

# **Neural Graph Embedding Methods for Natural Language Processing**

A THESIS  
SUBMITTED FOR THE DEGREE OF  
**Doctor of Philosophy**  
IN THE  
**Faculty of Engineering**

BY

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November, 2019

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I, **Shikhar Vashishth**, with SR No. **04-04-00-15-12-16-1-13374** hereby declare that the material presented in the thesis titled

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DEDICATED TO

*My Teachers*

*who enlightened me with all knowledge.*

# Acknowledgements

I want to offer my sincere thanks to my advisors Dr. Partha Pratim Talukdar and Prof. Chiranjib Bhattacharyya, who gave me the freedom to work in my area of interest and have been very supportive throughout the course of my PhD. I also want to thank Dr. Manaal Faruqui, who accepted the role of being my mentor and guided me in my research. I am very grateful to all my teachers at Indian Institute of Science for giving me a clear understanding of the basics which were essential for completing this work.

I feel fortunate to get the opportunity to collaborate with several researchers during my PhD. I want to thank Prince Jain, Shib Sankar Das, Swayambhu Nath, Rishabh Joshi, Sai Suman, Manik Bhandari, Prateek Yadav, Soumya Sanyal, Vikram Nitin, Shyam Upadhyay, Gaurav Singh Tomar and all the members of MALL Lab for their support and help. I am also thankful to my parents and friends for their support throughout my stay in Bangalore. Finally, I would like to thank Almighty God for all His blessings without which this would not have been possible.

# Abstract

Graphs are all around us, ranging from citation and social networks to Knowledge Graphs (KGs). They are one of the most expressive data structures which have been used to model a variety of problems. Embedding graphs involve learning a representation of all nodes (and relations) in the graph which allows to effectively utilize graphs for various downstream problems. In this thesis, we explore such techniques for alleviating sparsity problem in knowledge graphs and for tasks such as document timestamping and word representation learning. We also list some of the limitations of existing graph embedding methods and propose solutions for addressing them.

Knowledge graphs are structured representations of facts in a graph, where nodes represent entities and edges represent relationships between them. Recent research has resulted in the development of several large KGs; examples include DBpedia, YAGO, NELL, and Freebase. However, all of them tend to be sparse with very few facts per entity. For instance, NELL KG consists of only 1.34 facts per entity. In the first part of the thesis, we explore three solutions to alleviate this problem through neural graph embedding based techniques: (1) KG Canonicalization, i.e., identifying and merging duplicate entities in a KG, (2) Relation Extraction which involves automating the process of extracting semantic relationships between entities from unstructured text, and (3) Link prediction which includes inferring missing facts based on the known facts in a KG. For KG Canonicalization, we propose CESI (Canonicalization using Embeddings and Side Information), a novel approach that performs canonicalization over learned embeddings of KGs. The method extends recent advances in KG embedding by incorporating relevant NP and relation phrase side information in a principled manner. For relation extraction, we propose RESIDE, a distantly-supervised neural relation extraction method which utilizes additional side information from KGs for improved relation extraction. Finally, for link prediction, we propose InteractE which extends ConvE, a convolutional neural network-based link prediction method. InteractE increases the number of feature interaction through three key ideas feature permutation, a novel feature reshaping, and circular convolution. Through extensive experiments on multiple datasets, we demonstrate the effectiveness of our proposed

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methods.

Traditional Neural Networks like Convolutional Networks and Recurrent Neural Networks are constrained to learn representation of Euclidean data. However, graphs in Natural Language Processing (NLP) are prominent. Recently, Graph Convolutional Networks (GCNs) have been proposed to allow Deep Learning models to exploit graph structures by embedding them in an end-to-end fashion. GCNs have been successfully applied for several problems in NLP and computer vision. In the second part of the thesis, we utilize GCNs for Document Timestamping problem, which forms an essential component of tasks like document retrieval, and summarization. For this, we propose NeuralDater which leverages GCNs for jointly exploiting syntactic and temporal graph structures of document for obtaining state-of-the-art performance on the problem. We also propose SynGCN, a flexible Graph Convolution based method for learning word embeddings which utilize the dependency context of a word instead of linear context for learning more meaningful word embeddings.

Finally, in the last part of the thesis, we address two limitations of existing GCN models, viz, (1) The standard neighborhood aggregation scheme puts no constraints on the number of nodes that can influence the representation of a target node. This leads to a noisy representation of hub-nodes which covers almost the entire graph in a few hops. To address this shortcoming, we propose ConfGCN (Confidence-based GCN) which estimates confidences to determine the importance of a node on another during aggregation, thus restricting its influence neighborhood. (2) Most of the existing GCN models are limited to handle undirected graphs. However, a more general and pervasive class of graphs are relational graphs where each edge has a label and direction associated with it. Existing approaches to handle such graphs suffer from over-parameterization and are restricted to learning representation of nodes only. We propose CompGCN, a novel Graph Convolutional framework which jointly embeds entity and relations in a relational graph. CompGCN is parameter efficient and scales with the number of relations. It leverages a variety of entity-relation composition operations from KG Embedding techniques and achieves demonstrably superior results on node classification, link prediction, and graph classification tasks.

# Publications based on this Thesis

The work in this dissertation is primarily related to the following peer-reviewed articles:

1. **Shikhar Vashishth**, Prince Jain, and Partha Talukdar. “CESI: Canonicalizing Open Knowledge Bases using Embeddings and Side Information”. In Proceedings of the World Wide Web Conference (WWW), 2018.
2. **Shikhar Vashishth**, Shib Shankar Dasgupta, Swayambhu Nath Ray, and Partha Talukdar. “Dating Documents using Graph Convolution Networks”. In Proceedings of the 56th Annual Meeting of the Association for Computational Linguistics (ACL), 2018.
3. **Shikhar Vashishth**, Rishabh Joshi, Sai Suman Prayaga, Chiranjib Bhattacharyya, and Partha Talukdar. “RESIDE: Improving Distantly-Supervised Neural Relation Extraction using Side Information”. In Proceedings of the 2018 Conference on Empirical Methods in Natural Language Processing (EMNLP), 2018.
4. **Shikhar Vashishth\***, Prateek Yadav\*, Manik Bhandari\*, and Partha Talukdar. “Confidence-based Graph Convolutional Networks for Semi-Supervised Learning”. In Proceedings of the International Conference on Artificial Intelligence and Statistics (AISTATS), 2019.
5. **Shikhar Vashishth**, Manik Bhandari, Prateek Yadav, Piyush Rai, Chiranjib Bhattacharyya, and Partha Talukdar. “Incorporating Syntactic and Semantic Information in Word Embeddings using Graph Convolutional Networks”. In Proceedings of the 57th Annual Meeting of the Association for Computational Linguistics (ACL), 2019.
6. **Shikhar Vashishth\***, Soumya Sanyal\*, Vikram Nitin, and Partha Talukdar. “InteractE: Improving Convolution-based Knowledge Graph Embeddings by Increasing Feature Interactions”. In Proceedings of 34th Conference on Artificial Intelligence (AAAI), 2020.

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\* Equal Contribution

## Publications based on this Thesis

The following articles are also discussed but are currently under review.

7. **Shikhar Vashishth\***, Soumya Sanyal\*, Vikram Nitin, and Partha Talukdar. “Composition-based Multi-Relational Graph Convolutional Networks”. Under review in International Conference on Learning Representations, 2020. *arXiv preprint arXiv:1911.03082*.

Finally, the following articles have also been completed over the course of the PhD but are not discussed in the thesis:

8. Prateek Yadav, Madhav Nimishakavi, Naganand Yadati, **Shikhar Vashishth**, Arun Rajkumar and Partha Talukdar. “Lovasz Convolutional Networks”. In Proceedings of the International Conference on Artificial Intelligence and Statistics (AISTATS), 2019.
9. **Shikhar Vashishth**, Shyam Upadhyay, Gaurav Singh Tomar, and Manaal Faruqui. “Attention Interpretability Across NLP Tasks”. *arXiv preprint arXiv:1909.11218*, 2019.
10. Zhiqing Sun\*, **Shikhar Vashishth\***, Soumya Sanyal\*, Partha Talukdar, and Yiming Yang. “A Re-evaluation of Knowledge Graph Completion Methods”. *arXiv preprint arXiv:1911.03903*, 2019.

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# Chapter 1

## Introduction

Graphs are pervasive data structures that have been used to model a variety of problems. Embedding graphs involves learning a vector representation for each node (and relation) in the graph. This enables to effectively make inference over the graph and solve various tasks such as link prediction and node classification. Such techniques have also been extensively explored for Knowledge Graphs (KGs), which are a large storehouse of world facts in a graph format. Instances of KGs include Freebase [21], WordNet [132], YAGO [185], and NELL [137]. KGs find application in a variety of tasks, such as relation extraction [133], question answering [24, 25], recommender systems [231], and dialog systems [122]. Formally, Knowledge graphs are defined as a structured representation of facts in a graph, where nodes represent entities and edges denote relationships between them. This can be represented as a collection of triples  $(s, r, o)$ , each representing a relation  $r$  between a “subject-entity”  $s$  and an “object-entity”  $o$ . A KG example is presented in Figure 1.1. Here, *John Lennon*, *Beatles*, and *Liverpool* are nodes in the knowledge graph, and edges indicate different facts about them. For instance, the fact that “*John Lennon was born in Liverpool*” is expressed through  $(\text{John Lennon}, \text{bornIn}, \text{Liverpool})$  triple in the KG.

Recently, neural graph embeddings are also being exploited for several Natural Language Processing applications. Graphs are prevalent in NLP, starting from dependency parse to temporal event ordering graphs, constituency parse graphs, etc. Neural graph embeddings methods allow utilizing such graph structures for a variety of downstream applications. An instance of such methods is Graph Convolutional Networks (GCNs) [47, 90], which allows embedding graphs based on the final objective in an end-to-end fashion. GCNs are a generalization of Convolutional Neural Networks (CNNs), which have been pivotal to achieve beyond human-like performance for tasks such as object recognition [96, 234] and speech recognition [176]. GCNs have been shown to be effective for tasks such as neural machine translation [15], se-

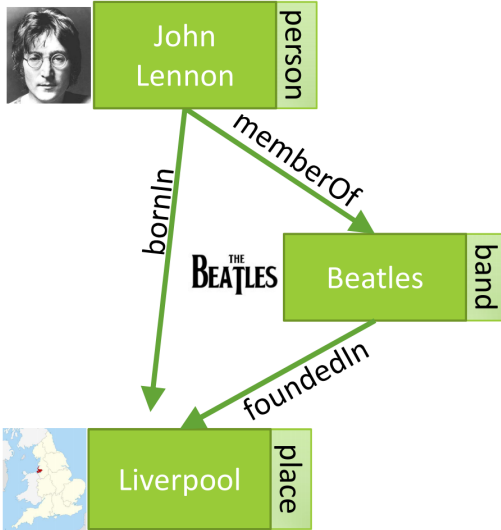


Figure 1.1: An instance of Knowledge Graph<sup>1</sup>. Here, nodes represent entities and edges indicate different relationships between them.

mantic role labeling [126], event detection [144]. However, the scope of their applicability for other NLP tasks is still an area of research. Furthermore, improving existing graph embedding methods also serves as an important research problem. Recent works [216, 217] have identified several limitations of Graph Convolutional Networks and have theoretically analyzed their potential.

Knowledge graphs have been utilized for a variety of tasks [122, 52]. However, most of the KGs are highly sparse with very few edges per entity, as also observed by [22]. For instance, NELL KG consists of only 1.34 facts per entity. This severely restricts their usage for several real-life applications. In the first part of the thesis, we present three graph embedding based techniques for addressing the sparsity problem in Knowledge Graphs. We discuss each of them below.

1. **Knowledge Graph Canonicalization** involves identifying duplicate or redundant nodes in a KG and merging them as a single node. This can be explained through a concrete example. Given two triples in a KG: (*Barack Obama, was president of, US*) and (*Obama, born in, US*). Identifying that *Barack Obama* and *Obama* refer to the same entity increases the overall facts per entity ratio. In our work, we focus on addressing such issues in Open KGs, which are automatically constructed knowledge graphs (using OpenIE algorithms [128, 40, 129]) without any pre-specified ontology. In spite of its importance, canonicalization is a relatively unexplored problem, especially in the case of Open KGs. In this work, we propose CESI

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<sup>1</sup>Image credits: <https://kgtutorial.github.io/>

(Canonicalization using Embeddings and Side Information), a novel approach which performs canonicalization over learned embeddings of Open KBs. CESI extends recent advances in KB embedding by incorporating relevant noun and relation phrase side information in a principled manner. More details are provided in Chapter 3.

2. **Relation Extraction** involves automatically extracting semantic relationships between entity pairs from unstructured text. Most of the existing KGs like Wikidata and Freebase are human-curated. Relation extraction offers a mechanism for automatically constructing these KGs without any supervision. Moreover, it can be used for further densifying existing KGs by extracting new facts from unstructured text. Since most supervised relation extraction methods require sizeable labeled training data which is expensive to construct, we utilize Distant Supervision (DS) [133] for automatically constructing a dataset. DS is based on the assumption that if two entities have a relationship in a KB, then all sentences mentioning those entities express the same relation. We propose a novel distantly-supervised neural relation extraction method, RESIDE which utilizes additional side information from KBs for improved relation extraction. It uses entity type and relation alias information for imposing soft constraints while predicting relations. RESIDE employs Graph Convolution Networks (GCN) to encode syntactic information from a text and improves performance even when limited side information is available. Please refer to Chapter 4 for more details.
3. **Link Prediction** is the task of inferring missing facts based on the known facts in a KG. For instance, if we have two triples in the KG: (*Michelle Obama, spouse of, Barack Obama*) and (*Sasha Obama, child of, Michelle Obama*). Now, based on these two facts (*Sasha Obama, child of, Barack Obama*) can be inferred. A popular approach for solving this problem involves learning a low-dimensional representation for all entities and relations and utilizing them to predict new facts. In general, most existing link prediction methods learn to embed KGs by optimizing a score function which assigns higher scores to actual facts than invalid ones. ConvE [50], a recently proposed approach, applies convolutional filters on 2D reshaping of entity and relation embeddings to capture rich interactions between their components. However, the number of interactions that ConvE can capture is limited. In this work, we analyze how increasing the number of these interactions affects link prediction performance, and utilize our observations to propose InteractE which is based on three key ideas feature permutation, a novel feature reshaping, and circular convolution. For more details, please refer to Chapter 5.

In the second part of the thesis, we focus on leveraging recently proposed Graph Convolutional Networks (GCNs) [47, 90] for exploiting different graph structures in NLP. Traditional

neural network architectures like Convolutional Neural Networks (CNNs) [99] and Recurrent Neural Networks (RNNs) [77] are limited to handle Euclidean data. GCNs have been proposed to address this shortcoming and have been successfully employed for improving performance on tasks such as semantic role labeling [126], neural machine translation [15], relation extraction [233], shape segmentation [226], and action recognition [79]. In this work, we begin with utilizing GCNs for Document Timestamping problem which is at the core of many essential tasks, such as, information retrieval [150, 106], temporal reasoning [123, 118], text summarization [207], and analysis of historical text [45]. For this, we propose NeuralDater, a neural network-based approach which leverages GCNs for jointly exploiting syntactic and temporal graph structures of a document. To the best of our knowledge, this is the first application of deep learning for the problem. The model is more elaborately described in Chapter 6. Next, we propose to use GCNs for utilizing syntactic context while learning word embeddings. Most existing word embedding methods are restricted to using the sequential context of a word. In this work, we overcome this problem by proposing SynGCN, a flexible Graph Convolution based method which utilizes the dependency context of a word without increasing the vocabulary size. We also propose SemGCN, an effective framework for incorporating diverse semantic knowledge for further enhancing learned word representations. Refer to Chapter 7 for details.

In the third part of the thesis, we address some of the significant limitations of the current Graph Convolution based models. Most of the existing GCN methods are an instantiation of *Message Passing Neural Networks* [66] which uses neighborhood aggregation scheme which puts no constraints on the number of nodes that can influence the representation of a given target node. In a  $k$ -layer model, each node is influenced by all the nodes in its  $k$ -hop neighborhood. This becomes a concern for hub nodes which covers almost the entire graph with a few hop neighbors. To alleviate this shortcoming, we propose ConfGCN, a Graph Convolutional Network which models label distribution and their confidences for each node in the graph. ConfGCN utilizes label confidences to estimate the influence of one node on another in a label-specific manner during neighborhood aggregation, thus controlling the influence neighborhood of nodes during GCN learning. Please refer to Chapter 8 for details. Apart from this, we also propose an extension of GCN models for multi-relational graphs. Most of the existing GCN models are limited to handle undirected graphs. However, a more general and pervasive class of graphs are relational graphs where each edge has a label and direction associated with it. Existing approaches to handle such graph data suffer from overparameterization and are restricted to learning representation of nodes only. We propose CompGCN, a novel Graph Convolutional framework which jointly embeds entity and relations in a relational graph. CompGCN is parameter efficient and scales with the number of relations. It leverages a variety of entity-relation

composition operations from Knowledge Graph Embedding techniques. CompGCN allows the application of GCNs for a problem which requires both node and edge embeddings such as drug discovery and KG link prediction. Through extensive experiments, we demonstrate the effectiveness of our proposed approaches. More details are presented in Chapter 9.

## 1.1 Summary of Contributions

Our contributions in the thesis can be grouped into the following three parts:

### 1.1.1 Addressing Sparsity in Knowledge Graphs

For addressing the sparsity problem in knowledge graphs, first we propose CESI (Canonicalization using Embeddings and Side Information), a novel method for canonicalizing Open KBs using learned embeddings. To the best of our knowledge, this is the first approach to use learned embeddings and side information for canonicalizing an Open KB. CESI models the problem of noun phrase (NP) and relation phrase canonicalization jointly using relevant side information in a principled manner. This is unlike prior approaches where NP and relation phrase canonicalization were performed sequentially. For densifying existing knowledge graphs using unstructured text, we propose RESIDE, a novel neural method which utilizes additional supervision from KB in a principled manner for improving distant supervised RE. RESIDE uses Graph Convolution Networks (GCN) for modeling syntactic information and has been shown to perform competitively even with limited side information. Finally, for inferring new relations based on the existing ones, we propose InteractE, a method that augments the expressive power of ConvE through three key ideas feature permutation, "checkered" feature reshaping, and circular convolution. We provide a precise definition of interaction, and theoretically analyze InteractE to show that it increases interactions compared to ConvE. Further, we establish a correlation between the number of *heterogeneous interactions* and link prediction performance. Through extensive evaluation on various benchmark datasets, we demonstrate the effectiveness of our proposed approaches.

### 1.1.2 Exploiting Graph Convolutional Networks in NLP

We leverage recently proposed Graph Convolutional Networks for exploiting several graph structures in NLP to improve performance on two tasks: Document Timestamping and Word embeddings. We propose NeuralDater, a Graph Convolution Network (GCN)-based approach for document dating. To the best of our knowledge, this is the first application of GCNs, and more broadly deep neural network-based methods, for the document dating problem. NeuralDater is the first document dating approach which exploits the syntactic as well as temporal structure of

the document, all within a principled joint model. Next, we propose SynGCN, a Graph Convolution based method for learning word embeddings. Unlike previous methods, SynGCN utilizes syntactic context for learning word representations without increasing vocabulary size. We also present SemGCN, a framework for incorporating diverse semantic knowledge (e.g., synonymy, antonymy, hyponymy, etc.) in learned word embeddings, without requiring relation-specific special handling as in previous methods. Through experiments on multiple intrinsic and extrinsic tasks, we demonstrate that our proposed methods obtain substantial improvement over state-of-the-art approaches, and also yield advantage when used with methods such as ELMo.

### 1.1.3 Addressing Limitations in Existing GCN Architectures

Finally, we address two limitations in existing Graph Convolutional Network (GCN) based methods. For this, We propose ConfGCN, a Graph Convolutional Network framework for semi-supervised learning which models label distribution and their confidences for each node in the graph. To the best of our knowledge, this is the first confidence enabled formulation of GCNs. ConfGCN utilizes label confidences to estimate the influence of one node on another in a label-specific manner during neighborhood aggregation of GCN learning. Next, we propose CompGCN, a novel framework for incorporating multi-relational information in Graph Convolutional Networks which leverages a variety of composition operations from knowledge graph embedding techniques. Unlike previous GCN based multi-relational graph embedding methods, COMPGCN jointly learns embeddings of both nodes and relations in the graph. Through extensive experiments on multiple tasks, we demonstrate the effectiveness of our proposed method. Through extensive experiments on multiple tasks, we demonstrate the effectiveness of our proposed methods.

## 1.2 Organization of Thesis

The rest of the thesis is organized as follows: In Chapter 2, we review some background on Knowledge Graphs and Graph Convolutional Networks. Then in Part 1 of the thesis, we present three methods for addressing sparsity problem in Knowledge Graph: Canonicalization (Chapter 3), Relation Extraction (Chapter 4), and Link prediction (Chapter 5). In Part 2, we present two novel applications of Graph Convolutional Networks in NLP for Document Timestamping (Chapter 6) and Word embedding (Chapter 7) tasks. Then, we address two limitations in existing GCN models in Part 3. We present ConfGCN for controlling influence neighborhood in GCN learning in Chapter 8 and an extension of GCNs for relational graphs in Chapter 9. Finally, we conclude in Chapter 10 by summarizing our contributions and discussing future directions.

# Chapter 2

## Background

In this chapter, we provide an overview of Graph Convolutional Networks (GCNs) which forms a necessary background material required for understanding the subsequent chapters of this work.

### 2.1 Graph Convolutional Networks

#### 2.1.1 Introduction

Convolutional Neural Networks (CNNs) have lead to major breakthroughs in the era of deep learning because of their ability to extract highly expressive features. However, CNNs are restricted to Euclidean data like images and text. Non-Euclidean data like graphs are more expressive and have been used to model a variety of problems. Graph Convolutional Networks (GCNs) address this shortcoming by generalizing CNNs' property of local receptive field, shared weights, and multiple layers to graphs. GCNs have been successfully applied to several domains such as social networks [37], knowledge graphs [171], natural language processing [126, 15], drug discovery [164] and natural sciences [108, 60]. In this chapter. we describe how CNN model for Euclidean graphs can be generalized for non-Euclidean data using Spectral Graph theory [181]. We acknowledge that most of the content of this chapter is adopted from Shuman et al. [175], Defferrard et al. [47], Kipf and Welling [90].

