark for the evaluation of link prediction model. This value means that the true facts are ranked, in average, in the *eighth* position. Our result is thus not so distant from the one obtained over the benchmarking dataset. Moreover, if we consider the number of nodes in the PKG dataset, the result obtained is particularly good, given that the model is able to rank (in average) the true triple in the eleventh position when ranked with 47.995 corrupted triples, while in FB15k-237 for each true fact only 14.951 corrupted ones are generated.

The Figure 6.1 shows how the testing triples are distributed over the possible ranking values. As can be seen, the model is able to correctly assign an high rank to the majority of the true facts.

When looking at the statistics of the Polito Knowledge Graph, we can see how a big part of the entities, and in particular of the edges, are related to the publications topics. In particular, as showed in Table 5.1, almost a third of the PKG edges connect a publication to a subject extracted by TellMeFirst.

Given that the amount of topics extracted by TMF for each publication is a Graph Builder parameter, we decided to generate three new graphs with different numbers of topics extracted for each publication, so to evaluate how having a different number of *TMFResource* entities inside the



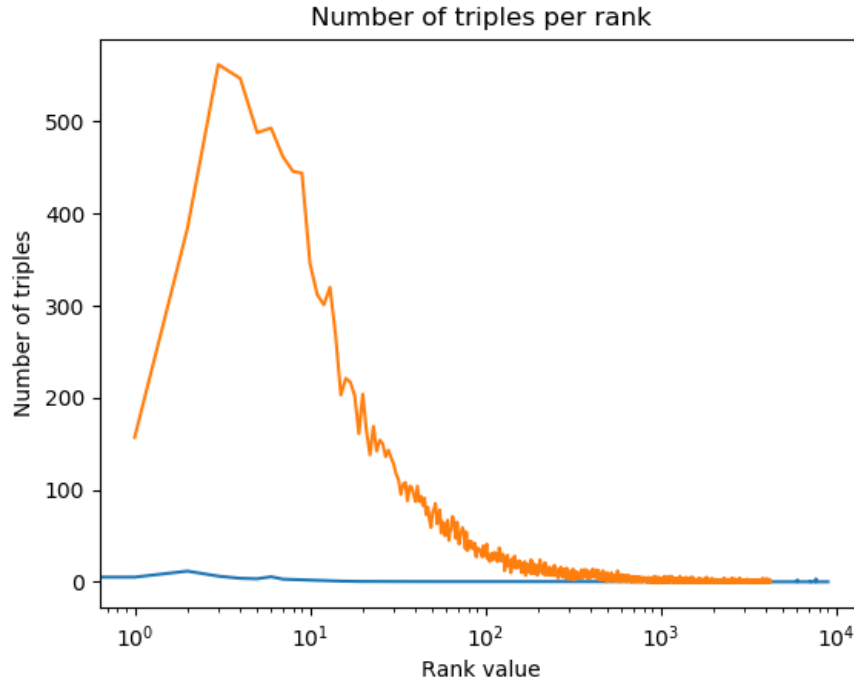


Figure 6.1: Number of triples obtained for each ranking position when evaluating the best model found over the test set.

knowledge graph could have an impact on the trained model.

As can be seen from the results showed in Table XXX.



## Chapter 7

# Conclusions and future work