#### 2.1.2 Preliminaries

**Notations:** We denote an undirected and connected graph as  $\mathcal{G} = (\mathcal{V}, \mathcal{E}, \mathbf{W})$ , where  $\mathcal{V}$  refers to the set of nodes ( $N = |\mathcal{V}|$ ),  $\mathcal{E} = \{(u, v) \mid u, v \in \mathcal{V}\}$  indicates the set of edges, and  $\mathbf{W}$  is a weighted adjacency matrix of the graph. If there does not exist an edge between node  $i$  and  $j$  then  $W_{ij}$  is set to 0.

**Graph Signal** refers to a function defined on the vertices of a graph  $\mathcal{G}$ , i.e.,  $f : \mathcal{V} \rightarrow \mathbb{R}$ . For the entire graph, it can be represented as a vector  $\mathbf{x} \in \mathbb{R}^N$ , where  $x_i$  denotes the function value at the  $i^{th}$  vertex. Figure 2.1 shows an illustration of a graph signal over a graph.

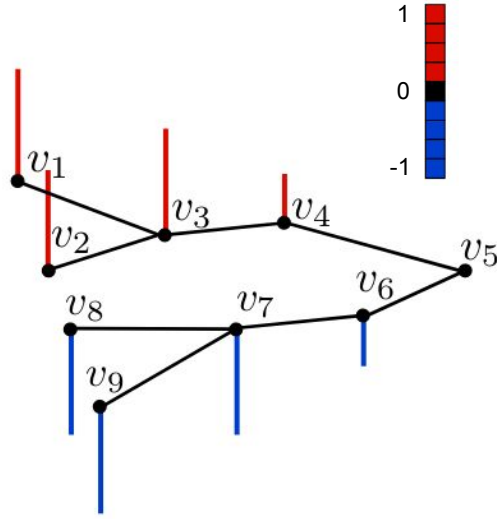


Figure 2.1: An Illustration of Graph signal over a graph. Refer to Section 2.1.2 for details.

**Graph Laplacian ( $\Delta$ )** for any graph signal  $f$  is defined as:

$$(\Delta f)(i) = \sum_{j \in \mathcal{N}_i} W_{i,j}[f(i) - f(j)],$$

where  $\mathcal{N}_i$  is the set of immediate neighbors of vertex  $i$  in  $\mathcal{G}$ . Graph Laplacian measures the difference between  $f$  and its local average. It is small for a smooth signal, i.e., connected vertices have similar values and is large when  $f$  frequently oscillates between connected vertices. Graph Laplacian can be represented as Laplacian matrix, i.e.,

$$\Delta = D - \mathbf{W},$$

where  $D$  is a degree matrix, i.e.,  $D = \text{diag} \left( \sum_{i \neq j} W_{i,j} \right)$ .  $\Delta$  is a real symmetric matrix, therefore, it has a complete set of orthonormal eigenvectors which we denote by  $\{\phi_0, \phi_1, \dots, \phi_{N-1}\}$ . Moreover, all its eigenvalues are real and non-negative, i.e.,  $\lambda_0, \lambda_1, \dots, \lambda_{N-1} \geq 0$ . Further, graph Laplacian ( $\Delta$ ) can be decomposed (Spectral Decomposition) as

$$\Delta = \Phi^T \Lambda \Phi,$$

where  $\Phi = [\phi_0, \phi_1, \dots, \phi_{N-1}]$  and  $\Lambda = \text{diag}(\lambda_0, \lambda_1, \dots, \lambda_{N-1})$ . In the graph setting, eigenvalues and eigenvectors provide a notion of frequency. The eigenvector corresponding to smaller eigenvalues are smoother compared to the eigenvectors with larger eigenvalues. For instance, if we count cross edges ( $\mathcal{Z}_G(f)$ ), i.e., the number of edges connecting vertices with opposite signal value which is defined as:

$$\mathcal{Z}_G(f) = \{e = (i, j) \in \mathcal{E} : f(i)f(j) < 0\},$$

then, we obtain the plot as shown in Figure 2.2. This shows that with the increase in eigenvalue, the number of such edges also increases.

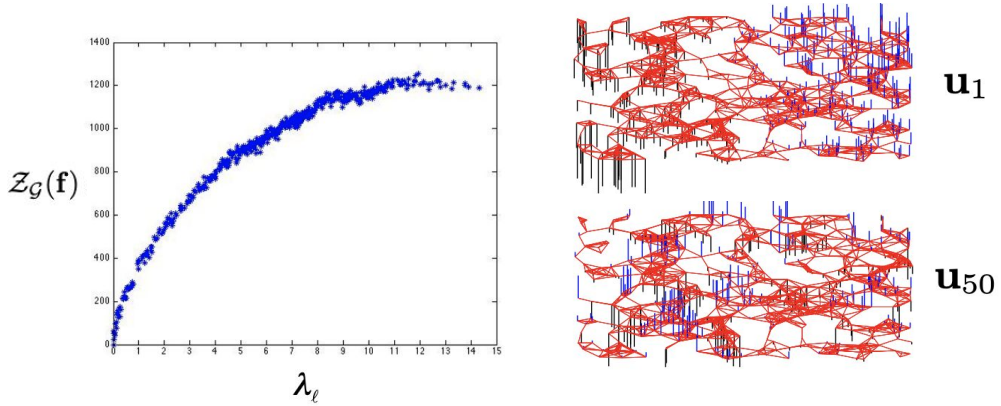


Figure 2.2: **(Left)** Shows change in cross edges with the increase in eigenvalues of graph Laplacian. **(Right)** demonstrates that an eigenvector corresponding to a smaller eigenvalue is smoother compared to the eigenvector corresponding to a larger eigenvalue.

### 2.1.3 Convolution in Euclidean space

Given two functions  $f, g : [-\pi, \pi] \rightarrow \mathbb{R}$ , their convolution is defined as

$$(f \star g)(x) = \int_{-\pi}^{\pi} f(t)g(x-t)dt, \quad (2.1)$$

the above formulation satisfies the following properties:

1. **Shift-invariance** means that the convolution result remains unchanged on translating either of the function, i.e.,  $f(x - x_0) \star g(x) = (f \star g)(x - x_0)$ .
2. **Convolutional Theorem:** Fourier transform diagonalizes the convolution operator which allows it to be computed in the Fourier domain as

$$\widehat{(f \star g)} = \hat{f} \cdot \hat{g},$$

where  $\hat{\cdot}$  indicates the Fourier transform of the function. Similarly, the convolution of two vectors  $\mathbf{f} = (f_1, f_2, \dots, f_n)$  and  $\mathbf{g} = (g_1, g_2, \dots, g_n)$  can be defined as

$$\mathbf{f} \star \mathbf{g} = \Phi(\Phi^T \mathbf{g} \circ \Phi^T \mathbf{f}) \quad (2.2)$$

3. **Computational Efficient:** Using Fast-Fourier Transform (FFT) [65], the Fourier transform can be computed efficiently in  $O(\log n)$ .

#### 2.1.4 Convolution in non-Euclidean space

The definition of convolution as given in Equation 2.1 cannot be directly generalized for the graph setting. This is because translation,  $f(x - t)$  is not defined on graphs. However, by analogy, one can define convolution operation for two vectors  $\mathbf{f}, \mathbf{g} : \mathcal{V} \rightarrow \mathbb{R}^N$  as

$$\begin{aligned} \mathbf{f} \star \mathbf{g} &= \Phi(\Phi^T \mathbf{g} \circ \Phi^T \mathbf{f}) \\ &= \Phi \operatorname{diag}(\hat{g}_1, \hat{g}_2, \dots, \hat{g}_N) \Phi^T \mathbf{f} \\ &= \Phi \hat{g}(\Lambda) \Phi^T \mathbf{f} \\ &= \hat{g}(\Phi \Lambda \Phi^T) \mathbf{f} \\ &= \hat{g}(\Delta) \mathbf{f} \end{aligned}$$

The above formulation unlike for Euclidean space suffers from non-shift invariance. Moreover, the filter coefficients depends on Fourier basis  $\{\phi_1, \phi_2, \dots, \phi_N\}$ , which is expensive to compute  $O(n^2)$  as FFT algorithm is not directly applicable.

To address some of these shortcomings,  $\hat{g}(\Lambda)$  can be defined as a polynomial function of Laplacian eigenvalues, i.e.,

$$\hat{g}(\Lambda) = \sum_{k=1}^K \boldsymbol{\alpha}_k \lambda^k,$$

where  $\boldsymbol{\alpha} = (\alpha_1, \alpha_2, \dots, \alpha_K)^T$  is a vector of filter parameters. This provides localization of convolution operation. The filters represented by  $K^{th}$ -order polynomials of the Laplacian are exactly  $K$ -localized. Moreover, this also reduces the learning complexity to  $O(K)$ , the support size of the filter which is the same complexity as the standard CNNs. The above formulation, however requires  $O(n^2)$  operation as  $\Phi \hat{g}(\Lambda) \Phi^T \mathbf{f}$  involves multiplication with Fourier basis. One solution for this is to use Chebyshev polynomial to parameterize  $\hat{g}(\Delta)$  and recursively compute it from  $\Delta$ , i.e.,

$$\hat{g}(\Delta) = \sum_{k=0}^K \theta_k T_k(\tilde{\Delta}) \mathbf{f}. \quad (2.3)$$

Here,  $T_{k+1}(x) = 2xT_k(x) - T_{k-1}(x)$  with  $T_0(x) = 1$  and  $T_1(x) = x$ .  $T_k(\tilde{\Delta})$  denotes the  $k^{th}$ -order Chebyshev polynomial evaluated at  $\tilde{\Delta} = 2\Delta/\lambda_{max} - I_N$ , Laplacian with eigenvalues constrained to  $[-1, 1]$ . This reduces the time complexity from  $O(n^2)$  to  $O(K|\mathcal{E}|)$  as it involves  $K$  multiplication with sparse  $\Delta$  matrix.

Kipf and Welling [90] defines a first-order approximation of the above formulation by taking  $K = 1$ . This reduces Equation 2.3 to

$$\hat{g}(\Delta)\mathbf{f} = (\theta_0 + \theta_1\tilde{\Delta})\mathbf{f}.$$

Now, approximating  $\lambda_{max} \approx 2$  and taking  $\theta_0 = -\theta_1 = \theta$  gives  $(\theta_0 + \theta_1\tilde{\Delta}) = (\theta - \theta(2\Delta/\lambda_{max} - I_N)) = \theta(I_N - (\Delta - I_N))$ . Thus, the above equation reduces to

$$\hat{g}(\Delta)\mathbf{f} = \theta(I_N + D^{-1/2}AD^{-1/2})\mathbf{f}, \quad (2.4)$$

here,  $\Delta$  is replaced with  $I_N - D^{-1/2}AD^{-1/2}$ , the normalized Laplacian operator. Note that since  $I_N + D^{-1/2}AD^{-1/2}$  has eigenvalues in range  $[0, 2]$ , repeated application of this operator can lead to numerical instabilities. To address this, re-normalization trick is used which replaces  $I_N + D^{-1/2}AD^{-1/2}$  with  $\tilde{D}^{-1/2}\tilde{A}\tilde{D}^{-1/2}$ , where  $\tilde{A} = A + I_N$  and  $\tilde{D}_{ii} = \sum_j \tilde{A}_{ij}$ . Thus, Equation 2.4 is reduced to

$$\hat{g}(\Delta)\mathbf{f} = \theta(\tilde{D}^{-1/2}\tilde{A}\tilde{D}^{-1/2})\mathbf{f}.$$

The above formulation can be generalized for a graph signal  $X \in \mathbb{R}^{N \times d}$  with  $d$ -dimensional feature vector for every node and  $F$  filters as

$$\mathbf{H} = f(\tilde{D}^{-1/2}\tilde{A}\tilde{D}^{-1/2}X\mathbf{W}), \quad (2.5)$$

where  $\mathbf{W} \in \mathbb{R}^{d \times F}$  is a filter parameter,  $f$  is any non-linearity and  $\mathbf{H} \in \mathbb{R}^{N \times F}$  is the convoluted signal matrix. For an undirected graph  $\mathcal{G}$ , the above equation can be re-written as

$$h_v = f \left( \sum_{u \in \mathcal{N}(v)} (\mathbf{W}x_u + b) \right), \quad \forall v \in \mathcal{V}. \quad (2.6)$$

Here,  $\mathcal{N}(v)$  refers to the set of neighbors of  $v$  and  $b \in \mathbb{R}^F$  are learned in a task-specific setting using first-order gradient optimization. In order to capture nodes many hops away, multiple GCN layers may be stacked one on top of another. In particular,  $h_v^{k+1}$ , representation

of node  $v$  after  $k^{th}$  GCN layer can be formulated as

$$h_v^{k+1} = f \left( \sum_{u \in \mathcal{N}(v)} (W^k h_u^k + b^k) \right), \forall v \in \mathcal{V}. \quad (2.7)$$

### 2.1.5 GCNs for Directed and Labeled Graphs

In this section, we consider GCN formulation over graphs where each edge is labeled as well as directed proposed by Marcheggiani and Titov [126]. In this setting, an edge from node  $u$  to  $v$  with label  $l(u, v)$  is denoted as  $(u, v, l(u, v))$ . Based on the assumption that the information in a directed edge need not only propagate along its direction, following Marcheggiani and Titov [126] we define an updated edge set  $\mathcal{E}'$  which expands the original set  $\mathcal{E}$  by incorporating inverse, as well self-loop edges.

$$\mathcal{E}' = \mathcal{E} \cup \{(v, u, l(u, v)^{-1}) \mid (u, v, l(u, v)) \in \mathcal{E}\} \cup \{(u, u, \top) \mid u \in \mathcal{V}\}.$$

$$h_v^{k+1} = f \left( \sum_{u \in \mathcal{N}(v)} (W_{l(u,v)}^k h_u^k + b_{l(u,v)}^k) \right). \quad (2.8)$$

We note that the parameters  $W_{l(u,v)}^k$  and  $b_{l(u,v)}^k$  in this case are edge label specific.

**Incorporating Edge Importance:** In many practical settings, we may not want to give equal importance to all the edges. For example, in case of automatically constructed graphs, some of the edges may be erroneous and we may want to automatically learn to discard them. Edge-wise gating may be used in a GCN to give importance to relevant edges and subdue the noisy ones. Nguyen and Grishman [144], Marcheggiani and Titov [126] used gating for similar reasons and obtained high performance gain. At  $k^{th}$  layer, we compute gating value for a particular edge  $(u, v, l(u, v))$  as:

$$g_{u,v}^k = \sigma \left( h_u^k \cdot \hat{w}_{l(u,v)}^k + \hat{b}_{l(u,v)}^k \right),$$

where,  $\sigma(\cdot)$  is the sigmoid function,  $\hat{w}_{l(u,v)}^k$  and  $\hat{b}_{l(u,v)}^k$  are label specific gating parameters. Thus, gating helps to make the model robust to the noisy labels and directions of the input graphs.

GCN embedding of a node while incorporating edge gating may be computed as follows.

$$h_v^{k+1} = f \left( \sum_{u \in \mathcal{N}(v)} g_{u,v}^k \times (W_{l(u,v)}^k h_u^k + b_{l(u,v)}^k) \right). \quad (2.9)$$

We utilize the GCN formulation for directed and labeled graph with (Equation 2.9) and without edge-wise gating (Equation 2.8) for most of the works in this thesis.

# Part I

## Addressing Sparsity in Knowledge Graphs

# Chapter 3

## Open Knowledge Base Canonicalization using Embeddings and Side Information

### 3.1 Introduction

In this chapter, we present our first solution to address the sparsity problem in Knowledge Graphs. Recent research has resulted in the development of several large *Ontological* Knowledge Bases (KBs), examples include DBpedia [6], YAGO [185], and Freebase [21]. These KBs are called ontological as the knowledge captured by them conform to a fixed ontology, i.e., pre-specified Categories (e.g., *person*, *city*) and Relations (e.g., *mayorOfCity(Person, City)*). Construction of such ontological KBs require significant human supervision. Moreover, due to the need for pre-specification of the ontology, such KB construction methods can't be quickly adapted to new domains and corpora. While other ontological KB construction approaches such as NELL [136] learn from limited human supervision, they still suffers from the quick adaptation bottleneck.

In contrast, Open Information Extraction (OpenIE) methods need neither supervision nor any pre-specified ontology. Given unstructured text documents, OpenIE methods readily extract triples of the form (*noun phrase*, *relation phrase*, *noun phrase*) from them, resulting in the development of large Open Knowledge Bases (Open KBs). Examples of Open KBs include TextRunner [11], ReVerb [55], and OLLIE [40, 170, 129]. While this makes OpenIE methods highly adaptable, they suffer from the following shortcoming: unlike Ontological KBs, the Noun Phrases (NPs) and relation phrases in Open KBs are not *canonicalized*. This results in storage of redundant and ambiguous facts.

Let us explain the need for canonicalization through a concrete example. Please consider the two sentences below.

*Barack Obama was the president of US.*  
*Obama was born in Honolulu.*

Given the two sentences above, an OpenIE method may extract the two triples below and store them in an Open KB.

(Barack Obama, was president of, US)  
(Obama, born in, Honolulu)

Unfortunately, neither such OpenIE methods nor the associated Open KBs have any knowledge that both *Barack Obama* and *Obama* refer to the same person. This can be a significant problem as Open KBs will not return all the facts associated with *Barack Obama* on querying for it. Such KBs will also contain redundant facts, which is undesirable. Thus, there is an urgent need to *canonicalize* noun phrases (NPs) and relations in Open KBs.

In spite of its importance, canonicalization of Open KBs is a relatively unexplored problem. In [62], canonicalization of Open KBs is posed as a clustering problem over *manually* defined feature representations. Given the costs and sub-optimality involved with manual feature engineering, and inspired by recent advances in knowledge base embedding [22, 147], we pose canonicalization of Open KBs as a clustering over *automatically learned* embeddings. We make the following contributions in this chapter.

- We propose Canonicalization using Embeddings and Side Information (CESI), a novel method for canonicalizing Open KBs using learned embeddings. To the best of our knowledge, this is the first approach to use learned embeddings and side information for canonicalizing an Open KB.
- CESI models the problem of noun phrase (NP) and relation phrase canonicalization *jointly* using relevant side information in a principled manner. This is unlike prior approaches where NP and relation phrase canonicalization were performed sequentially.
- We build and experiment with ReVerb45K, a new dataset for Open KB canonicalization. ReVerb45K consists of 20x more NPs than the previous biggest dataset for this task. Through extensive experiments on this and other real-world datasets, we demonstrate CESI’s effectiveness (Section 3.4).

CESI’s source code and datasets used in the chapter are available at <https://github.com/mallabiisc/cesi>.

## 3.2 Related Work

**Entity Linking:** One traditional approach to canonicalizing noun phrases is to map them to an existing KB such as Wikipedia or Freebase. This problem is known as Entity Linking (EL) or Named Entity Disambiguation (NED). Most approaches generate a list of candidate entities for each NP and re-rank them using machine learning techniques. Entity linking has been an active area of research in the NLP community [194, 111, 165]. A major problem with these kind of approaches is that many NPs may refer to new and emerging entities which may not exist in KBs. One approach to resolve these noun phrases is to map them to NIL or an OOKB (Out of Knowledge Base) entity, but the problem still remains as to how to cluster these NIL mentions. Although entity linking is not the best approach to NP canonicalization, we still leverage signals from entity linking systems for improved canonicalization in CESI.

**Canonicalization in Ontological KBs:** Concept Resolver [95] is used for clustering NP mentions in NELL [136]. It makes “one sense per category” assumption which states that a noun phrase can refer to at most one concept in each category of NELL’s ontology. For example, the noun phrase “Apple” can either refer to a company or a fruit, but it can refer to only one company and only one fruit. Another related problem to NP canonicalization is Knowledge Graph Identification [161], where given a noisy extraction graph, the task is to produce a consistent Knowledge Graph (KG) by performing entity resolution, entity classification and link prediction jointly. Pujara et al. [161] incorporate information from multiple extraction sources and use ontological information to infer the most probable knowledge graph using probabilistic soft logic (PSL) [26]. However, both of these approaches require additional information in the form of an ontology of relations, which is not available in the Open KB setting.

**Relation Taxonomy Induction:** SICTF [149] tries to learn relation schemas for different OpenIE relations. It is built up on RESCAL [146], and uses tensor factorization methods to cluster noun phrases into *categories* (such as “person”, “disease”, etc.). We, however, are interested in clustering noun phrases into entities.

There has been relatively less work on the task of relation phrase canonicalization. Some of the early works include DIRT [110], which proposes an unsupervised method for discovering inference rules of the form “*X is the author of Y*  $\approx$  *X wrote Y*” using paths in dependency trees; and the PATTY system [142], which tries to learn subsumption rules among relations (such as *son-of*  $\subset$  *child-of*) using techniques based on frequent itemset mining. These approaches are more focused on finding a taxonomy of relation phrases, while we are looking at finding equivalence between relation phrases.

**Knowledge Base Embedding:** KB embedding techniques such as TransE [22], HolE [147]

try to learn vector space embeddings for entities and relations present in a KB. TransE makes the assumption that for any  $\langle \text{subject}, \text{relation}, \text{object} \rangle$  triple, the relation vector is a translation from the subject vector to the object vector. HolE, on the other hand, uses non-linear operators to model a triple. These embedding methods have been successfully applied for the task of link prediction in KBs. In this work, we build up on HolE while exploiting relevant side information for the task of Open KB canonicalization. We note that, even though KB embedding techniques like HolE have been applied to ontological KBs, CESI might be the first attempt to use them in the context of Open KBs.

**Canonicalizing Open KBs:** The RESOLVER system [224] uses string similarity based features to cluster phrases in TextRunner [11] triples. String similarity features, although being effective, fail to handle synonymous phrases which have completely different surface forms, such as *Myopia* and *Near-sightedness*.

KB-Unify [49] addresses the problem of unifying multiple Ontological and Open KBs into one KB. However, KB-Unify requires a pre-determined sense inventory which is not available in the setting CESI operates.

The most closely related work to ours is [62]. They perform NP canonicalization by performing Hierarchical Agglomerative Clustering (HAC) [190] over manually-defined feature spaces, and subsequently perform relation phrase clustering by using the AMIE algorithm [63]. CESI significantly outperforms this prior method (Section 3.4).

### 3.3 Proposed Approach: CESI

#### 3.3.1 Overview

Overall architecture and dataflow of CESI is shown in Figure 3.1. The input to CESI is an un-canonicalized Open Knowledge Base (KB) with source information for each triple. The output is a list of canonicalized noun and relation phrases, which can be used to identify equivalent entities and relations or canonicalize the KB. CESI achieves this through its three step procedure:

1. **Side Information Acquisition:** The goal of this step is to gather various NP and relation phrase side information for each triple in the input by running several standard algorithms on the source text of the triples. More details can be found in Section 3.3.2.
2. **Embedding NP and Relation Phrases:** In this step, CESI learns specialized vector embeddings for all NPs and relation phrases in the input by making principled use of side information available from the previous step.

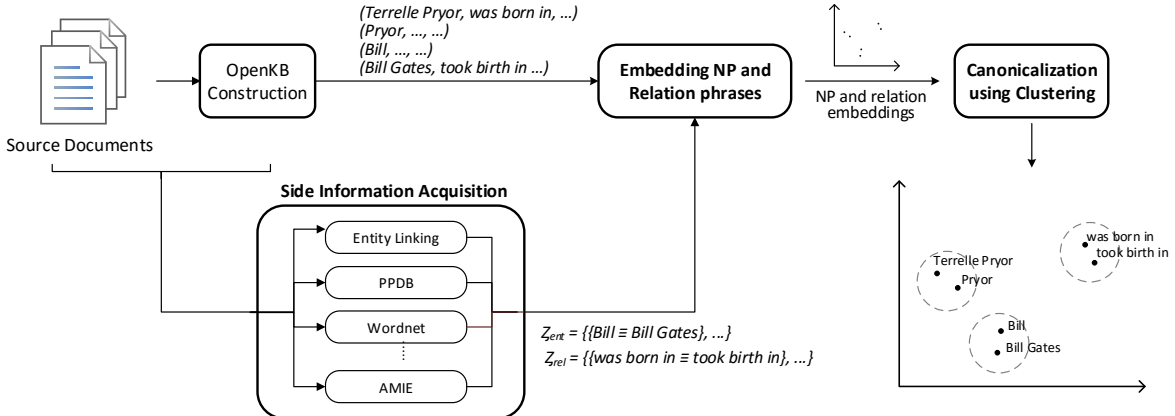


Figure 3.1: Overview of CESI. CESI first acquires side information of noun and relation phrases of Open KB triples. In the second step, it learns embeddings of these NPs and relation phrases while utilizing the side information obtained in previous step. In the third step, CESI performs clustering over the learned embeddings to canonicalize NP and relation phrases. Please see Section 3.3 for more details.

**3. Clustering Embeddings and Canonicalization:** Goal of this step is to cluster the NPs and relation phrases on the basis of their distance in the embedding space. Each cluster represents a specific entity or relation. Based on certain relevant heuristics, we assign a representative to each NP and relation phrase cluster.

Details of different steps of CESI are described next.

### 3.3.2 Side Information Acquisition

Noun and relation phrases in Open KBs often have relevant side information in the form of useful context in the documents from which the triples were extracted. Sometimes, such information may also be present in other related KBs. Previous Open KB canonicalization methods [62] ignored such available side information and performed canonicalization in isolation focusing only on the Open KB triples. CESI attempts to exploit such side information to further improve the performance on this problem. In CESI, we make use of five types of NP side information to get equivalence relations of the form  $e_1 \equiv e_2$  between two entities  $e_1$  and  $e_2$ . Similarly, relation phrase side information is used to derive relation equivalence,  $r_1 \equiv r_2$ . All equivalences are used as soft constraints in later steps of CESI (details in Section 3.3.3).

#### 3.3.2.1 Noun Phrase side Information

In the present version of CESI, we make use of the following five types of NP side information:

1. **Entity Linking:** Given unstructured text, entity linking algorithms identify entity mentions and link them to Ontological KBs such as Wikipedia, Freebase etc. We make use of Stanford CoreNLP entity linker which is based on [182] for getting NP to Wikipedia entity linking. Roughly, in about 30% cases, we get this information for NPs. If two NPs are linked to the same Wikipedia entity, we assume them to be equivalent as per this information. For example, *US* and *America* can get linked to the same Wikipedia entity *United\_States*.
2. **PPDB Information:** We make use of PPDB 2.0 [155], a large collection of paraphrases in English, for identifying equivalence relation among NPs. We first extracted high confidence paraphrases from the dataset while removing duplicates. Then, using union-find, we clustered all the equivalent phrases and randomly assigned a representative to each cluster. Using an index created over the obtained clusters, we find cluster representative for each NP. If two NPs have the same cluster representative then they are considered to be equivalent. NPs not present in the dataset are skipped. This information helps us identifying equivalence between NPs such as *management* and *administration*.
3. **WordNet with Word-sense Disambiguation:** Using word-sense disambiguation [10] with Wordnet [132], we identify possible synsets for a given NP. If two NPs share a common synset, then they are marked as similar as per this side information. For example, *picture* and *image* can get linked to the same synset *visualize.v.01*.
4. **IDF Token Overlap:** NPs sharing infrequent terms give a strong indication of them referring to the same entity. For example, it is very likely for *Warren Buffett* and *Buffett* to refer to the same person. In [62], IDF token overlap was found to be the most effective feature for canonicalization. We assign a score for every pair of NPs based on the standard IDF formula:
$$score_{idf}(n, n') = \frac{\sum_{x \in w(n) \cap w(n')} \log(1 + f(x))^{-1}}{\sum_{x \in w(n) \cup w(n')} \log(1 + f(x))^{-1}}$$

Here,  $w(\cdot)$  for a given NP returns the set of its terms, excluding stop words.  $f(\cdot)$  returns the document frequency for a token.
5. **Morph Normalization:** We make use of multiple morphological normalization operations like tense removal, pluralization, capitalization and others as used in [55] for finding out equivalent NPs. We show in Section 3.4.2.2 that this information helps in improving performance.

### 3.3.2.2 Relation Phrase Side Information

Similar to noun phrases, we make use of PPDB and WordNet side information for relation phrase canonicalization as well. Apart from these, we use the following two additional types of side information involving relation phrases.

1. **AMIE Information:** AMIE algorithm [63] tries to learn implication rules between two relations  $r$  and  $r'$  of the form  $r \Rightarrow r'$ . These rules are detected based on statistical rule mining, for more details refer [62]. It declares two relations  $r$  and  $r'$  to be equivalent if both  $r \Rightarrow r'$  and  $r' \Rightarrow r$  satisfy support and confidence thresholds. AMIE accepts a semi-canonicalized KB as input, i.e., a KB where NPs are already canonicalized. Since this is not the case with Open KBs, we first canonicalized NPs morphologically and then applied AMIE over the NP-canonicalized KB. We chose morphological normalization for this step as such normalization is available for all NPs, and also because we found this side information to be quite effective in large Open KBs.
2. **KBP Information:** Given unstructured text, Knowledge Base Population (KBP) systems detect relations between entities and link them to relations in standard KBs. For example, “*Obama was born in Honolulu*” contains “*was born in*” relation between *Obama* and *Honolulu*, which can be linked to *per:city\_of\_birth* relation in KBs. In CESI, we use Stanford KBP [187] to categorize relations. If two relations fall in the same category, then they are considered equivalent as per this information.

The given list can be further extended based on the availability of other side information. For the experiments in this chapter, we have used the above mentioned NP and relation phrase side information. Some of the equivalences derived from different side information might be erroneous, therefore, instead of using them as hard constraints, we try to use them as supplementary information as described in the next section. Even though side information might be available only for a small fraction of NPs and relation phrases, the hypothesis is that it will result in better overall canonicalization. We find this to be true, as shown in Section 3.4.2.

### 3.3.3 Embedding NP and Relation Phrases

For learning embeddings of NPs and relation phrases in a given Open KB, CESI optimizes Hole’s [147] objective function along with terms for penalizing violation of equivalence conditions from the NP and relation phrase side information. Since the conditions from side information might be spurious, a factor ( $\lambda_{\text{ent/rel},\theta}$ ) is multiplied with each term, which acts as a hyper-parameter and is tuned on a held out validation set. We also keep a constant ( $\lambda_{\text{str}}$ ) with

Hole objective function, to make selective use of structural information from KB for canonicalization. We choose HolE because it is one of the best performing KB embeddings techniques for tasks like link prediction in knowledge graphs. Since KBs store only true triples, we generate negative examples using local closed world heuristic [52]. To keep the rank of true triples higher than the non-existing ones, we use pairwise ranking loss function. The final objective function is described below.

$$\begin{aligned} \min_{\Theta} \quad & \lambda_{str} \sum_{i \in D_+} \sum_{j \in D_-} \max(0, \gamma + \sigma(\eta_j) - \sigma(\eta_i)) \\ & + \sum_{\theta \in \mathcal{C}_{ent}} \frac{\lambda_{ent,\theta}}{|\mathcal{Z}_{ent,\theta}|} \sum_{v,v' \in \mathcal{Z}_{ent,\theta}} \|e_v - e_{v'}\|^2 \\ & + \sum_{\phi \in \mathcal{C}_{rel}} \frac{\lambda_{rel,\phi}}{|\mathcal{Z}_{rel,\phi}|} \sum_{u,u' \in \mathcal{Z}_{rel,\phi}} \|r_u - r_{u'}\|^2 \\ & + \lambda_{reg} \left( \sum_{v \in V} \|e_v\|^2 + \sum_{r \in R} \|e_r\|^2 \right). \end{aligned}$$

The objective function, consists of three main terms, along with one term for regularization. Optimization parameter,  $\Theta = \{e_v\}_{v \in V} \cup \{r_u\}_{u \in R}$ , is the set of all NP ( $e_v$ ) and relation phrase ( $r_u$ )  $d$ -dimensional embeddings, where,  $V$  and  $R$  denote the set of all NPs and relation phrases in the input. In the first term,  $D_+, D_-$  specify the set of positive and negative examples and  $\gamma > 0$  refers to the width of the margin [22]. Further,  $\sigma(\cdot)$  denotes the logistic function and for a triple  $t_i (s, p, o)$ ,  $\eta_i = r_p^T (e_s \star e_o)$ , where  $\star : R^d \times R^d \rightarrow R^d$  is the circular correlation operator defined as follows.

$$[a \star b]_k = \sum_{i=0}^{d-1} a_i b_{(k+i) \bmod d}.$$

The first index of  $(a \star b)$  measures the similarity between  $a$  and  $b$ , while other indices capture the interaction of features from  $a$  and  $b$ , in a particular order. Please refer to [147] for more details.

In the second and third terms,  $\mathcal{C}_{ent}$  and  $\mathcal{C}_{rel}$  are the collection of all types of NP and relation side information available from the previous step (Section 3.3.2), i.e.,  $\mathcal{C}_{ent} = \{\text{Entity Linking, PPDB, ..}\}$  and  $\mathcal{C}_{rel} = \{\text{AMIE, KBP, ..}\}$ . Further,  $\lambda_{ent,\theta}$  and  $\lambda_{rel,\phi}$  denote the constants associated with entity and relation side information. Their value is tuned using grid search on a held out validation set. The set of all equivalence conditions from a particular side information is denoted

by  $\mathcal{Z}_{\text{ent},\theta}$  and  $\mathcal{Z}_{\text{rel},\phi}$ . The rationale behind putting these terms is to allow inclusion of side information while learning embeddings, by enforcing two NPs or relations close together if they are equivalent as per the available side information. Since the side information is available for a fraction of NPs and relation phrases in the input, including these terms in the objective does not slow down the training of embeddings significantly.

The last term adds L2 regularization on the embeddings. All embeddings are initialized by averaging GloVe vectors [157]. We use mini-batch gradient descent for optimization.

### 3.3.4 Clustering Embeddings and Canonicalization

CESI clusters NPs and relation phrases by performing Hierarchical Agglomerative Clustering (HAC) using cosine similarity over the embeddings learned in the previous step (Section 3.3.3). HAC was preferred over other clustering methods because the number of clusters are not known beforehand. Complete linkage criterion is used for calculating the similarity between intermediate clusters as it gives smaller sized clusters, compared to single and average linkage criterion. This is more reasonable for canonicalization problem, where cluster sizes are expected to be small. The threshold value for HAC was chosen based on held out validation dataset.

The time complexity of HAC with complete linkage criterion is  $O(n^2)$  [46]. For scaling up CESI to large knowledge graphs, one may go for modern variants of approximate Hierarchical clustering algorithms [92] at the cost of some loss in performance.

Finally, we decide a representative for each NP and relation phrase cluster. For each cluster, we compute a mean of all elements' embeddings weighted by the frequency of occurrence of each element in the input. NP or relation phrase which lies closest to the weighted cluster mean is chosen as the representative of the cluster.

## 3.4 Experiments

### 3.4.1 Experimental Setup

#### 3.4.1.1 Datasets

Statistics of the three datasets used in the experiments of this chapter are summarized in Table 3.1. We present below brief summary of each dataset.

- 1. Base and Ambiguous Datasets:** We obtained the Base and Ambiguous datasets from the authors of [62]. Base dataset was created by collecting triples containing 150 sampled Freebase entities that appear with at least two aliases in ReVerb Open KB. The same dataset was further enriched with mentions of homonym entities to create the Ambiguous dataset. Please see [62] for more details.

Datasets	# Gold Entities	#NPs	#Relations	#Triples
Base	150	290	3K	9K
Ambiguous	446	717	11K	37K
ReVerb45K	7.5K	15.5K	22K	45K

Table 3.1: Details of datasets used. ReVerb45K is the new dataset we propose in this chapter. Please see Section 3.4.1.1 for details.

2. **ReVerb45K:** This is the new Open KB canonicalization dataset we propose in this work. ReVerb45K is a significantly extended version of the Ambiguous dataset, containing more than 20x NPs. ReVerb45K is constructed by intersecting information from the following three sources: ReVerb Open KB [55], Freebase entity linking information from [61], and Clueweb09 corpus [31]. Firstly, for every triple in ReVerb, we extracted the source text from Clueweb09 corpus from which the triple was generated. In this process, we rejected triples for which we could not find any source text. Then, based on the entity linking information from [61], we linked all subjects and objects of triples to their corresponding Freebase entities. If we could not find high confidence linking information for both subject and object in a triple, then it was rejected. Further, following the dataset construction procedure adopted by [62], we selected triples associated with all Freebase entities with at least two aliases occurring as subject in our dataset. Through these steps, we obtained 45K high-quality triples which we used for evaluation. We call this resulting dataset ReVerb45K.

In contrast to Base and Ambiguous datasets, the number of entities, NPs and relation phrases in ReVerb45K are significantly larger. Please see Table 3.1 for a detailed comparison. This better mimics real-world KBs which tend to be sparse with very few edges per entity, as also observed by [22].

For getting test and validation set for each dataset, we randomly sampled 20% Freebase entities and called all the triples associated with them as validation set and rest was used as the test set.

### 3.4.1.2 Evaluation Metrics

Following [62], we use macro-, micro- and pairwise metrics for evaluating Open KB canonicalization methods. We briefly describe below these metrics for completeness. In all cases,  $C$  denotes the clusters produced by the algorithm to be evaluated, and  $E$  denotes the gold

standard clusters. In all cases, F1 measure is given as the harmonic mean of precision and recall.

**Macro:** Macro precision ( $P_{\text{macro}}$ ) is defined as the fraction of pure clusters in  $C$ , i.e., clusters in which all the NPs (or relations) are linked to the same gold entity (or relation). Macro recall ( $R_{\text{macro}}$ ) is calculated like macro precision but with the roles of  $E$  and  $C$  interchanged.

$$\begin{aligned} P_{\text{macro}}(C, E) &= \frac{|\{c \in C : \exists e \in E : e \supseteq c\}|}{|C|} \\ R_{\text{macro}}(C, E) &= P_{\text{macro}}(E, C) \end{aligned}$$

**Micro:** Micro precision ( $P_{\text{micro}}$ ) is defined as the purity of  $C$  clusters [124] based on the assumption that the most frequent gold entity (or relation) in a cluster is correct. Micro recall ( $R_{\text{micro}}$ ) is defined similarly as macro recall.

$$\begin{aligned} P_{\text{micro}}(C, E) &= \frac{1}{N} \sum_{c \in C} \max_{e \in E} |c \cap e| \\ R_{\text{micro}}(C, E) &= P_{\text{micro}}(E, C) \end{aligned}$$

**Pairwise:** Pairwise precision ( $P_{\text{pair}}$ ) is measured as the ratio of the number of hits in  $C$  to the total possible pairs in  $C$ . Whereas, pairwise recall ( $R_{\text{pair}}$ ) is the ratio of number of hits in  $C$  to all possible pairs in  $E$ . A pair of elements in a cluster in  $C$  produce a hit if they both refer to the same gold entity (or relation).

$$\begin{aligned} P_{\text{pair}}(C, E) &= \frac{\sum_{c \in C} |\{(v, v') \in e, \exists e \in E, \forall (v, v') \in c\}|}{\sum_{c \in C} |c| C_2} \\ R_{\text{pair}}(C, E) &= \frac{\sum_{c \in C} |\{(v, v') \in e, \exists e \in E, \forall (v, v') \in c\}|}{\sum_{e \in E} |e| C_2} \end{aligned}$$

Let us illustrate these metrics through a concrete NP canonicalization example shown in Figure 3.2. In this Figure, we can see that only  $c_2$  and  $c_3$  clusters in  $C$  are pure because they contain mentions of only one entity, and hence,  $P_{\text{macro}} = \frac{2}{3}$ . On the other hand, we have  $e_1$  and  $e_3$  as pure clusters if we interchange the roles of  $E$  and  $C$ . So,  $R_{\text{macro}} = \frac{2}{3}$  in this case. For micro precision, we can see that *America*, *New York*, and *California* are the most frequent gold entities in  $C$  clusters. Hence,  $P_{\text{micro}} = \frac{6}{7}$ . Similarly,  $R_{\text{micro}} = \frac{6}{7}$  in this case. For pairwise analysis, we need to first calculate the number of hits in  $C$ . In  $c_1$  we have 3 possible pairs out of which only 1, (*America*, *USA*) is a hit as they belong to same gold cluster  $e_1$ . Similarly, we

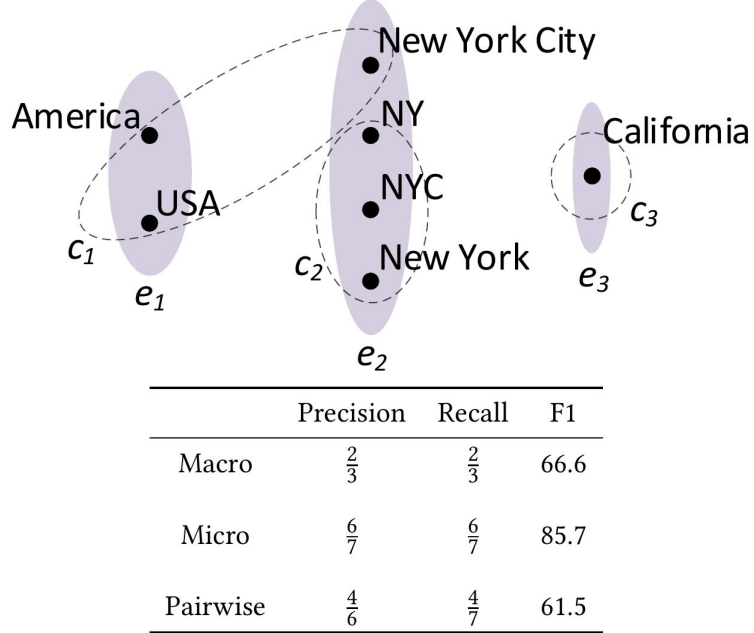


Figure 3.2: Top: Illustrative example for different evaluation metrics.  $e_i$  denotes actual clusters, whereas  $c_i$  denotes predicted clusters. Bottom: Metric results for the above example. Please see Section 3.4.1.2 for details.

have 3 hits in  $c_2$  and 0 hits in  $c_3$ . Hence,  $P_{\text{pair}} = \frac{4}{6}$ . To compute  $R_{\text{pair}}$ , we need total number of pairwise decisions in  $E$ , which is  $1 + 6 + 0$ , thus,  $R_{\text{pair}} = \frac{4}{7}$ . All the results are summarized in Table 3.2.

For evaluating NP canonicalization, we use Macro, Micro and Pairwise F1 score. However, in the case of relations, where gold labels are not available, we use macro, micro and pairwise precision values based on the scores given by human judges.

### 3.4.1.3 Methods Compared

**Noun Phrase Canonicalization:** For NP canonicalization, CESI has been compared against the following methods:

- **Morphological Normalization:** As used in [55], this involves applying simple normalization operations like removing tense, pluralization, capitalization etc. over NPs and relation phrases.
- **Paraphrase Database (PPDB):** Using PPDB 2.0 [155], we clustered two NPs together if they happened to share a common paraphrase. NPs which could not be found in PPDB are put into singleton clusters.

- **Entity Linking:** Since the problem of NP canonicalization is closely related to entity linking, we compare our method against Stanford CoreNLP Entity Linker [182]. Two NPs linked to the same entity are clustered together.
- **Galárraga-IDF** [62]: IDF Token Overlap was the best performing method proposed in [62] for NP canonicalization. In this method, IDF token similarity is defined between two NPs as in Section 3.3.2.1, and HAC is used to cluster the mentions.
- **Galárraga-StrSim** [62]: This method is similar to Galarraga-IDF, but with similarity metric being the Jaro-Winkler [213] string similarity measure.
- **Galárraga-Attr** [62]: Again, this method is similar to the Galarraga-IDF, except that Attribute Overlap is used as the similarity metric between two NPs in this case. Attribute for a NP  $n$ , is defined as the set of relation-NP pairs which co-occur with  $n$  in the input triples. Attribute overlap similarity between two NPs, is defined as the Jaccard coefficient of the set of attributes:

$$f_{\text{attr}}(n, n') = \frac{|A \cap A'|}{|A \cup A'|}$$

where,  $A$  and  $A'$  denote the set of attributes associated with  $n$  and  $n'$ .

Since canonicalization methods using above similarity measures were found to be most effective in [62], even outperforming Machine Learning-based alternatives, we consider these three baselines as representatives of state-of-the-art in Open KB canonicalization.

- **GloVe**: In this scheme, each NP and relation phrase is represented by a 300 dimensional GloVe embedding [157] trained on Wikipedia 2014 and Gigaword 5 [153] datasets with 400k vocabulary size. Word vectors were averaged together to get embeddings for multi-word phrases. These GloVE embeddings were then clustered for final canonicalization.
- **Hole**: In this method, embeddings of NPs and relation phrases in an Open KB are obtained by applying Hole [147] over the Open KB. These embeddings are then clustered to obtain the final canonicalized groupings. Based on the initialization of embeddings, we differentiate between **Hole(Random)** and **Hole(GloVe)**.
- **CESI**: This is the method proposed in this chapter, please see Section 3.3 for more details.

**Hyper-parameters:** Following [62], we used Hierarchical Agglomerative Clustering (HAC) as the default clustering method across all methods (wherever necessary). For all methods, grid search over the hyperparameter space was performed, and results for the best performing setting are reported. This process was repeated for each dataset.

### 3.4.1.4 Relation Phrase Canonicalization

AMIE [63] was found to be effective for relation phrase canonicalization in [62]. We thus consider AMIE<sup>1</sup> as the state-of-the-art baseline for relation phrase canonicalization and compare against CESI. We note that AMIE requires NPs of the input Open KB to be already canonicalized. In all our evaluation datasets, we already have *gold* NP canonicalization available. We provide this gold NP canonicalization information as input to AMIE. Please note that CESI doesn't require such pre-canonicalized NP as input, as it performs *joint* NP and relation phrase canonicalization. Moreover, providing gold NP canonicalization information to AMIE puts CESI at a disadvantage. We decided to pursue this choice anyways in the interest of stricter evaluation. However, in spite of starting from this disadvantageous position, CESI significantly outperforms AMIE in relation phrase canonicalization, as we will see in Section 3.4.2.1.

For evaluating performance of both algorithms, we randomly sampled 25 non-singleton relation clusters for each of the three datasets and gave them to five different human evaluators<sup>2</sup> for assigning scores to each cluster. The setting was kept blind, i.e., identity of the algorithm producing a cluster was not known to the evaluators. Based on the average of evaluation scores, precision values were calculated. Only non-singleton clusters were sampled, as singleton clusters will always give a precision of one.

## 3.4.2 Results

In this section, we evaluate the following questions.

- Q1. Is CESI effective in Open KB canonicalization? (Section 3.4.2.1)
- Q2. What is the effect of side information in CESI's performance? (Section 3.4.2.2)
- Q3. Does addition of entity linking side information degrade CESI's ability to canonicalize unlinked NPs (i.e., NPs missed by the entity linker)? (Section 3.4.2.3)

Finally, in Section 3.4.2.4, we present qualitative examples and discussions.

### 3.4.2.1 Evaluating Effectiveness of CESI in Open KB Canonicalization

**Noun Phrase Canonicalization:** Results for NP canonicalization are summarized in Table 3.2. Overall, we find that CESI performs well consistently across the datasets. Morphological Normalization failed to give competitive performance in presence of homonymy. PPDB, in spite of being a vast reservoir of paraphrases, lacks information about real-world entities like people,

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<sup>1</sup>We use support and confidence values of 2 and 0.2 for all the experiments in this chapter.

<sup>2</sup>Authors did not participate in this evaluation.

Method	Base Dataset			Ambiguous Dataset			ReVerb45K			Row Average
	Macro	Micro	Pair.	Macro	Micro	Pair.	Macro	Micro	Pair.	
Morph Norm	58.3	88.3	83.5	49.1	57.2	70.9	1.4	77.7	75.1	62.3
PPDB	42.4	46.9	32.2	37.3	60.2	69.3	46.0	45.4	64.2	49.3
EntLinker	54.9	65.1	75.2	49.7	83.2	68.8	62.8	81.8	80.4	69.1
Galárraga-StrSim	88.2	96.5	97.7	66.6	85.3	82.2	69.9	51.7	0.5	70.9
Galárraga-IDF	94.8	97.9	98.3	67.9	82.9	79.3	71.6	50.8	0.5	71.5
Galárraga-Attr	76.1	51.4	18.1	<b>82.9</b>	27.7	8.4	<b>75.1</b>	20.1	0.2	40.0
GloVe	95.7	97.2	91.1	65.9	89.9	90.1	56.5	82.9	75.3	82.7
HolE (Random)	69.5	91.3	86.6	53.3	85.0	75.1	5.4	74.6	50.9	65.7
HolE (GloVe)	75.2	93.6	89.3	53.9	85.4	76.7	33.5	75.8	51.0	70.4
CESI	<b>98.2</b>	<b>99.8</b>	<b>99.9</b>	66.2	<b>92.4</b>	<b>91.9</b>	62.7	<b>84.4</b>	<b>81.9</b>	<b>86.3</b>

Table 3.2: NP Canonicalization Results. CESI outperforms all other methods across datasets (Best in 7 out of 9 cases. Section 3.4.2.1)

places etc. Therefore, its performance remained weak throughout all datasets. Entity linking methods make use of contextual information from source text of each triple to link a NP to a KB entity. But their performance is limited because they are restricted by the entities in KB.

String similarity also gave decent performance in most cases but since they solely rely on surface form of NPs, they are bound to fail with NPs having dissimilar mentions.

Methods such as Galárraga-IDF, Galárraga-StrSim, and Galárraga-Attr performed poorly on ReVerb45K. Although, their performance is considerably better on the other two datasets. This is because of the fact that in contrast to Base and Ambiguous datasets, ReVerb45K has considerably large number of entities and comparatively fewer triples (Table 3.1). Galárraga-IDF token overlap is more likely to put two NPs together if they share an uncommon token, i.e., one with high IDF value. Hence, accuracy of the method relies heavily on the quality of document frequency estimates which may be quite misleading when we have smaller number of triples. Similar is the case with Galárraga-Attr which decides similarity of NPs based on the set of shared attributes. Since, attributes for a NP is defined as a set of relation-NP pairs occurring with it across all triples, sparse data also results in poor performance for this method.

GloVe captures semantics of NPs and unlike string similarity it doesn't rely on the surface form of NPs. Therefore, its performance has been substantial across all the datasets. HolE captures structural information from the given triples and uses it for learning embeddings. Through our experiments, we can see that solely structural information from KB is quite effective for NP canonicalization. CESI performs the best across the datasets in 7 out of the 9 settings, as it incorporates the strength of all the listed methods. The superior performance of CESI compared to HolE clearly indicates that the side information is indeed helpful for

	Macro Precision	Micro Precision	Pairwise Precision	Induced Relation Clusters
<b>Base Dataset</b>				
AMIE	42.8	63.6	43.0	7
CESI	<b>88.0</b>	<b>93.1</b>	<b>88.1</b>	<b>210</b>
<b>Ambiguous Dataset</b>				
AMIE	55.8	64.6	23.4	46
CESI	<b>76.0</b>	<b>91.9</b>	<b>80.9</b>	<b>952</b>
<b>ReVerb45K</b>				
AMIE	69.3	84.2	66.2	51
CESI	<b>77.3</b>	<b>87.8</b>	<b>72.6</b>	<b>2116</b>

Table 3.3: Relation canonicalization results. Compared to AMIE, CESI canonicalizes more number of relation phrases at higher precision. Please see Section 3.4.2.1 for details.

canonicalization task. Results of GloVe, HolE and CESI suggest that embeddings based method are much more effective for Open KB canonicalization.

**Relation Phrase Canonicalization** Results for relation phrase canonicalization are presented in Table 3.3. For all experiments, in spite of using quite low values for minimum support and confidence, AMIE was unable to induce any reasonable number of non-singleton clusters (e.g., only 51 clusters out of the 22K relation phrases in the ReVerb45K dataset). For relation canonicalization experiments, AMIE was evaluated on gold NP canonicalized data as the algorithm requires NPs to be already canonicalized. CESI, on the other hand, was tested on all the datasets without making use of gold NP canonicalization information.

Based on the results in Table 3.3, it is quite evident that AMIE induces too few relation clusters to be of value in practical settings. On the other hand, CESI consistently performs well across all the datasets and induces significantly larger number of clusters.

#### 3.4.2.2 Effect of Side Information in CESI

In this section, we evaluate the effect of various side information in CESI’s performance. For this, we evaluated the performances of various versions of CESI, each one of them obtained by ablating increasing amounts of side information from the full CESI model. Experimental results comparing these ablated versions on the ReVerb45K are presented in Figure 3.3a. From this figure, we observe that while macro performance benefits most from different forms of

	Macro F1	Micro F1	Pairwise F1
CESI	81.7	87.6	81.5
CESI w/o EL	81.3	87.3	80.7

Table 3.4: CESI’s performance in canonicalizing unlinked NPs, with and without Entity Linking (EL) side information, in the ReVerb45K dataset. We observe that CESI does not overfit to EL side information, and thereby helps prevent performance degradation in unlinked NP canonicalization (in fact it even helps a little). Please see Section 3.4.2.3 for details.

side information, micro and pairwise performance also show increased performance in the presence of various side information. This validates one of the central thesis of this chapter: side information, along with embeddings, can result in improved Open KB canonicalization.

### 3.4.2.3 Effect of Entity Linking Side Information on Unlinked NP

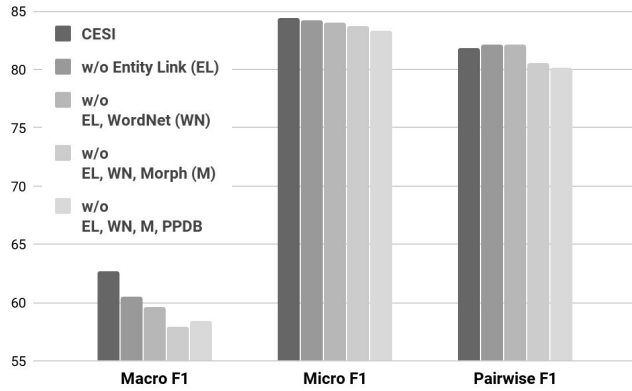
From experiments in Section 3.4.2.2, we find that Entity Linking (EL) side information (see Section 3.3.2.1) is one of the most useful side information that CESI exploits. However, such side information is not available in case of unlinked NPs, i.e., NPs which were not linked by the entity linker. So, this naturally raises the following question: does CESI overfit to the EL side information and ignore the unlinked NPs, thereby resulting in poor canonicalization of such unlinked NPs?

In order to evaluate this question, we compared CESI’s performance on unlinked NPs in the ReVerb45K dataset, with and without EL side information. We note that triples involving unlinked NPs constitute about 25% of the entire dataset. Results are presented in Table 3.4. From this table, we observe that CESI doesn’t overfit to EL side information, and it selectively uses such information when appropriate (i.e., for linked NPs). Because of this robust nature, presence of EL side information in CESI doesn’t have an adverse effect on the unlinked NPs, in fact there is a small gain in performance.

### 3.4.2.4 Qualitative Evaluation

Figure 3.3b shows some of the NP and relation phrase clusters detected by CESI in ReVerb45K dataset. These results highlight the efficacy of algorithm in canonicalizing non-trivial NPs and relation phrases. The figure shows t-SNE [197] visualization of NP and relation phrase (marked in ‘< … >’) embeddings for a few examples. We can see that the learned embeddings are actually able to capture equivalence of NPs and relation phrases. The algorithm is able to correctly embed *Prozac*, *Sarafem* and *Fluoxetine* together (different names of the same drug), despite their having completely different surface forms.

Figure 3.3b also highlights the failures of CESI. For example, *Toyota* and *Nissan* have been



(a) Performance comparison of various side information-ablated versions of CESI for NP phrase canonicalization in the ReVerb45K dataset. Overall, side information helps CESI improve performance. Please see Section 3.4.2.2 for details.



(b) t-SNE visualization of NP and relation embeddings learned by CESI for ReVerb45K dataset. We observe that CESI is able to induce non-trivial canonical clusters. More details in Sec. 3.4.2.4.

embedded together although the two being different companies. Another case is with *Pablo* and *Juan Pablo Angel*, which refer to different entities. The latter case can be avoided by keeping track of the source domain type information of each NP for disambiguation. In this if we know that *Juan Pablo Angel* has come from *SPORTS* domain, whereas *Pablo* has come from a different domain then we can avoid putting them together. We tried using DMOZ [180] dataset, which provide mapping from URL domain to their categories, for handling such errors. But, because of poor coverage of URLs in DMOZ dataset, we couldn't get significant improvement in canonicalization results. We leave this as a future work.

### 3.5 Conclusion

Canonicalizing Open Knowledge Bases (KBs) is an important but underexplored problem. In this chapter, we proposed CESI, a novel method for canonicalizing Open KBs using learned embeddings and side information. CESI solves a joint objective to learn noun and relation phrase embeddings, while utilizing relevant side information in a principled manner. These learned embeddings are then clustered together to obtain canonicalized noun and relation phrase clusters. We also propose ReVerb45K, a new and larger dataset for Open KB canonicalization. Through extensive experiments on this and other real-world datasets, we demonstrate CESI's effectiveness over state-of-the-art baselines. CESI's source code and all data used in this work are publicly available.

# Chapter 4

## Improving Distantly-Supervised Relation Extraction using Graph Convolutional Networks and Side Information

### 4.1 Introduction

The construction of large-scale Knowledge Bases (KBs) like Freebase [21] and Wikidata [206] has proven to be useful in many natural language processing (NLP) tasks like question-answering, web search, etc. However, these KBs are not exhaustive. Relation Extraction (RE) attempts to fill this gap by extracting semantic relationships between entity pairs from plain text. This task can be modeled as a simple classification problem after the entity pairs are specified. Formally, given an entity pair  $(e_1, e_2)$  from the KB and an entity annotated sentence (or instance), we aim to predict the relation  $r$ , from a predefined relation set, that exists between  $e_1$  and  $e_2$ . If no relation exists, we simply label it  $NA$ .

Most supervised relation extraction methods require large labeled training data which is expensive to construct. Distant Supervision (DS) [133] helps with the construction of this dataset automatically, under the assumption that if two entities have a relationship in a KB, then all sentences mentioning those entities express the same relation. While this approach works well in generating large amounts of training instances, the DS assumption does not hold in all cases. Riedel et al. [167], Hoffmann et al. [78], Surdeanu et al. [187] propose multi-instance based learning to relax this assumption. However, they use NLP tools to extract features, which can be noisy.

Recently, neural models have demonstrated promising performance on RE. Zeng et al. [229, 230] employ Convolutional Neural Networks (CNN) to learn representations of instances. For alleviating noise in distant supervised datasets, attention has been utilized by [113, 80]. Syntactic information from dependency parses has been used by [133, 74] for capturing long-range dependencies between tokens. Recently proposed Graph Convolution Networks (GCN) [47] have been effectively employed for encoding this information [126, 15]. However, all the above models rely only on the noisy instances from distant supervision for RE.

Relevant side information can be effective for improving RE. For instance, in the sentence, *Microsoft was started by Bill Gates.*, the type information of *Bill Gates (person)* and *Microsoft (organization)* can be helpful in predicting the correct relation *funderOfCompany*. This is because every relation constrains the type of its target entities. Similarly, relation phrase “*was started by*” extracted using Open Information Extraction (Open IE) methods can be useful, given that the aliases of relation *funderOfCompany*, e.g., *founded*, *co-founded*, etc., are available. KBs used for DS readily provide such information which has not been completely exploited by current models.

In this chapter, we propose RESIDE, a novel distant supervised relation extraction method which utilizes additional supervision from KB through its neural network based architecture. RESIDE makes principled use of entity type and relation alias information from KBs, to impose soft constraints while predicting the relation. It uses encoded syntactic information obtained from Graph Convolution Networks (GCN), along with embedded side information, to improve neural relation extraction. Our contributions can be summarized as follows:

- We propose RESIDE, a novel neural method which utilizes additional supervision from KB in a principled manner for improving distant supervised RE.
- RESIDE uses Graph Convolution Networks (GCN) for modeling syntactic information and has been shown to perform competitively even with limited side information.
- Through extensive experiments on benchmark datasets, we demonstrate RESIDE’s effectiveness over state-of-the-art baselines.

RESIDE’s source code and datasets used in the chapter are available at <http://github.com/mallabiisc/RESIDE>.

## 4.2 Related Work

**Distant supervision:** Relation extraction is the task of identifying the relationship between two entity mentions in a sentence. In supervised paradigm, the task is considered as a multi-class classification problem but suffers from lack of large labeled training data. To address this

limitation, [133] propose distant supervision (DS) assumption for creating large datasets, by heuristically aligning text to a given Knowledge Base (KB). As this assumption does not always hold true, some of the sentences might be wrongly labeled. To alleviate this shortcoming, Riedel et al. [167] relax distant supervision for multi-instance single-label learning. Subsequently, for handling overlapping relations between entities [78, 187] propose multi-instance multi-label learning paradigm.

**Neural Relation Extraction:** The performance of the above methods strongly rely on the quality of hand engineered features. Zeng et al. [229] propose an end-to-end CNN based method which could automatically capture relevant lexical and sentence level features. This method is further improved through piecewise max-pooling by [230]. Lin et al. [113], Nagarajan et al. [141] use attention [7] for learning from multiple valid sentences. We also make use of attention for learning sentence and bag representations.

Dependency tree based features have been found to be relevant for relation extraction [133]. He et al. [74] use them for getting promising results through a recursive tree-GRU based model. In RESIDE, we make use of recently proposed Graph Convolution Networks [47, 91], which have been found to be quite effective for modelling syntactic information [126, 144, 198].

**Side Information in RE:** Entity description from KB has been utilized for RE [81], but such information is not available for all entities. Type information of entities has been used by Ling and Weld [114], Liu et al. [117] as features in their model. Yaghoobzadeh et al. [219] also attempt to mitigate noise in DS through their joint entity typing and relation extraction model. However, KBs like Freebase readily provide reliable type information which could be directly utilized. In our work, we make principled use of entity type and relation alias information obtained from KB. We also use unsupervised Open Information Extraction (Open IE) methods [128, 4], which automatically discover possible relations without the need of any predefined ontology, which is used as a side information as defined in Section 4.3.3.

## 4.3 Proposed Method: RESIDE

### 4.3.1 Overview

In multi-instance learning paradigm, we are given a bag of sentences (or instances)  $\{s_1, s_2, \dots, s_n\}$  for a given entity pair, the task is to predict the relation between them. RESIDE consists of three components for learning a representation of a given bag, which is fed to a softmax classifier. We briefly present the components of RESIDE below. Each component will be described in detail in the subsequent sections. The overall architecture of RESIDE is shown in Figure 4.1.

1. **Syntactic Sentence Encoding:** RESIDE uses a Bi-GRU over the concatenated positional

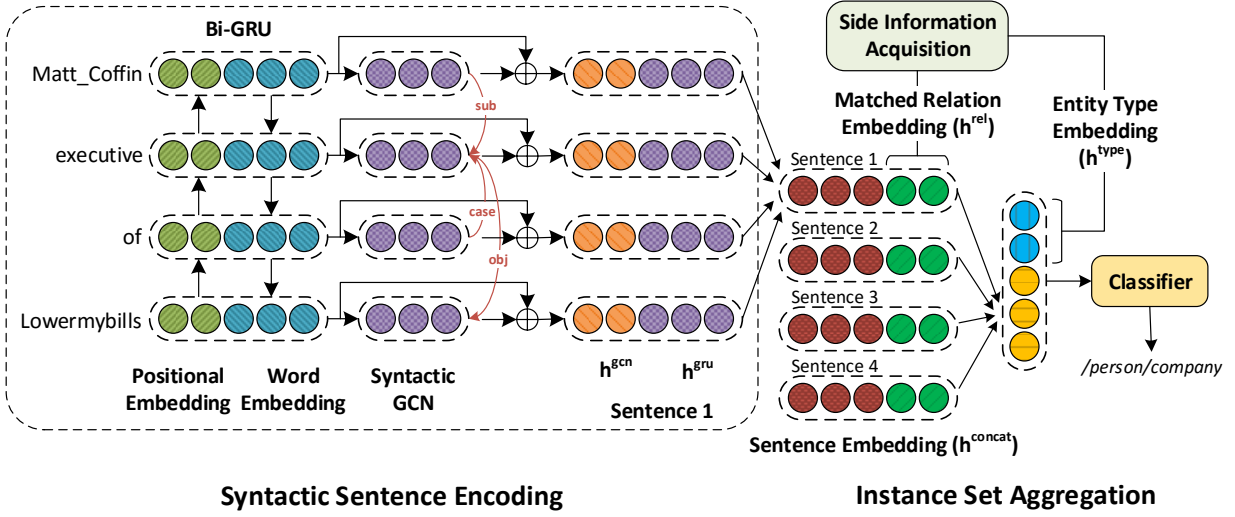


Figure 4.1: Overview of RESIDE. RESIDE first encodes each sentence in the bag by concatenating embeddings (denoted by  $\oplus$ ) from Bi-GRU and Syntactic GCN for each token, followed by word attention. Then, sentence embedding is concatenated with relation alias information, which comes from the Side Information Acquisition Section (Figure 4.2), before computing attention over sentences. Finally, bag representation with entity type information is fed to a softmax classifier. Please see Section 4.3 for more details.

and word embedding for encoding the local context of each token. For capturing long-range dependencies, GCN over dependency tree is employed and its encoding is appended to the representation of each token. Finally, attention over tokens is used to subdue irrelevant tokens and get an embedding for the entire sentence. More details in Section 4.3.2.

2. **Side Information Acquisition:** In this module, we use additional supervision from KBs and utilize Open IE methods for getting relevant side information. This information is later utilized by the model as described in Section 4.3.3.
3. **Instance Set Aggregation:** In this part, sentence representation from syntactic sentence encoder is concatenated with the *matched relation embedding* obtained from the previous step. Then, using attention over sentences, a representation for the entire bag is learned. This is then concatenated with *entity type embedding* before feeding into the softmax classifier for relation prediction. Please refer to Section 4.3.4 for more details.

Below, we provide the detailed description of the components of RESIDE.

### 4.3.2 Syntactic Sentence Encoding

For each sentence in the bag  $s_i$  with  $m$  tokens  $\{w_1, w_2, \dots, w_m\}$ , we first represent each token by  $k$ -dimensional GloVe embedding [157]. For incorporating relative position of tokens with respect to target entities, we use  $p$ -dimensional position embeddings, as used by [229]. The combined token embeddings are stacked together to get the sentence representation  $\mathcal{H} \in \mathbb{R}^{m \times (k+2p)}$ . Then, using Bi-GRU [38] over  $\mathcal{H}$ , we get the new sentence representation  $\mathcal{H}^{gru} \in \mathbb{R}^{m \times d_{gru}}$ , where  $d_{gru}$  is the hidden state dimension. Bi-GRUs have been found to be quite effective in encoding the context of tokens in several tasks [188, 69].

Although Bi-GRU is capable of capturing local context, it fails to capture long-range dependencies which can be captured through dependency edges. Prior works [133, 74] have exploited features from syntactic dependency trees for improving relation extraction. Motivated by their work, we employ Syntactic Graph Convolution Networks for encoding this information. For a given sentence, we generate its dependency tree using Stanford CoreNLP [125]. We then run GCN over the dependency graph and use Equation 2.9 for updating the embeddings, taking  $\mathcal{H}^{gru}$  as the input. Since dependency graph has 55 different edge labels, incorporating all of them over-parameterizes the model significantly. Therefore, following [126, 144, 198] we use only three edge labels based on the direction of the edge  $\{\text{forward } (\rightarrow), \text{backward } (\leftarrow), \text{self-loop } (\top)\}$ . We define the new edge label  $L_{uv}$  for an edge  $(u, v, l_{uv})$  as follows:

$$L_{uv} = \begin{cases} \rightarrow & \text{if edge exists in dependency parse} \\ \leftarrow & \text{if edge is an inverse edge} \\ \top & \text{if edge is a self-loop} \end{cases}$$

For each token  $w_i$ , GCN embedding  $h_{i_{k+1}}^{gcn} \in \mathbb{R}^{d_{gcn}}$  after  $k^{th}$  layer is defined as:

$$h_{i_{k+1}}^{gcn} = f \left( \sum_{u \in \mathcal{N}(i)} g_{iu}^k \times (W_{L_{iu}}^k h_{u_k}^{gcn} + b_{L_{iu}}^k) \right).$$

Here,  $g_{iu}^k$  denotes edgewise gating as defined in Equation 2.9 and  $L_{iu}$  refers to the edge label defined above. We use ReLU as activation function  $f$ , throughout our experiments. The syntactic graph encoding from GCN is appended to Bi-GRU output to get the final token representation,  $h_i^{concat}$  as  $[h_i^{gru}; h_{i_{k+1}}^{gcn}]$ . Since, not all tokens are equally relevant for RE task, we calculate the degree of relevance of each token using attention as used in [80]. For token  $w_i$  in

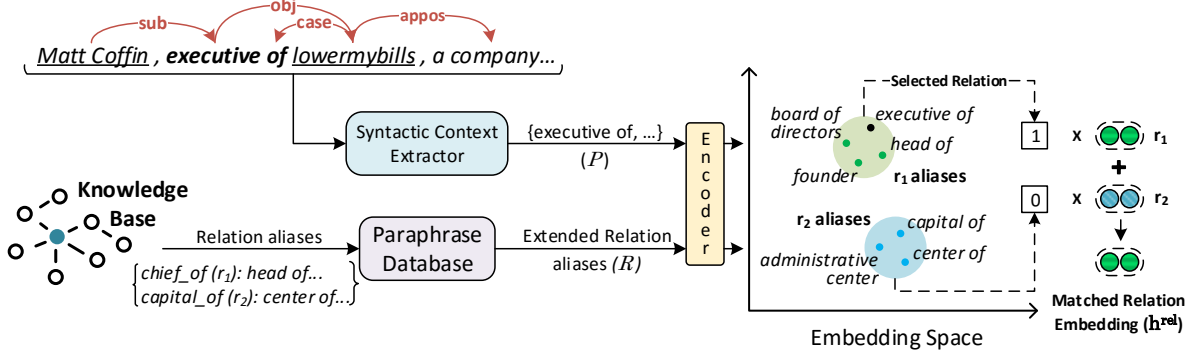


Figure 4.2: Relation alias side information extraction for a given sentence. First, Syntactic Context Extractor identifies relevant relation phrases  $\mathcal{P}$  between target entities. They are then matched in the embedding space with the extended set of relation aliases  $\mathcal{R}$  from KB. Finally, the relation embedding corresponding to the closest alias is taken as relation alias information. Please refer Section 4.3.3.

the sentence, attention weight  $\alpha_i$  is calculated as:

$$\alpha_i = \frac{\exp(u_i)}{\sum_{j=1}^m \exp(u_j)} \text{ where, } u_i = h_i^{concat} \cdot r.$$

where  $r$  is a random query vector and  $u_i$  is the relevance score assigned to each token. Attention values  $\{\alpha_i\}$  are calculated by taking softmax over  $\{u_i\}$ . The representation of a sentence is given as a weighted sum of its tokens,  $s = \sum_{j=1}^m \alpha_j h_j^{concat}$ .

### 4.3.3 Side Information Acquisition

Relevant side information has been found to improve performance on several tasks [114, 199]. In distant supervision based relation extraction, since the entities are from a KB, knowledge about them can be utilized to improve relation extraction. Moreover, several unsupervised relation extraction methods (Open IE) [4, 128] allow extracting relation phrases between target entities without any predefined ontology and thus can be used to obtain relevant side information. In RESIDE, we employ Open IE methods and additional supervision from KB for improving neural relation extraction.

#### Relation Alias Side Information

RESIDE uses Stanford Open IE [4] for extracting relation phrases between target entities, which we denote by  $\mathcal{P}$ . As shown in Figure 4.2, for the sentence *Matt Coffin, executive of lowermybills, a company..*, Open IE methods extract “executive of” between *Matt Coffin* and

*lowermybills*. Further, we extend  $\mathcal{P}$  by including tokens at one hop distance in dependency path from target entities. Such features from dependency parse have been exploited in the past by [133, 74]. The degree of match between the extracted phrases in  $\mathcal{P}$  and aliases of a relation can give important clues about the relevance of that relation for the sentence. Several KBs like Wikidata provide such relation aliases, which can be readily exploited. In RESIDE, we further expand the relation alias set using Paraphrase database (PPDB) [155]. We note that even for cases when aliases for relations are not available, providing only the names of relations give competitive performance. We shall explore this point further in Section 4.4.2.3.

For matching  $\mathcal{P}$  with the PPDB expanded relation alias set  $\mathcal{R}$ , we project both in a  $d$ -dimensional space using GloVe embeddings [157]. Projecting phrases using word embeddings helps to further expand these sets, as semantically similar words are closer in embedding space [130, 157]. Then, for each phrase  $p \in \mathcal{P}$ , we calculate its cosine distance from all relation aliases in  $\mathcal{R}$  and take the relation corresponding to the closest relation alias as a matched relation for the sentence. We use a threshold on cosine distance to remove noisy aliases. In RESIDE, we define a  $k_r$ -dimensional embedding for each relation which we call as *matched relation embedding* ( $h^{rel}$ ). For a given sentence,  $h^{rel}$  is concatenated with its representation  $s$ , obtained from syntactic sentence encoder (Section 4.3.2) as shown in Figure 4.1. For sentences with  $|\mathcal{P}| > 1$ , we might get multiple matched relations. In such cases, we take the average of their embeddings. We hypothesize that this helps in improving the performance and find it to be true as shown in Section 4.4.2.

## Entity Type Side Information

Type information of target entities has been shown to give promising results on relation extraction [114, 219]. Every relation puts some constraint on the type of entities which can be its subject and object. For example, the relation *person/place\_of\_birth* can only occur between a *person* and a *location*. Sentences in distance supervision are based on entities in KBs, where the type information is readily available.

In RESIDE, we use types defined by FIGER [114] for entities in Freebase. For each type, we define a  $k_t$ -dimensional embedding which we call as *entity type embedding* ( $h^{type}$ ). For cases when an entity has multiple types in different contexts, for instance, *Paris* may have types *government* and *location*, we take the average over the embeddings of each type. We concatenate the *entity type embedding* of target entities to the final bag representation before using it for relation classification. To avoid over-parameterization, instead of using all fine-grained 112 entity types, we use 38 coarse types which form the first hierarchy of FIGER types.

Datasets	Split	# Sentences	# Entity-pairs
Riedel (# Relations: 53)	Train	455,771	233,064
	Valid	114,317	58,635
	Test	172,448	96,678
GDS (# Relations: 5)	Train	11,297	6,498
	Valid	1,864	1,082
	Test	5,663	3,247

Table 4.1: Details of datasets used. Please see Section 4.4.1.1 for more details.

#### 4.3.4 Instance Set Aggregation

For utilizing all valid sentences, following [113, 80], we use attention over sentences to obtain a representation for the entire bag. Instead of directly using the sentence representation  $s_i$  from Section 4.3.2, we concatenate the embedding of each sentence with *matched relation embedding*  $h_i^{rel}$  as obtained from Section 4.3.3. The attention score  $\alpha_i$  for  $i^{th}$  sentence is formulated as:

$$\alpha_i = \frac{\exp(\hat{s}_i \cdot q)}{\sum_{j=1}^n \exp(\hat{s}_j \cdot q)} \text{ where, } \hat{s}_i = [s_i; h_i^{rel}].$$

here  $q$  denotes a random query vector. The bag representation  $\mathcal{B}$ , which is the weighted sum of its sentences, is then concatenated with the *entity type embeddings* of the subject ( $h_{sub}^{type}$ ) and object ( $h_{obj}^{type}$ ) from Section 4.3.3 to obtain  $\hat{\mathcal{B}}$ .

$$\hat{\mathcal{B}} = [\mathcal{B}; h_{sub}^{type}; h_{obj}^{type}] \text{ where, } \mathcal{B} = \sum_{i=1}^n \alpha_i \hat{s}_i.$$

Finally,  $\hat{\mathcal{B}}$  is fed to a softmax classifier to get the probability distribution over the relations.

$$p(y) = \text{Softmax}(W \cdot \hat{\mathcal{B}} + b).$$

## 4.4 Experiments

### 4.4.1 Experimental Setup

#### 4.4.1.1 Datasets

In our experiments, we evaluate the models on Riedel and Google Distant Supervision (GDS) dataset. Statistics of the datasets is summarized in Table 4.1. Below we described each in

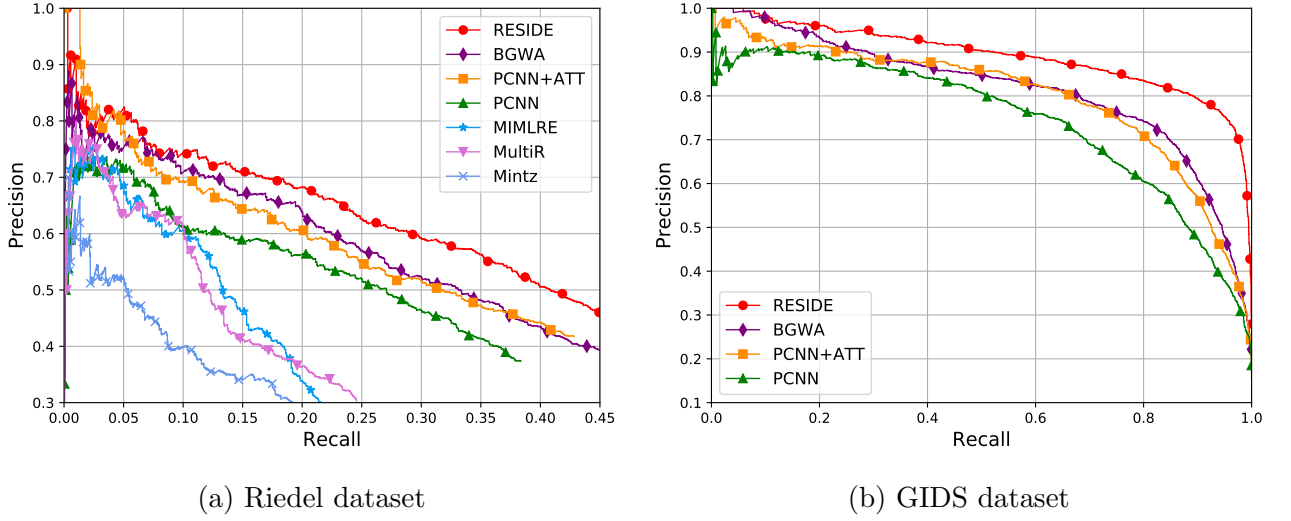


Figure 4.3: Comparison of Precision-recall curve. RESIDE achieves higher precision over the entire range of recall than all the baselines on both datasets. Please refer Section 4.4.2.1 for more details.

detail.

1. **Riedel:** The dataset is developed by [167] by aligning Freebase relations with New York Times (NYT) corpus, where sentences from the year 2005-2006 are used for creating the training set and from the year 2007 for the test set. The entity mentions are annotated using Stanford NER [58] and are linked to Freebase. The dataset has been widely used for RE by [78, 187] and more recently by [113, 57, 74].
2. **GIDS:** Jat et al. [80] created Google IISc Distant Supervision (GIDS) dataset by extending the Google relation extraction corpus<sup>1</sup> with additional instances for each entity pair. The dataset assures that the at-least-one assumption of multi-instance learning, holds. This makes automatic evaluation more reliable and thus removes the need for manual verification.

#### 4.4.1.2 Baselines

For evaluating RESIDE, we compare against the following baselines:

- **Mintz:** Multi-class logistic regression model proposed by [133] for distant supervision paradigm.
- **MultiR:** Probabilistic graphical model for multi instance learning by [78]

<sup>1</sup><https://research.googleblog.com/2013/04/50000-lessions-on-how-to-read-relation.html>

	One			Two			All		
	P@100	P@200	P@300	P@100	P@200	P@300	P@100	P@200	P@300
PCNN	73.3	64.8	56.8	70.3	67.2	63.1	72.3	69.7	64.1
PCNN+ATT	73.3	69.2	60.8	77.2	71.6	66.1	76.2	73.1	67.4
BGWA	78.0	71.0	63.3	81.0	73.0	64.0	82.0	75.0	72.0
RESIDE	<b>80.0</b>	<b>75.5</b>	<b>69.3</b>	<b>83.0</b>	<b>73.5</b>	<b>70.6</b>	<b>84.0</b>	<b>78.5</b>	<b>75.6</b>

Table 4.2: P@N for relation extraction using variable number of sentences in bags (with more than one sentence) in Riedel dataset. Here, One, Two and All represents the number of sentences randomly selected from a bag. RESIDE attains improved precision in all settings. More details in Section 4.4.2.1

- **MIMLRE:** A graphical model which jointly models multiple instances and multiple labels. More details in [187].
- **PCNN:** A CNN based relation extraction model by [230] which uses piecewise max-pooling for sentence representation.
- **PCNN+ATT:** A piecewise max-pooling over CNN based model which is used by [113] to get sentence representation followed by attention over sentences.
- **BGWA:** Bi-GRU based relation extraction model with word and sentence level attention [80].
- **RESIDE:** The method proposed in this chapter, please refer Section 4.3 for more details.

#### 4.4.1.3 Evaluation Criteria

Following the prior works [113, 57], we evaluate the models using held-out evaluation scheme. This is done by comparing the relations discovered from test articles with those in Freebase. We evaluate the performance of models with Precision-Recall curve and top-N precision (P@N) metric in our experiments.

## 4.4.2 Results

In this section we attempt to answer the following questions:

- Q1. Is RESIDE more effective than existing approaches for distant supervised RE? (4.4.2.1)
- Q2. What is the effect of ablating different components on RESIDE’s performance? (4.4.2.2)
- Q3. How is the performance affected in the absence of relation alias information? (4.4.2.3)

#### 4.4.2.1 Performance Comparison

For evaluating the effectiveness of our proposed method, RESIDE, we compare it against the baselines stated in Section 4.4.1.2. We use only the neural baselines on GDS dataset. The Precision-Recall curves on Riedel and GDS are presented in Figure 4.3. Overall, we find that RESIDE achieves higher precision over the entire recall range on both the datasets. All the non-neural baselines could not perform well as the features used by them are mostly derived from NLP tools which can be erroneous. RESIDE outperforms PCNN+ATT and BGWA which indicates that incorporating side information helps in improving the performance of the model. The higher performance of BGWA and PCNN+ATT over PCNN shows that attention helps in distant supervised RE. Following [113, 116], we also evaluate our method with different number of sentences. Results summarized in Table 4.2, show the improved precision of RESIDE in all test settings, as compared to the neural baselines, which demonstrates the efficacy of our model.

#### 4.4.2.2 Ablation Results

In this section, we analyze the effect of various components of RESIDE on its performance. For this, we evaluate various versions of our model with cumulatively removed components. The experimental results are presented in Figure 4.4a. We observe that on removing different components from RESIDE, the performance of the model degrades drastically. The results validate that GCNs are effective at encoding syntactic information. Further, the improvement from side information shows that it is complementary to the features extracted from text, thus validating the central thesis of this chapter, that inducing side information leads to improved relation extraction.

#### 4.4.2.3 Effect of Relation Alias Side Information

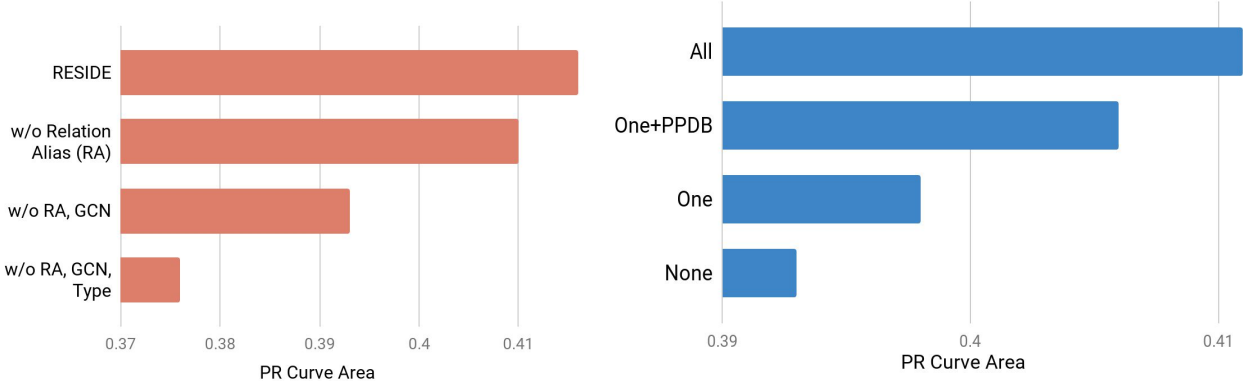
In this section, we test the performance of the model in setting where relation alias information is not readily available. For this, we evaluate the performance of the model on four different settings:

- **None:** Relation aliases are not available.
- **One:** The name of relation is used as its alias.
- **One+PPDB:** Relation name extended using Paraphrase Database (PPDB).
- **All:** Relation aliases from Knowledge Base<sup>1</sup>

The overall results are summarized in Figure 4.4b. We find that the model performs best

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<sup>1</sup>Each relation in Riedel dataset is manually mapped to corresponding Wikidata property for getting relation aliases. Few examples are presented in supplementary material.



(a) Performance comparison of different ablated version of RESIDE on Riedel dataset. Overall, [4.4.2.3](#) with respect to the presence of relation alias information helps RESIDE improve performance. Refer Section [4.4.2.2](#).

(b) Performance on settings defined in Section [4.4.2.3](#) with respect to the presence of relation alias side information on Riedel dataset. RESIDE performs comparably in the absence of relations from KB.

when aliases are provided by the KB itself. Overall, we find that RESIDE gives competitive performance even when very limited amount of relation alias information is available. We observe that performance improves further with the availability of more alias information.

## 4.5 Conclusion

In this chapter, we propose RESIDE, a novel neural network based model which makes principled use of relevant side information, such as entity type and relation alias, from Knowledge Base, for improving distant supervised relation extraction. RESIDE employs Graph Convolution Networks for encoding syntactic information of sentences and is robust to limited side information. Through extensive experiments on benchmark datasets, we demonstrate RESIDE’s effectiveness over state-of-the-art baselines. We have made RESIDE’s source code publicly available to promote reproducible research.

# Chapter 5

## Improving Convolution-based Knowledge Graph Embeddings by Increasing Feature Interactions

### 5.1 Introduction

In this chapter, we address the sparsity problems in Knowledge Graphs by inferring missing facts based on the known facts. Most existing KGs are incomplete [52]. The task of *link prediction* alleviates this problem by inferring missing facts based on the known facts in a KG. A popular approach for solving this problem involves learning a low-dimensional representation for all entities and relations and utilizing them to predict new facts. In general, most existing link prediction methods learn to embed KGs by optimizing a score function which assigns higher scores to true facts than invalid ones. These score functions can be classified as *translation distance based* [22, 214, 211] or *semantic matching based* [147, 115].

Recently, neural networks have also been utilized to learn the score function [179, 166, 50]. The motivation behind these approaches is that shallow methods like TransE [22] and DistMult [221] are limited in their expressiveness. As noted in Dettmers et al. [50], the only way to remedy this is to increase the size of their embeddings, which leads to an enormous increase in the number of parameters and hence limits their scalability to larger knowledge graphs.

Convolutional Neural Networks (CNN) have the advantage of using multiple layers, thus increasing their expressive power, while at the same time remaining parameter-efficient. Dettmers et al. [50] exploit these properties and propose ConvE - a model which applies convolutional filters on stacked 2D reshapings of entity and relation embeddings. Through this, they aim to increase the number of interactions between components of these embeddings.

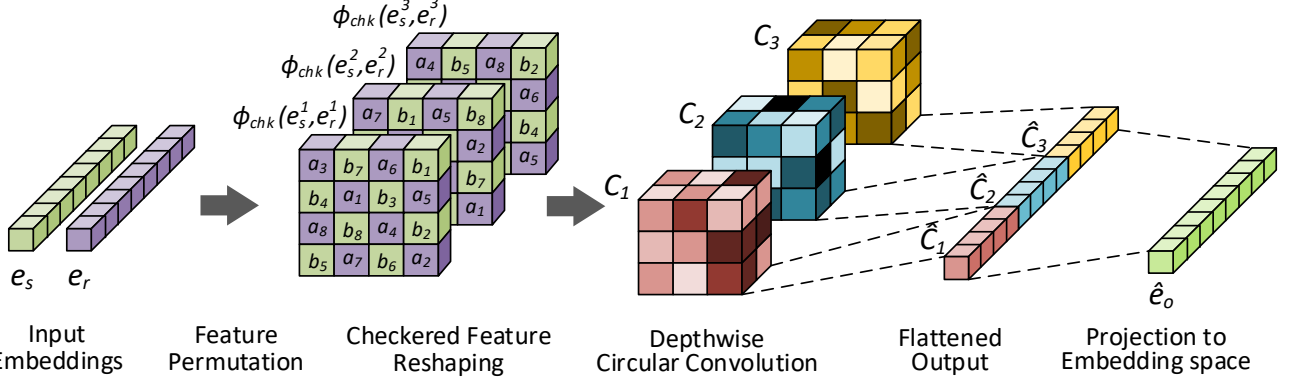


Figure 5.1: Overview of InteractE. Given entity and relation embeddings ( $e_s$  and  $e_r$  respectively), InteractE generates multiple permutations of these embeddings and reshapes them using a "Checkered" reshaping function ( $\phi_{chk}$ ). Depth-wise circular convolution is employed to convolve each of the reshaped permutations ( $C_i$ ), which are then flattened ( $\hat{C}_i$ ) and fed to a fully-connected layer to generate the predicted object embedding ( $\hat{e}_o$ ). Please refer to Section 5.4 for details.

In this chapter, we conclusively establish that increasing the number of such interactions is beneficial to link prediction performance, and show that the number of interactions that ConvE can capture is limited. We propose InteractE, a novel CNN based KG embedding approach which aims to further increase the interaction between relation and entity embeddings. Our contributions are summarized as follows:

1. We propose InteractE, a method that augments the expressive power of ConvE through three key ideas – feature permutation, "checkered" feature reshaping, and circular convolution.
2. We provide a precise definition of an interaction, and theoretically analyze InteractE to show that it increases interactions compared to ConvE. Further, we establish a correlation between the number of *heterogeneous interactions* (refer to Def. 5.2) and link prediction performance.
3. Through extensive evaluation on various link prediction datasets, we demonstrate InteractE's effectiveness (Section 5.7.2).

## 5.2 Related Work

**Non-Neural:** Starting with TransE [22], there have been multiple proposed approaches that use simple operations like dot products and matrix multiplications to compute a score function. Most approaches embed entities as vectors, whereas for relations, vector [22, 147], matrix [221,

[115](#) and tensor [112](#) representations have been explored. For modeling uncertainty of learned representations, Gaussian distributions [73, 214](#) have also been utilized. Methods like TransE [\[22\]](#) and TransH [\[211\]](#) utilize a translational objective for their score function, while DistMult [\[221\]](#) and ComplEx [\[195\]](#) use a bilinear diagonal based model.

**Neural Network based:** Recently, Neural Network (NN) based score functions have also been proposed. Neural Tensor Network [\[179\]](#) combines entity and relation embeddings by a relation-specific tensor which is given as input to a non-linear hidden layer for computing the score. Dong et al. [\[52\]](#), Ravishankar et al. [\[166\]](#) also utilize a Multi-Layer Perceptron for modeling the score function.

**Convolution based:** Convolutional Neural Networks (CNN) have also been employed for embedding Knowledge Graphs. ConvE [\[50\]](#) uses convolutional filters over reshaped subject and relation embeddings to compute an output vector and compares this with all other entities in the knowledge graph. [Shang et al.](#) propose ConvTransE a variant of the ConvE score function. They eschew 2D reshaping in favor of directly applying convolution on the stacked subject and relation embeddings. Further, they propose SACN which utilizes *weighted graph convolution* along with ConvTransE.

ConvKB [\[143\]](#) is another convolution based method which applies convolutional filters of width 1 on the stacked subject, relation and object embeddings for computing score. As noted in [\[173\]](#), although ConvKB was claimed to be superior to ConvE, its performance is not consistent across different datasets and metrics. Further, there have been concerns raised about the validity of its evaluation procedure<sup>1</sup> Hence, we do not compare against it in this chapter. A survey of all variants of existing KG embedding techniques can be found in [\[145, 208\]](#).

### 5.3 Notation and Definitions

Let  $\mathbf{e}_s = (a_1, \dots, a_d), \mathbf{e}_r = (b_1, \dots, b_d)$ , where  $a_i, b_i \in \mathbb{R} \forall i$ , be an entity and a relation embedding respectively, and let  $\mathbf{w} \in \mathbb{R}^{k \times k}$  be a convolutional kernel of size  $k$ . Further, we define that a matrix  $M_k \in \mathbb{R}^{k \times k}$  is a  $k$ -submatrix of another matrix  $N \in \mathbb{R}^{m \times n}$  if  $\exists i, j$  such that  $M_k = N_{i:i+k, j:j+k}$ . We denote this by  $M_k \subseteq N$ .

**Definition 5.1 (Reshaping Function)** A *reshaping function*  $\phi : \mathbb{R}^d \times \mathbb{R}^d \rightarrow \mathbb{R}^{m \times n}$  transforms embeddings  $\mathbf{e}_s$  and  $\mathbf{e}_r$  into a matrix  $\phi(\mathbf{e}_s, \mathbf{e}_r)$ , where  $m \times n = 2d$ . For conciseness, we abuse notation and represent  $\phi(\mathbf{e}_s, \mathbf{e}_r)$  by  $\phi$ . We define three types of reshaping functions.

- **Stack** ( $\phi_{stk}$ ) reshapes each of  $\mathbf{e}_s$  and  $\mathbf{e}_r$  into a matrix of shape  $(m/2) \times n$ , and stacks them along their height to yield an  $m \times n$  matrix (Fig. [5.2a](#)). This is the reshaping function used

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<sup>1</sup><https://openreview.net/forum?id=HkgEQnRqYQ&noteId=HklyVUAX2m>

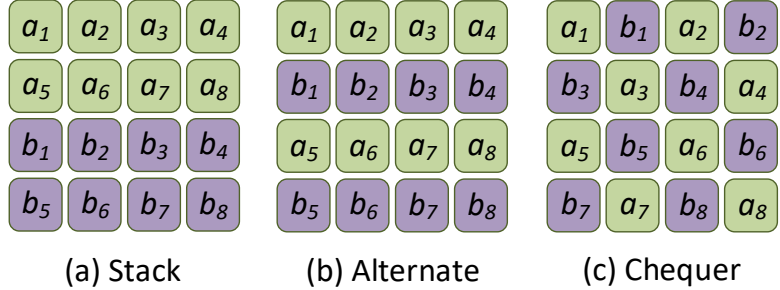


Figure 5.2: Different types of reshaping functions we analyze in this chapter. Here,  $e_s = (a_1, \dots, a_8)$ ,  $e_r = (b_1, \dots, b_8)$ , and  $m = n = 4$ . Please refer to Section 5.3 for more details.

in Dettmers et al. [50].

- **Alternate** ( $\phi_{alt}^\tau$ ) reshapes  $e_s$  and  $e_r$  into matrices of shape  $(m/2) \times n$ , and stacks  $\tau$  rows of  $e_s$  and  $e_r$  alternately. In other words, as we decrease  $\tau$ , the "frequency" with which rows of  $e_s$  and  $e_r$  alternate increases. We denote  $\phi_{alt}^1$  as  $\phi_{alt}$  for brevity (Fig. 5.2b).
- **Chequer** ( $\phi_{chk}$ ) arranges  $e_s$  and  $e_r$  such that no two adjacent cells are occupied by components of the same embedding (Fig. 5.2c).

**Definition 5.2 (Interaction)** An interaction is defined as a triple  $(x, y, M_k)$ , such that  $M_k \subseteq \phi(e_s, e_r)$  is a  $k$ -submatrix of the reshaped input embeddings;  $x, y \in M_k$  and are distinct components of  $e_s$  or  $e_r$ . The number of interactions  $\mathcal{N}(\phi, k)$  is defined as the cardinality of the set of all possible triples. Note that  $\phi$  can be replaced with  $\Omega(\phi)$  for some padding function  $\Omega$ .

An interaction  $(x, y, M_k)$  is called **heterogeneous** if  $x$  and  $y$  are components of  $e_s$  and  $e_r$  respectively, or vice-versa. Otherwise, it is called **homogeneous**. Let  $\mathcal{N}_{het}(\phi, k)$  and  $\mathcal{N}_{homo}(\phi, k)$  denote the number of heterogeneous and homogeneous interactions respectively. For example, in a  $3 \times 3$  matrix  $M_3$ , if there are 5 components of  $e_s$  and 4 of  $e_r$ , then the number of interactions within this matrix is  $\mathcal{N}_{het} = 2(5 \times 4) = 40$ , and  $\mathcal{N}_{homo} = 2 \left[ \binom{5}{2} + \binom{4}{2} \right] = 32$ .

## 5.4 InteractE Overview

Recent methods [221, 147] have demonstrated that expressiveness of a model can be enhanced by increasing the possible interactions between embeddings. ConvE [50] also exploits the same principle albeit in a limited way, using convolution on 2D reshaped embeddings. InteractE extends this notion of capturing entity and relation feature interactions using the following three ideas:

- **Feature Permutation:** Instead of using one fixed order of the input, we utilize multiple permutations to capture more possible interactions.

- **Checkered Reshaping:** We substitute simple feature reshaping of ConvE with checked reshaping and prove its superiority over other possibilities.
- **Circular Convolution:** Compared to the standard convolution, circular convolution allows to capture more feature interactions as depicted in Figure 5.3. The convolution is performed in a depth-wise manner [39] on different input permutations.

## 5.5 InteractE Details

In this section, we provide a detailed description of the various components of InteractE. The overall architecture is depicted in Fig. 5.1. InteractE learns a  $d$ -dimensional vector representation ( $\mathbf{e}_s, \mathbf{e}_r \in \mathbb{R}^d$ ) for each entity and relation in the knowledge graph, where  $d = d_w d_h$ .

### 5.5.1 Feature Permutation

To capture a variety of heterogeneous interactions, InteractE first generates  $t$ -random permutations of both  $e_s$  and  $e_r$ , denoted by  $\mathcal{P}_t = [(\mathbf{e}_s^1, \mathbf{e}_r^1); \dots; (\mathbf{e}_s^t, \mathbf{e}_r^t)]$ . Note that with high probability, the sets of interactions within  $\phi(\mathbf{e}_s^i, \mathbf{e}_r^i)$  for different  $i$  are disjoint. This is evident because the number of distinct interactions across all possible permutations is very large. So, for  $t$  different permutations, we can expect the total number of interactions to be approximately  $t$  times the number of interactions for one permutation.

### 5.5.2 Checkered Reshaping

Next, we apply the reshaping operation  $\phi_{chk}(\mathbf{e}_s^i, \mathbf{e}_r^i), \forall i \in \{1, \dots, t\}$ , and define  $\phi(\mathcal{P}_t) = [\phi(\mathbf{e}_s^1, \mathbf{e}_r^1); \dots; \phi(\mathbf{e}_s^t, \mathbf{e}_r^t)]$ . ConvE [50] uses  $\phi_{stk}(\cdot)$  as a reshaping function which has limited interaction capturing ability. On the basis of Proposition 5.3, we choose to utilize  $\phi_{chk}(\cdot)$  as the reshaping function in InteractE, which captures maximum *heterogeneous* interactions between entity and relation features.

### 5.5.3 Circular Convolution

Motivated by our analysis in Proposition 5.4, InteractE uses *circular convolution*, which further increases interactions compared to the standard convolution. This has been successfully applied for tasks like image recognition [209]. Circular convolution on a 2-dimensional input  $\mathbf{I} \in \mathbb{R}^{m \times n}$  with a filter  $w \in \mathbb{R}^{k \times k}$  is defined as:

$$[\mathbf{I} * \mathbf{w}]_{p,q} = \sum_{i=-\lfloor k/2 \rfloor}^{\lfloor k/2 \rfloor} \sum_{j=-\lfloor k/2 \rfloor}^{\lfloor k/2 \rfloor} \mathbf{I}_{[p-i]_m, [q-j]_n} \mathbf{w}_{i,j},$$

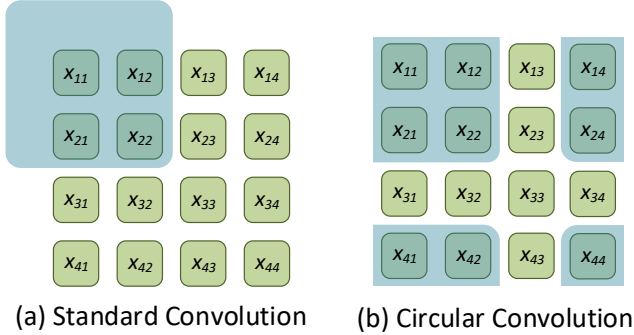


Figure 5.3: Circular convolution induces more interactions than standard convolution. Here,  $X$  is a  $4 \times 4$  input matrix with components  $x_{ij}$ . The shaded region depicts where the filter is applied. Please refer to Section 5.5.3 for more details.

where,  $[x]_n$  denotes  $x$  modulo  $n$  and  $\lfloor \cdot \rfloor$  denotes the floor function. Figure 5.3 and Proposition 5.4 show how circular convolution captures more interactions compared to standard convolution with zero padding.

InteractE stacks each reshaped permutation as a separate channel. For convolving permutations, we apply circular convolution in a *depth-wise* manner [39]. Although different sets of filters can be applied for each permutation, in practice we find that sharing filters across channels works better as it allows a single set of kernel weights to be trained on more input instances.

#### 5.5.4 Score Function

The output of each circular convolution is flattened and concatenated into a vector. InteractE then projects this vector to the embedding space ( $\mathbb{R}^d$ ). Formally, the score function used in InteractE is defined as follows:

$$\psi(s, r, o) = g(\text{vec}(f(\phi(\mathcal{P}_k) \circledast \mathbf{w})) \mathbf{W}) \mathbf{e}_o,$$

where  $\circledast$  denotes depth-wise circular convolution,  $\text{vec}(\cdot)$  denotes vector concatenation,  $\mathbf{e}_o$  represents the object entity embedding matrix and  $W$  is a learnable weight matrix. Functions  $f$  and  $g$  are chosen to be ReLU and sigmoid respectively. For training, we use the standard binary cross entropy loss with label smoothing.

## 5.6 Theoretical Analysis

In this section, we analyze multiple variants of 2D reshaping with respect to the number of interactions they induce. We also examine the advantages of using circular padded convolution

over the standard convolution.

For simplicity, we restrict our analysis to the case where the output of the reshaping function is a square matrix, i.e.,  $m = n$ . Note that our results can be extended to the general case as well. Proofs of all propositions herein are included in the supplementary material.

**Proposition 5.1** *For any kernel  $\mathbf{w}$  of size  $k$ , for all  $n \geq (\frac{5k}{3} - 1)$  if  $k$  is odd and  $n \geq \frac{(5k+2)(k-1)}{3k}$  if  $k$  is even, the following statement holds:*

$$\mathcal{N}_{het}(\phi_{alt}, k) \geq \mathcal{N}_{het}(\phi_{stk}, k)$$

**Proof:** Any  $M_k \in \mathbb{R}^{k \times k}$ ,  $M_k \subseteq \phi_{alt}$  contains  $\lfloor \frac{k}{2} \rfloor$  rows of elements of  $\mathbf{e}_s$ , and  $\lfloor \frac{k+1}{2} \rfloor$  rows of elements of  $\mathbf{e}_r$ , or vice-versa.

For a single fixed  $M_k$ , the total number of triples  $(a_i, b_j, M_k)$  and  $(b_j, a_i, M_k)$  is

$$2 \times k \left\lfloor \frac{k}{2} \right\rfloor \times k \left\lfloor \frac{k+1}{2} \right\rfloor$$

The number of possible  $M_k$  matrices is  $(n-k+1)^2$ . Hence the total number of heterogeneous interactions is

$$\mathcal{N}_{het}(\phi_{alt}, k) = (n-k+1)^2 k^2 \times 2 \left\lfloor \frac{k}{2} \right\rfloor \left\lfloor \frac{k+1}{2} \right\rfloor$$

Any  $M_k \in \mathbb{R}^{k \times k}$ ,  $M_k \subseteq \phi_{stk}$  contains  $l$  rows of elements of  $\mathbf{e}_s$ , and  $k-l$  rows of elements of  $\mathbf{e}_r$ , where  $0 \leq l \leq k$ .

For a fixed  $l$ , the number of different possible  $M_k$  matrices is  $(n-k+1)$ . Hence, the number of heterogeneous interactions is

$$\begin{aligned} \mathcal{N}_{het}(\phi_{stk}, k) &= (n-k+1) \left( \sum_{l=0}^k 2 \times kl \times k(k-l) \right) \\ &= (n-k+1) \cdot k^2 \cdot \left( k^2(k+1) - \frac{k(k+1)(2k+1)}{3} \right) \\ &= (n-k+1) \cdot k^2 \cdot \left( \frac{k(k+1)(k-1)}{3} \right) \end{aligned} \tag{5.1}$$

We need to check whether,

$$\begin{aligned} \mathcal{N}_{het}(\phi_{alt}, k) &\geq \mathcal{N}_{het}(\phi_{stk}, k) \\ \implies (n-k+1) \left\lfloor \frac{k}{2} \right\rfloor \left\lfloor \frac{k+1}{2} \right\rfloor &\geq \frac{k(k+1)(k-1)}{6} \end{aligned}$$

For odd  $k$ , this becomes

$$\begin{aligned} (n - k + 1) \left( \frac{k - 1}{2} \right) \left( \frac{k + 1}{2} \right) &\geq \frac{k(k + 1)(k - 1)}{6} \\ n - k + 1 &\geq \frac{2k}{3} \\ n &\geq \frac{5k}{3} - 1 \end{aligned}$$

For even  $k$ ,

$$\begin{aligned} (n - k + 1) \left( \frac{k}{2} \right)^2 &\geq \frac{k(k + 1)(k - 1)}{6} \\ nk - k(k - 1) &\geq \frac{2(k + 1)(k - 1)}{3} \\ n &\geq \frac{(5k + 2)(k - 1)}{3k} \end{aligned}$$

□

**Proposition 5.2** *For any kernel  $\mathbf{w}$  of size  $k$  and for all  $\tau < \tau'$  ( $\tau, \tau' \in \mathbb{N}$ ), the following statement holds:*

$$\mathcal{N}_{het}(\phi_{alt}^\tau, k) \geq \mathcal{N}_{het}(\phi_{alt}^{\tau'}, k)$$

**Proof:** For simplicity, let us assume that  $\alpha = n/(2\tau) \in \mathbb{N}$ , i.e.,  $\phi_{alt}^\tau$  is composed of exactly  $\alpha$  blocks of  $\tau$  rows of  $\mathbf{e}_s$  and  $\mathbf{e}_r$  stacked alternately. Also, when  $\tau < k$ , we assume that  $k/\tau \in \mathbb{N}$ . Now, for any  $M_k \in \mathbb{R}^{k \times k}$ ,  $M_k \subseteq \phi_{alt}^\tau$ , we consider the following two cases:

**Case 1  $\tau \geq k - 1$ :** It is easy to see that this case can be split into  $n/\tau$  subproblems, each of which is similar to  $\phi_{stk}$ . Hence,

$$\mathcal{N}_{het}(\phi_{alt}^\tau, k) = \left( \frac{n}{\tau} \right) \mathcal{N}_{het}(\phi_{stk}, k)$$

Clearly,  $\mathcal{N}_{het}(\phi_{alt}^\tau, k)$  is monotonically decreasing with increasing  $\tau$ .

**Case 2  $\tau < k - 1$ :** As shown in Fig. 5.4, let  $T_a, T_b \in \mathbb{R}^{\tau \times k}$  denote a submatrix formed by components of  $\mathbf{e}_s, \mathbf{e}_r$  respectively. Note that if  $k$  is even, then for any  $M_k \subseteq \phi_{alt}^\tau$ , the number of components of  $\mathbf{e}_s$  and  $\mathbf{e}_r$  are always equal to  $k^2/2$  each. For odd  $k$ , the number of  $T_a$ 's and  $T_b$ 's are  $\frac{(k/\tau)+1}{2}$  and  $\frac{(k/\tau)-1}{2}$  in some order. Now, if we move  $M_k$  down by  $i$  rows ( $i \leq \tau$ ), the

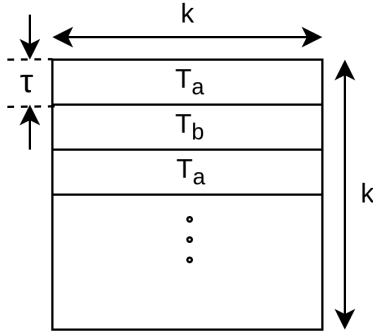


Figure 5.4: The figure depicts a  $k \times k$  matrix  $M_k$ .  $T_a, T_b$  are reshaped matrices each containing  $\tau k$  components of  $\mathbf{e}_s, \mathbf{e}_r$  respectively.

total number of heterogeneous interactions across all such positions is:

$$\begin{aligned}
& \frac{n}{\tau} \sum_{i=0}^{\tau-1} k^2 \left( \left( \frac{k/\tau + 1}{2} \right) \tau - i \right) \left( \left( \frac{k/\tau - 1}{2} \right) \tau + i \right) \\
&= \frac{nk^2}{\tau} \sum_{i=0}^{\tau-1} \frac{(k + \tau - 2i)(k - \tau + 2i)}{4} \\
&= \frac{nk^2}{4\tau} \sum_{i=0}^{\tau-1} k^2 - (\tau^2 + 4i^2 - 4i\tau) \\
&= \frac{nk^2}{4} \left( (k^2 - \tau^2) - \frac{4(\tau-1)(2\tau-1)}{6} + \frac{4\tau(\tau-1)}{2} \right) \\
&= C \left( k^2 - \frac{\tau^2}{3} - \frac{2}{3} \right)
\end{aligned}$$

We can see that this is also monotonically decreasing with increasing  $\tau$ . It is also evident that the above expression is maximum at  $\tau = 1$  (since  $\tau \in \mathbb{N}$ ).

□

**Proposition 5.3** For any kernel  $\mathbf{w}$  of size  $k$  and for all reshaping functions  $\phi : \mathbb{R}^d \times \mathbb{R}^d \rightarrow \mathbb{R}^{n \times n}$ , the following statement holds:

$$\mathcal{N}_{het}(\phi_{chk}, k) \geq \mathcal{N}_{het}(\phi, k)$$

**Proof:** For any  $\phi$ , and for any  $M_k \in \mathbb{R}^{k \times k}$  such that  $M_k \subseteq \phi$ , let  $x, y$  be the number of components of  $\mathbf{e}_s$  and  $\mathbf{e}_r$  in  $M_k$  respectively. Then  $\mathcal{N}_{het}(M_k, k) = 2xy$ . Also, since total number of elements in  $M_k$  is fixed, we have  $x + y = k^2$ .

Using the AM-GM inequality on  $x, y$  we have,

$$xy \leq \left( \frac{x+y}{2} \right)^2 = \frac{k^4}{4}$$

If  $k$  is odd, since  $x, y \in \mathbb{N}$ ,

$$xy \leq \frac{k^4 - 1}{4}$$

Therefore, the maximum interaction occurs when  $x = y = \frac{k^2}{2}$  (for even  $k$ ), or  $x = \frac{k^2+1}{2}, y = \frac{k^2-1}{2}$  (for odd  $k$ ). It can be easily verified that this property holds for all  $M_k \subseteq \phi_{chk}$ . Hence,

$$\mathcal{N}_{het}(\phi, k) = \sum_{M_k} 2xy \leq \sum_{M_k} \frac{2k^4}{4} = \mathcal{N}_{het}(\phi_{chk}, k)$$

□

**Lemma 5.1** Consider a matrix  $M_k \in \mathbb{R}^{k \times k}$ . Create another matrix  $M'_k$  by replacing some  $p$  components ( $p \geq 0$ ) of  $\mathbf{e}_s$  or  $\mathbf{e}_r$  in  $M_k$  with 0. Then,

$$\mathcal{N}_{het}(M_k, k) \geq \mathcal{N}_{het}(M'_k, k),$$

**Proof:** If  $M_k$  contains  $x$  components of  $\mathbf{e}_s$  and  $y$  components of  $\mathbf{e}_r$ , then  $\mathcal{N}_{het}(M_k, k) = 2xy$ , and  $\mathcal{N}_{het}(M'_k, k) = 2(x-l)(y-(p-l))$  for some  $l \leq p$  and  $l \leq x$ . We observe that

$$\begin{aligned} \mathcal{N}_{het}(M'_k, k) &= \mathcal{N}_{het}(M_k, k) - 2(x-l)(p-l) - 2ly \\ &\leq \mathcal{N}_{het}(M_k, k) \end{aligned}$$

□

**Proposition 5.4** Let  $\Omega_0, \Omega_c : \mathbb{R}^{n \times n} \rightarrow \mathbb{R}^{(n+p) \times (n+p)}$  denote zero padding and circular padding functions respectively, for some  $p > 0$ . Then for any reshaping function  $\phi$ ,

$$\mathcal{N}_{het}(\Omega_c(\phi), k) \geq \mathcal{N}_{het}(\Omega_0(\phi), k)$$

**Proof:** Given  $\Omega_c(\phi)$ , we know that we can obtain  $\Omega_0(\phi)$  by replacing certain components of  $\Omega_c(\phi)$  with 0. So for every  $M_k \subseteq \Omega_c(\phi)$ , there is a corresponding  $M'_k \subseteq \Omega_0(\phi)$  which is obtained by replacing some  $p$  components ( $p \geq 0$ ) of  $M_k$  with 0.

Dataset	$ \mathcal{E} $	$ \mathcal{R} $	# Triples		
			Train	Valid	Test
FB15k-237	14,541	237	272,115	17,535	20,466
WN18RR	40,943	11	86,835	3,034	3,134
YAGO3-10	123,182	37	1,079,040	5,000	5,000

Table 5.1: Details of the datasets used. Please see Section 5.7.1.1 for more details.

Using the above Lemma, we can see that

$$\begin{aligned}
\mathcal{N}_{het}(\Omega_c(\phi), k) &= \sum_{M_k \subseteq \Omega_c(\phi)} \mathcal{N}_{het}(M_k, k)) \\
&\geq \sum_{M'_k \subseteq \Omega_0(\phi)} \mathcal{N}_{het}(M'_k, k)) \\
&= \mathcal{N}_{het}(\Omega_0(\phi), k))
\end{aligned}$$

□

## 5.7 Experiments

### 5.7.1 Experimental Setup

#### 5.7.1.1 Datasets

In our experiments, following Dettmers et al. [50], Sun et al. [186], we evaluate on the three most commonly used link prediction datasets. A summary statistics of the datasets is presented in Table 5.1.

- **FB15k-237** [193] is a improved version of FB15k [22] dataset where all inverse relations are deleted to prevent direct inference of test triples by reversing training triples.
- **WN18RR** [50] is a subset of WN18 [22] derived from WordNet [132], with deleted inverse relations similar to FB15k-237.
- **YAGO3-10** is a subset of YAGO3 [185] constitutes entities with at least 10 relations. Triples consist of descriptive attributes of people.

#### 5.7.1.2 Evaluation protocol

Following [22], we use the filtered setting, i.e., while evaluating on test triples, we filter out all the valid triples from the candidate set, which is generated by either corrupting the head or tail entity of a triple. The performance is reported on the standard evaluation metrics: Mean

	FB15k-237				WN18RR				YAGO3-10			
	MRR	MR	H@10	H@1	MRR	MR	H@10	H@1	MRR	MR	H@10	H@1
DistMult [221]	.241	254	.419	.155	.430	5110	.49	.39	.34	5926	.54	.24
ComplEx [195]	.247	339	.428	.158	.44	5261	.51	.41	.36	6351	.55	.26
R-GCN [171]	.248	-	.417	.151	-	-	-	-	-	-	-	-
KBGAN [30]	.278	-	.458	-	.214	-	.472	-	-	-	-	-
KBLRN [64]	.309	209	.493	.219	-	-	-	-	-	-	-	-
ConvTransE [173]	.33	-	.51	.24	.46	-	.52	.43	-	-	-	-
SACN [173]	.35	-	.54	.26	.47	-	.54	.43	-	-	-	-
RotatE [186]	.338	177	.533	.241	<b>.476</b>	<b>3340</b>	<b>.571</b>	.428	.495	1767	.670	.402
ConvE [50]	.325	244	.501	.237	.43	4187	.52	.40	.44	<b>1671</b>	.62	.35
InteractE (Proposed Method)	<b>.354</b>	<b>172</b>	<b>.535</b>	<b>.263</b>	.463	5202	.528	<b>.430</b>	<b>.541</b>	2375	<b>.687</b>	<b>.462</b>

Table 5.2: Link prediction results of several models evaluated on FB15k-237, WN18RR and YAGO3-10. We find that InteractE outperforms all other methods across metrics on FB15k-237 and in 3 out of 4 settings on YAGO3-10. Since InteractE generalizes ConvE, we highlight performance comparison between the two methods specifically in the table above. Please refer to Section 5.7.2.1 for more details.

Reciprocal Rank (MRR), Mean Rank (MR) and Hits@1, and Hits@10. We report average results across 5 runs. We note that the variance is substantially low on all the metrics and hence omit it.

### 5.7.1.3 Baselines

In our experiments, we compare InteractE against a variety of baselines which can be categorized as:

- **Non-neural:** Methods that use simple vector based operations for computing score. For instance, DistMult [221], ComplEx [195], KBGAN [30], KBLRN [64] and RotatE [186].
- **Neural:** Methods which leverage a non-linear neural network based architecture in their scoring function. This includes R-GCN [171], ConvE [50], ConvTransE [173], and SACN [173].

## 5.7.2 Results

In this section, we attempt to answer the questions below:

- Q1. How does InteractE perform in comparison to the existing approaches? (Section 5.7.2.1)
- Q2. What is the effect of different feature reshaping and circular convolution on link prediction performance? (Section 5.7.2.2)
- Q3. How does the performance of our model vary with number of feature permutations? (Section 5.7.2.3)

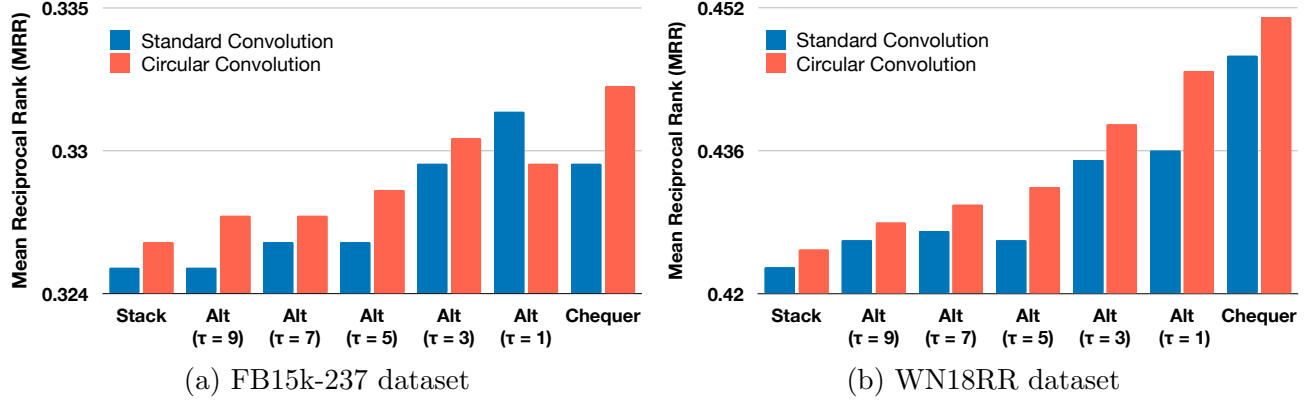


Figure 5.5: Performance with different feature reshaping and convolution operation on validation data of FB15k-237 and WN18RR. Stack and Alt denote *Stacked* and *Alternate* reshaping as defined in Section 5.3. As we decrease  $\tau$  the number of heterogeneous interactions increases (refer to Proposition 5.2). The results empirically verify our theoretical claim in Section 5.6 and validate the central thesis of this chapter that increasing *heterogeneous interactions* improves link prediction performance. Please refer to Section 5.7.2.2 for more details.

Q4. What is the performance of InteractE on different relation types? (Section 5.7.2.4)

### 5.7.2.1 Performance Comparison

In order to evaluate the effectiveness of InteractE, we compare it against the existing knowledge graph embedding methods listed in Section 5.7.1.3. The results on three standard link prediction datasets are summarized in Table 5.2. The scores of all the baselines are taken directly from the values reported in the papers [50, 186, 173, 30, 64]. Since our model builds on ConvE, we specifically compare against it, and find that InteractE outperforms ConvE on all metrics for FB15k-237 and WN18RR and on three out of four metrics on YAGO3-10. On an average, InteractE obtains an improvement of 9%, 7.5%, and 23% on FB15k-237, WN18RR, and YAGO3-10 on MRR over ConvE. This validates our hypothesis that increasing heterogeneous interactions help improve performance on link prediction. For YAGO3-10, we observe that the MR obtained from InteractE is worse than ConvE although it outperforms ConvE on all other metrics. Similar trend has been observed in [50, 186].

Compared to other baseline methods, InteractE outperforms them on FB15k-237 across all the metrics and on 3 out of 4 metrics on YAGO3-10 dataset. The below-par performance of InteractE on WN18RR can be attributed to the fact that this dataset is more suitable for shallow models as it has very low average relation-specific in-degree. This is consistent with the observations of Dettmers et al. [50].

		RotatE			ConvE			InteractE		
		MRR	MR	H@10	MRR	MR	H@10	MRR	MR	H@10
Head Pred	1-1	<b>0.498</b>	359	<b>0.593</b>	0.374	223	0.505	0.386	<b>175</b>	0.547
	1-N	0.092	614	0.174	0.091	700	0.17	<b>0.106</b>	<b>573</b>	<b>0.192</b>
	N-1	<b>0.471</b>	108	<b>0.674</b>	0.444	73	0.644	0.466	<b>69</b>	0.647
	N-N	0.261	<b>141</b>	<b>0.476</b>	0.261	158	0.459	<b>0.276</b>	148	<b>0.476</b>
Tail Pred	1-1	<b>0.484</b>	307	<b>0.578</b>	0.366	<b>261</b>	0.51	0.368	308	0.547
	1-N	0.749	41	0.674	0.762	33	0.878	<b>0.777</b>	<b>27</b>	<b>0.881</b>
	N-1	<b>0.074</b>	<b>578</b>	0.138	0.069	682	0.15	<b>0.074</b>	625	<b>0.141</b>
	N-N	0.364	<b>90</b>	0.608	0.375	100	0.603	<b>0.395</b>	92	<b>0.617</b>

Table 5.3: Link prediction results by relation category on FB15k-237 dataset for RotatE, ConvE, and InteractE. Following (Wang et al., 2014b), the relations are categorized into one-to-one (1-1), one-to-many (1-N), many-to-one (N-1), and many-to-many (N-N). We observe that InteractE is effective at capturing complex relations compared to RotatE. Refer to Section 5.7.2.4 for details.

### 5.7.2.2 Effect of Feature Reshaping and Circular Convolution

In this section, we empirically test the effectiveness of different reshaping techniques we analyzed in Section 5.6. For this, we evaluate different variants of InteractE on validation data of FB15k-237 and WN18RR with the number of feature permutations set to 1. We omit the analysis on YAGO3-10 given its large size. The results are summarized in Figure 5.5. We find that the performance with *Stacked* reshaping is the worst, and it improves when we replace it with *alternate* reshaping. This observation is consistent with our findings in Proposition 5.1. Further, we find that MRR improves on decreasing the value of  $\tau$  in alternate reshaping, which empirically validates Proposition 5.2. Finally, we observe that *checkered* reshaping gives the best performance across all reshaping functions for most scenarios, thus justifying Proposition 5.3.

We also compare the impact of using circular and standard convolution on link prediction performance. The MRR scores are reported in Figure 5.5. The results show that circular convolution is consistently better than the standard convolution. This also verifies our statement in Proposition 5.4. Overall, we find that increasing interaction helps improve performance on the link prediction task, thus validating the central thesis of our work.

### 5.7.2.3 Effect of Feature Permutations

In this section, we analyze the effect of increasing the number of feature permutations on InteractE’s performance on validation data of FB15k-237, WN18RR, and YAGO3-10. The overall

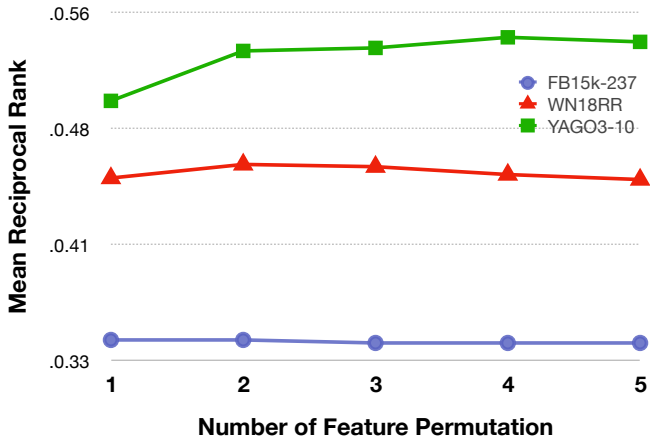


Figure 5.6: Performance on the validation data of FB15k-237, WN18RR, and YAGO3-10 with different numbers of feature permutations. We find that although increasing the number of permutations improves performance, it saturates as we exceed a certain limit. Please see Section 5.7.2.3 for details.

results are summarized in Figure 5.6. We observe that on increasing the number of permutations although on FB15k-237, MRR remains the same, it improves on WN18RR and YAGO3-10 datasets. However, it degrades as the number of permutations is increased beyond a certain limit. We hypothesize that this is due to over-parameterization of the model. Moreover, since the number of relevant interactions are finite, increasing the number of permutations could become redundant beyond a limit.

#### 5.7.2.4 Evaluation on different Relation Types

In this section, we analyze the performance of InteractE on different relation categories of FB15k-237. We chose FB15k-237 for analysis over other datasets because of its more and diverse set of relations. Following [210], we classify the relations based on the average number of tails per head and heads per tail into four categories: one-to-one, one-to-many, many-to-one, and many-to-many. The results are presented in Table 5.3. Overall, we find that InteractE is effective at modeling complex relation types like one-to-many and many-to-many whereas, RotatE captures simple relations like one-to-one better. This demonstrates that an increase in interaction allows the model to capture more complex relationships.

## 5.8 Conclusion

In this chapter, we propose InteractE, a novel knowledge graph embedding method which alleviates the limitations of ConvE by capturing additional heterogeneous feature interactions. InteractE is able to achieve this by utilizing three central ideas, namely feature permutation,

checkered feature reshaping, and circular convolution. Through extensive experiments, we demonstrate that InteractE achieves a consistent improvement on link prediction performance on multiple datasets. We also theoretically analyze the effectiveness of the components of InteractE, and provide empirical validation of our hypothesis that increasing heterogeneous feature interaction is beneficial for link prediction with ConvE. This work demonstrates a possible scope for improving existing knowledge graph embedding methods by leveraging rich heterogenous interactions.

## **Part II**

# **Exploiting Graph Convolutional Networks in NLP**

# Chapter 6

## Documents Timestamping using Graph Convolutional Networks

### 6.1 Introduction

Date of a document, also referred to as the Document Creation Time (DCT), is at the core of many important tasks, such as, information retrieval [150, 106, 43], temporal reasoning [123, 118], text summarization [207], event detection [1], and analysis of historical text [45], among others. In all such tasks, the document date is assumed to be available and also accurate – a strong assumption, especially for arbitrary documents from the Web. Thus, there is a need to automatically predict the date of a document based on its content. This problem is referred to as *Document Dating*.

Initial attempts on automatic document dating started with generative models by [45]. This model is later improved by [86] who incorporate additional features such as POS tags, collocations, etc. Chambers [32] shows significant improvement over these prior efforts through their discriminative models using handcrafted temporal features. Kotsakos et al. [94] propose a statistical approach for document dating exploiting term burstiness [98].

Document dating is a challenging problem which requires extensive reasoning over the temporal structure of the document. Let us motivate this through an example shown in Figure 6.1. In the document, *four years after* plays a crucial role in identifying the creation time of the document. The existing approaches give higher confidence for timestamp immediate to the year mention 1995. NeurdalDater exploits the syntactic and temporal structure of the document to predict the right timestamp (1999) for the document. With the exception of [32], all prior works on the document dating problem ignore such informative temporal structure within the document.

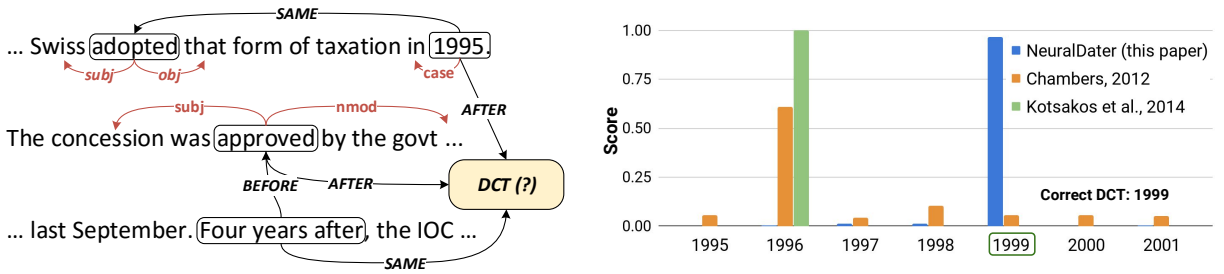


Figure 6.1: **Top:** An example document annotated with syntactic and temporal dependencies. In order to predict the right value of 1999 for the Document Creation Time (DCT), inference over these document structures is necessary. **Bottom:** Document date prediction by two state-of-the-art baselines and NeurdalDater, the method proposed in this chapter. While the two previous methods are getting misled by the temporal expression (1995) in the document, NeurdalDater is able to use the syntactic and temporal structure of the document to predict the right value (1999).

Research in document event extraction and ordering have made it possible to extract such temporal structures involving events, temporal expressions, and the (unknown) document date in a document [135, 35]. While methods to perform reasoning over such structures exist [203, 204, 196, 119, 162], none of them have exploited advances in deep learning [96, 76, 68]. In particular, recently proposed Graph Convolution Networks (GCN) [48, 91] have emerged as a way to learn graph representation while encoding structural information and constraints represented by the graph. We adapt GCNs for the document dating problem and make the following contributions:

- We propose NeurdalDater, a Graph Convolution Network (GCN)-based approach for document dating. To the best of our knowledge, this is the first application of GCNs, and more broadly deep neural network-based methods, for the document dating problem.
- NeurdalDater is the first document dating approach which exploits syntactic as well temporal structure of the document, all within a principled joint model.
- Through extensive experiments on multiple real-world datasets, we demonstrate Neurdal-Dater's effectiveness over state-of-the-art baselines.

NeurdalDater's source code and datasets used in the chapter are available at <http://github.com/mallabiisc/NeurdalDater>.

## 6.2 Related Work

**Automatic Document Dating:** de Jong et al. [45] propose the first approach for automating

document dating through a statistical language model. Kanhabua and Nørvåg [86] further extend this work by incorporating semantic-based preprocessing and temporal entropy [86] based term-weighting. Chambers [32] proposes a MaxEnt based discriminative model trained on hand-crafted temporal features. He also proposes a model to learn probabilistic constraints between year mentions and the actual creation time of the document. We draw inspiration from his work for exploiting temporal reasoning for document dating. Kotsakos et al. [94] propose a purely statistical method which considers lexical similarity alongside burstiness [98] of terms for dating documents. To the best of our knowledge, NeurdalDater, our proposed method, is the first method to utilize deep learning techniques for the document dating problem.

**Event Ordering Systems:** Temporal ordering of events is a vast research topic in NLP. The problem is posed as a temporal relation classification between two given temporal entities. Machine Learned classifiers and well crafted linguistic features for this task are used in [34, 134]. D’Souza and Ng [54] use a hybrid approach by adding 437 hand-crafted rules. Chambers and Jurafsky [33], Yoshikawa et al. [227] try to classify with many more temporal constraints, while utilizing integer linear programming and Markov logic.

CAEVO, a CAscading EVent Ordering architecture [35] use sieve-based architecture [100] for temporal event ordering for the first time. They mix multiple learners according to their precision based ranks and use transitive closure for maintaining consistency of temporal graph. Mirza and Tonelli [135] recently propose CATENA (CAusal and TEmporal relation extraction from NAtural language texts), the first integrated system for the temporal and causal relations extraction between pre-annotated events and time expressions. They also incorporate sieve-based architecture which outperforms existing methods in temporal relation classification domain. We make use of CATENA for temporal graph construction in our work.

**Graph Convolutional Networks (GCN):** GCNs generalize Convolutional Neural Network (CNN) over graphs. GCN is introduced by [29], and later extended by [48] with efficient localized filter approximation in spectral domain. Kipf and Welling [91] propose a first-order approximation of localized filters through layer-wise propagation rule. GCNs over syntactic dependency trees have been recently exploited in the field of semantic-role labeling [126], neural machine translation [15], event detection [144]. In our work, we successfully use GCNs for document dating.

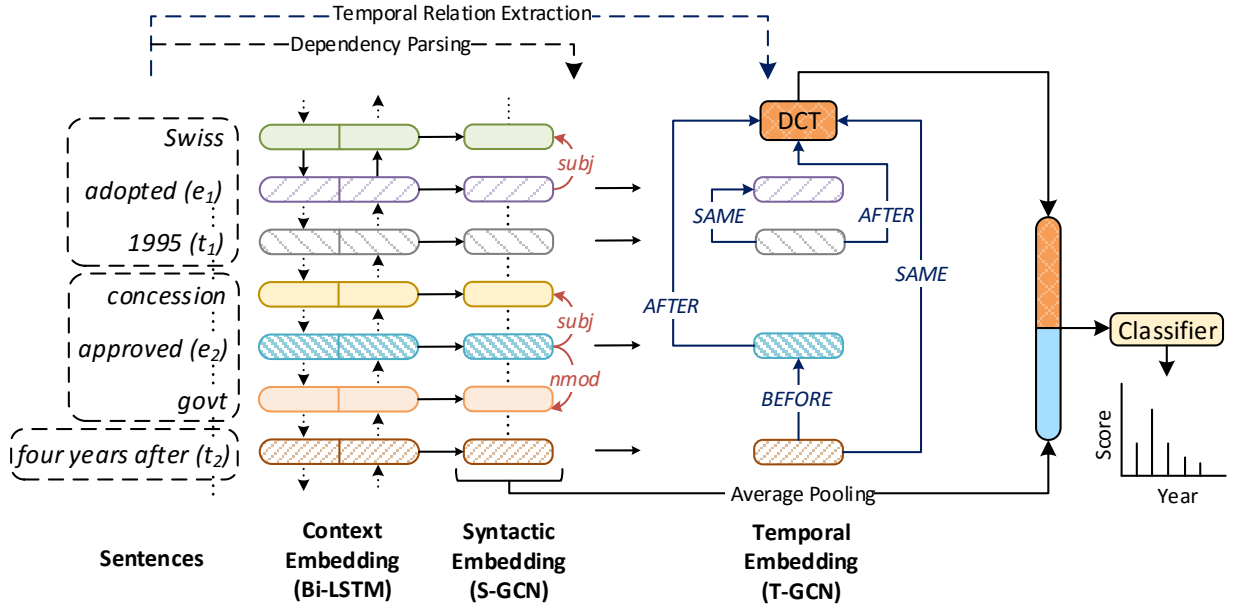


Figure 6.2: Overview of NeurdalDater. NeurdalDater exploits syntactic and temporal structure in a document to learn effective representation, which in turn are used to predict the document time. NeurdalDater uses a Bi-directional LSTM (Bi-LSTM), two Graph Convolution Networks (GCN) – one over the dependency tree and the other over the document’s temporal graph – along with a softmax classifier, all trained end-to-end jointly. Please see Section 6.3 for more details.

## 6.3 Proposed Approach: NeuralDater

### 6.3.1 Overview

The Documents Dating problem may be cast as a multi-class classification problem [94, 32]. In this section, we present an overview of NeurdalDater, the document dating system proposed in this chapter. Architectural overview of NeurdalDater is shown in Figure 6.2.

NeurdalDater is a deep learning-based multi-class classification system. It takes in a document as input and returns its predicted date as output by exploiting the syntactic and temporal structure of document.

NeurdalDater network consists of three layers which learn an embedding for the Document Creation Time (DCT) node corresponding to the document. This embedding is then fed to a softmax classifier which produces a distribution over timestamps. Following prior research [32, 94], we work with year granularity for the experiments in this chapter. We, however, note that NeuralDater can be trained for finer granularity with appropriate training data. The

NeurdalDater network is trained end-to-end using training data. We briefly present NeurdalDater’s various components below. Each component is described in greater detail in subsequent sections.

- **Context Embedding:** In this layer, NeurdalDater uses a Bi-directional LSTM (Bi-LSTM) to learn embedding for each token in the document. Bi-LSTMs have been shown to be quite effective in capturing local context inside token embeddings [188].
- **Syntactic Embedding:** In this step, NeurdalDater revises token embeddings from the previous step by running a GCN over the dependency parses of sentences in the document. We refer to this GCN as **Syntactic GCN** or **S-GCN**. While the Bi-LSTM captures immediate local context in token embeddings, S-GCN augments them by capturing syntactic context.
- **Temporal Embedding:** In this step, NeurdalDater further refines embeddings learned by S-GCN to incorporate cues from temporal structure of event and times in the document. NeurdalDater uses state-of-the-art causal and temporal relation extraction algorithm [135] for extracting temporal graph for each document. A GCN is then run over this temporal graph to refine the embeddings from the previous layer. We refer to this GCN as **Temporal GCN** or **T-GCN**. In this step, a special DCT node is introduced whose embedding is also learned by the T-GCN.
- **Classifier:** Embedding of the DCT node along with average pooled embeddings learned by S-GCN are fed to a fully connected softmax classifier which makes the final prediction about the date of the document.

Even though the previous discussion is presented in a sequential manner, the whole network is trained in a joint end-to-end manner using backpropagation. Below, we present detailed description of various components of NeurdalDater.

### 6.3.2 Context Embedding (Bi-LSTM)

Let us consider a document  $D$  with  $n$  tokens  $w_1, w_2, \dots, w_n$ . We first represent each token by a  $k$ -dimensional word embedding. For the experiments in this chapter, we use GloVe [157] embeddings. These token embeddings are stacked together to get the document representation  $\mathcal{X} \in \mathbb{R}^{n \times k}$ . We then employ a Bi-directional LSTM (Bi-LSTM) [77] on the input matrix  $\mathcal{X}$  to obtain contextual embedding for each token. After stacking contextual embedding of all these tokens, we get the new document representation matrix  $\mathcal{H}^{cntx} \in \mathbb{R}^{n \times r_{cntx}}$ . In this new

representation, each token is represented in a  $r_{cntx}$ -dimensional space. Our choice of LSTMs for learning contextual embeddings for tokens is motivated by the previous success of LSTMs in this task [188].

### 6.3.3 Syntactic Embedding (S-GCN)

While the Bi-LSTM is effective at capturing immediate local context of a token, it may not be as effective in capturing longer range dependencies among words in a sentence. For example, in Figure 6.1, we would like the embedding of token *approved* to be directly affected by *govt*, even though they are not immediate neighbors. A dependency parse may be used to capture such longer-range connections. In fact, similar features were exploited by [32] for the document dating problem. NeurdalDater captures such longer-range information by using another GCN run over the syntactic structure of the document. We describe this in detail below.

The context embedding,  $\mathcal{H}^{cntx} \in \mathbb{R}^{n \times r_{cntx}}$  learned in the previous step is used as input to this layer. For a given document, we first extract its syntactic dependency structure by applying the Stanford CoreNLP’s dependency parser [125] on each sentence in the document individually. We now employ the Graph Convolution Network (GCN) over this dependency graph using the GCN formulation presented in Section 2.1.5. We call this GCN the Syntactic GCN or S-GCN, as mentioned in Section 6.3.

Since S-GCN operates over the dependency graph and uses Equation 2.9 for updating embeddings, the number of parameters in S-GCN is directly proportional to the number of dependency edge types. Stanford CoreNLP’s dependency parser returns 55 different dependency edge types. This large number of edge types is going to significantly over-parameterize S-GCN, thereby increasing the possibility of overfitting. In order to address this, we use only three edge types in S-GCN. For each edge connecting nodes  $w_i$  and  $w_j$  in  $\mathcal{E}'$  (see Equation 2.1.5), we determine its new type  $L(w_i, w_j)$  as follows:

- $L(w_i, w_j) = \rightarrow$  if  $(w_i, w_j, l(w_i, w_j)) \in \mathcal{E}'$ , i.e., if edge is an original dependency parse edge
- $L(w_i, w_j) = \leftarrow$  if  $(w_i, w_j, l(w_i, w_j)^{-1}) \in \mathcal{E}'$ , i.e., if the edges is an inverse edge
- $L(w_i, w_j) = \top$  if  $(w_i, w_j, \top) \in \mathcal{E}'$ , i.e., if the edge is a self-loop with  $w_i = w_j$

S-GCN now estimates embedding  $h_{w_i}^{syn} \in \mathbb{R}^{r_{syn}}$  for each token  $w_i$  in the document using the formulation shown below.

$$h_{w_i}^{syn} = f \left( \sum_{w_j \in \mathcal{N}(w_i)} \left( W_{L(w_i, w_j)} h_{w_j}^{cntx} + b_{L(w_i, w_j)} \right) \right)$$

Please note S-GCN’s use of the new edge types  $L(w_i, w_j)$  above, instead of the  $l(w_i, w_j)$  types used in Equation 2.9. By stacking embeddings for all the tokens together, we get the new embedding matrix  $\mathcal{H}^{syn} \in \mathbb{R}^{n \times r_{syn}}$  representing the document.

**AveragePooling:** We obtain an embedding  $h_D^{avg}$  for the whole document by average pooling of every token representation.

$$h_D^{avg} = \frac{1}{n} \sum_{i=1}^n h_{w_i}^{syn}. \quad (6.1)$$

### 6.3.4 Temporal Embedding (T-GCN)

In this layer, NeurdalDater exploits temporal structure of the document to learn an embedding for the Document Creation Time (DCT) node of the document. First, we describe the construction of temporal graph, followed by GCN-based embedding learning over this graph.

**Temporal Graph Construction:** NeurdalDater uses Stanford’s SUTime tagger [36] for date normalization and the event extraction classifier of [35] for event detection. The annotated document is then passed to CATENA [135], current state-of-the-art temporal and causal relation extraction algorithm, to obtain a temporal graph for each document. Since our task is to predict the creation time of a given document, we supply DCT as unknown to CATENA. We hypothesize that the temporal relations extracted in absence of DCT are helpful for document dating and we indeed find this to be true, as shown in Section 6.4.2. Temporal graph is a directed graph, where nodes correspond to events, time mentions, and the Document Creation Time (DCT). Edges in this graph represent causal and temporal relationships between them. Each edge is attributed with a label representing the type of the temporal relation. CATENA outputs 9 different types of temporal relations, out of which we selected five types, viz., *AFTER*, *BEFORE*, *SAME*, *INCLUDES*, and *IS\_INCLUDED*. The remaining four types were ignored as they were substantially infrequent.

Please note that the temporal graph may involve only a small number of tokens in the document. For example, in the temporal graph in Figure 6.2, there are a total of 5 nodes: two temporal expression nodes (*1995* and *four years after*), two event nodes (*adopted* and *approved*), and a special DCT node. This graph also consists of temporal relation edges such as (*four years after*, *approved*, *BEFORE*).

**Temporal Graph Convolution:** NeurdalDater employs a GCN over the temporal graph constructed above. We refer to this GCN as the Temporal GCN or T-GCN, as mentioned in Section 6.3. T-GCN is based on the GCN formulation presented in Section 2.1.5. Unlike S-GCN, here we consider label and direction specific parameters as the temporal graph consists of only five types of edges.

Datasets	# Docs	Start Year	End Year
APW	675k	1995	2010
NYT	647k	1987	1996

Table 6.1: Details of datasets used. Please see Section 6.4.1 for details.

Let  $n_T$  be the number of nodes in the temporal graph. Starting with  $\mathcal{H}^{syn}$  (Section 6.3.3), T-GCN learns a  $r_{temp}$ -dimensional embedding for each node in the temporal graph. Stacking all these embeddings together, we get the embedding matrix  $\mathcal{H}^{temp} \in \mathbb{R}^{n_T \times r_{temp}}$ . T-GCN embeds the temporal constraints induced by the temporal graph in  $h_{DCT}^{temp} \in \mathbb{R}^{r_{temp}}$ , embedding of the DCT node of the document.

### 6.3.5 Classifier

Finally, the DCT embedding  $h_{DCT}^{temp}$  and average-pooled syntactic representation  $h_D^{avg}$  (see Equation 6.1) of document  $D$  are concatenated and fed to a fully connected feed forward network followed by a softmax. This allows the NeurdalDater to exploit context, syntactic, and temporal structure of the document to predict the final document date  $y$ .

$$\begin{aligned} h_D^{avg+temp} &= [h_{DCT}^{temp} ; h_D^{avg}] \\ p(y|D) &= \text{Softmax}(W \cdot h_D^{avg+temp} + b). \end{aligned}$$

## 6.4 Experiments

### 6.4.1 Experimental Setup

**Datasets:** We experiment on Associated Press Worldstream (APW) and New York Times (NYT) sections of Gigaword corpus [152]. The original dataset contains around 3 million documents of APW and 2 million documents of NYT from span of multiple years. From both sections, we randomly sample around 650k documents while maintaining balance among years. Documents belonging to years with substantially fewer documents are omitted. Details of the dataset can be found in Table 6.1. For train, test and validation splits, the dataset was randomly divided in 80:10:10 ratio.

**Evaluation Criteria:** Given a document, the model needs to predict the year in which the document was published. We measure performance in terms of overall accuracy of the model.

**Baselines:** For evaluating NeurdalDater, we compared against the following methods:

Method	APW	NYT
BurstySimDater	45.9	38.5
MaxEnt-Time+NER	52.5	42.3
MaxEnt-Joint	52.5	42.5
MaxEnt-Uni-Time	57.5	50.5
CNN	56.3	50.4
NeurdalDater	<b>64.1</b>	<b>58.9</b>

Table 6.2: Accuracies of different methods on APW and NYT datasets for the document dating problem (higher is better). NeurdalDater significantly outperforms all other competitive baselines. This is our main result. Please see Section 6.4.2.1 for more details.

- **BurstySimDater** Kotsakos et al. [94]: This is a purely statistical method which uses lexical similarity and term burstiness [98] for dating documents in arbitrary length time frame. For our experiments, we took the time frame length as 1 year. Please refer to [94] for more details.
- **MaxEnt-Time-NER**: Maximum Entropy (MaxEnt) based classifier trained on hand-crafted temporal and Named Entity Recognizer (NER) based features. More details in [32].
- **MaxEnt-Joint**: Refers to MaxEnt-Time-NER combined with year mention classifier as described in [32].
- **MaxEnt-Uni-Time**: MaxEnt based discriminative model which takes bag-of-words representation of input document with normalized time expression as its features.
- **CNN**: A Convolution Neural Network (CNN) [99] based text classification model proposed by [88], which attained state-of-the-art results in several domains.
- **NeurdalDater**: Our proposed method, refer Section 6.3.

**Hyperparameters:** By default, edge gating (Section 2.1.5) is used in all GCNs. The parameter  $K$  represents the number of layers in T-GCN (Section 6.3.4). We use 300-dimensional GloVe embeddings and 128-dimensional hidden state for both GCNs and BiLSTM with 0.8 dropout. We used Adam [89] with 0.001 learning rate for training.

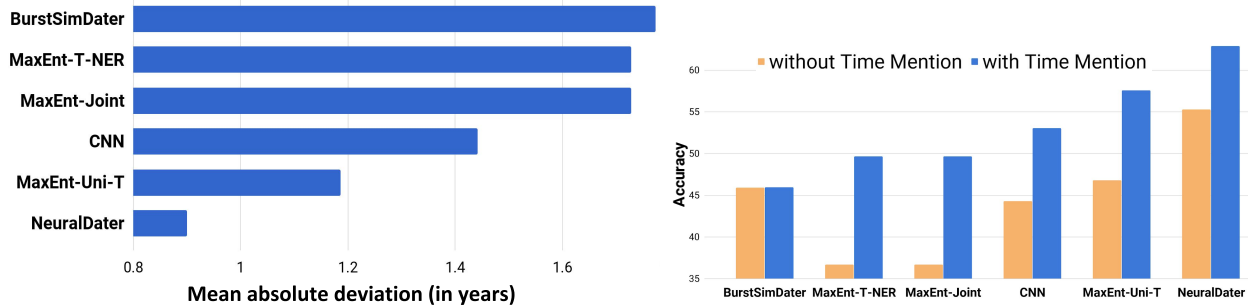
Method	Accuracy
T-GCN	57.3
S-GCN + T-GCN ( $K = 1$ )	57.8
S-GCN + T-GCN ( $K = 2$ )	58.8
S-GCN + T-GCN ( $K = 3$ )	<b>59.1</b>
Bi-LSTM	58.6
Bi-LSTM + CNN	59.0
Bi-LSTM + T-GCN	60.5
Bi-LSTM + S-GCN + T-GCN (no gate)	62.7
Bi-LSTM + S-GCN + T-GCN ( $K = 1$ )	<b>64.1</b>
Bi-LSTM + S-GCN + T-GCN ( $K = 2$ )	63.8
Bi-LSTM + S-GCN + T-GCN ( $K = 3$ )	63.3

Table 6.3: Accuracies of different ablated methods on the APW dataset. Overall, we observe that incorporation of context (Bi-LSTM), syntactic structure (S-GCN) and temporal structure (T-GCN) in NeurdalDater achieves the best performance. Please see Section 6.4.2.1 for details.

## 6.4.2 Results

### 6.4.2.1 Performance Comparison

In order to evaluate the effectiveness of NeurdalDater, our proposed method, we compare it against existing document dating systems and text classification models. The final results are summarized in Table 6.2. Overall, we find that NeurdalDater outperforms all other methods with a significant margin on both datasets. Compared to the previous state-of-the-art in document dating, BurstySimDater [94], we get 19% average absolute improvement in accuracy across both datasets. We observe only a slight gain in the performance of MaxEnt-based model (MaxEnt-Time+NER) of [32] on combining with temporal constraint reasoner (MaxEnt-Joint). This may be attributed to the fact that the model utilizes only year mentions in the document, thus ignoring other relevant signals which might be relevant to the task. BurstySimDater performs considerably better in terms of precision compared to the other baselines, although it significantly underperforms in accuracy. We note that NeuralDater outperforms all these prior models both in terms of precision and accuracy. We find that even generic deep-learning based text classification models, such as CNN [88], are quite effective for the problem. However, since such a model doesn't give specific attention to temporal features in the document, its performance remains limited. From Figure 6.3a, we observe that NeurdalDater's top prediction achieves on average the lowest deviation from the true year.



(a) Mean absolute deviation (in years; lower is better) between a model’s top prediction and the true odds on dating documents with and without year in the APW dataset. We find that Neurdal-Dater, the proposed method, achieves the least deviation. Please see Section 6.4.2.1 for details.

(b) Evaluating performance of different methods on dating documents with and without time mentions. Please see Section 6.4.2.3 for details.

#### 6.4.2.2 Ablation Comparisons

For demonstrating the efficacy of GCNs and BiLSTM for the problem, we evaluate different ablated variants of NeurdalDater on the APW dataset. Specifically, we validate the importance of using syntactic and temporal GCNs and the effect of eliminating BiLSTM from the model. Overall results are summarized in Table 6.3. The first block of rows in the table corresponds to the case when BiLSTM layer is excluded from NeurdalDater, while the second block denotes the case when BiLSTM is included. We also experiment with multiple stacked layers of T-GCN (denoted by  $K$ ) to observe its effect on the performance of the model.

We observe that embeddings from Syntactic GCN (S-GCN) are much better than plain GloVe embeddings for T-GCN as S-GCN encodes the syntactic neighborhood information in event and time embeddings which makes them more relevant for document dating task.

Overall, we observe that including BiLSTM in the model improves performance significantly. Single BiLSTM model outperforms all the models listed in the first block of Table 6.3. Also, some gain in performance is observed on increasing the number of T-GCN layers ( $K$ ) in absence of BiLSTM, although the same does not follow when BiLSTM is included in the model. This observation is consistent with [126], as multiple GCN layers become redundant in the presence of BiLSTM. We also find that eliminating edge gating from our best model deteriorates its overall performance.

In summary, these results validate our thesis that joint incorporation of syntactic and temporal structure of a document in NeurdalDater results in improved performance.

#### 6.4.2.3 Discussion and Error Analysis

In this section, we list some of our observations while trying to identify pros and cons of NeurdalDater, our proposed method. We divided the development split of the APW dataset into two sets – those with and without any mention of time expressions (year). We apply NeurdalDater and other methods to these two sets of documents and report accuracies in Figure 6.3b. We find that overall, NeurdalDater performs better in comparison to the existing baselines in both scenarios. Even though the performance of NeurdalDater degrades in the absence of time mentions, its performance is still the best relatively. Based on other analysis, we find that NeurdalDater fails to identify timestamp of documents reporting local infrequent incidents without explicit time mention. NeurdalDater becomes confused in the presence of multiple misleading time mentions; it also loses out on documents discussing events which are outside the time range of the text on which the model was trained. In future, we plan to eliminate these pitfalls by incorporating additional signals from Knowledge Graphs about entities mentioned in the document. We also plan to utilize free text temporal expression [97] in documents for improving performance on this problem.

## 6.5 Conclusion

We propose NeurdalDater, a Graph Convolutional Network (GCN) based method for document dating which exploits syntactic and temporal structures in the document in a principled way. To the best of our knowledge, this is the first application of deep learning techniques for the problem of document dating. Through extensive experiments on real-world datasets, we demonstrate the effectiveness of NeurdalDater over existing state-of-the-art approaches. We are hopeful that the representation learning techniques explored in this chapter will inspire further development and adoption of such techniques in the temporal information processing research community.

# Chapter 7

## Incorporating Syntactic and Semantic Information in Word Embeddings using Graph Convolutional Networks

### 7.1 Introduction

As we saw in the last chapter, Graph Convolutional Networks prove to be very effective for exploiting different graph structure in NLP. Here, we utilize them for learning word representation. Representing words as real-valued vectors is an effective and widely adopted technique in NLP. Such representations capture properties of words based on their usage and allow them to generalize across tasks. Meaningful word embeddings have been shown to improve performance on several relevant tasks, such as named entity recognition (NER) [18], parsing [177], and part-of-speech (POS) tagging [121]. Using word embeddings for initializing Deep Neural Networks has also been found to be quite useful [42, 84, 183].

Most popular methods for learning word embeddings are based on the distributional hypothesis, which utilizes the co-occurrence statistics from *sequential* context of words for learning word representations [130, 157]. More recently, this approach has been extended to include syntactic contexts [103] derived from dependency parse of text. Higher order dependencies have also been exploited by Komninos and Manandhar [93], Li et al. [104]. Syntax-based embeddings encode functional similarity (in-place substitutable words) rather than topical similarity (topically related words) which provides an advantage on specific tasks like question classification [93]. However, current approaches incorporate syntactic context by concatenating words with their dependency relations. For instance, in Figure 7.1 *scientists\_subj*, *water\_obj*, and *mars\_nmod* needs to be included as a part of vocabulary for utilizing the dependency context

of *discover*. This severely expands the vocabulary, thus limiting the scalability of models on large corpora. For instance, in Levy and Goldberg [103] and Komninos and Manandhar [93], the context vocabulary explodes to around 1.3 million for learning embeddings of 220k words.

Incorporating relevant signals from semantic knowledge sources such as WordNet [132], FrameNet [8], and Paraphrase Database (PPDB) [156] has been shown to improve the quality of word embeddings. Recent works utilize these by incorporating them in a neural language modeling objective function [228, 3], or as a post-processing step [56, 140]. Although existing approaches improve the quality of word embeddings, they require explicit modification for handling different types of semantic information.

Recently proposed Graph Convolutional Networks (GCN) [47, 90] have been found to be useful for encoding structural information in graphs. Even though GCNs have been successfully employed for several NLP tasks such as machine translation [15], semantic role labeling [126], document dating [198] and text classification [223], they have so far not been used for learning word embeddings, especially leveraging cues such as syntactic and semantic information. GCNs provide flexibility to represent diverse syntactic and semantic relationships between words all within one framework, without requiring relation-specific special handling as in previous methods. Recognizing these benefits, we make the following contributions in this chapter.

1. We propose SynGCN, a Graph Convolution based method for learning word embeddings. Unlike previous methods, SynGCN utilizes syntactic context for learning word representations without increasing vocabulary size.
2. We also present SemGCN, a framework for incorporating diverse semantic knowledge (e.g., synonymy, antonymy, hyponymy, etc.) in learned word embeddings, without requiring relation-specific special handling as in previous methods.
3. Through experiments on multiple intrinsic and extrinsic tasks, we demonstrate that our proposed methods obtain substantial improvement over state-of-the-art approaches, and also yield an advantage when used in conjunction with methods such as ELMo [159].

## 7.2 Related Work

**Word Embeddings:** Recently, there has been much interest in learning meaningful word representations such as neural language modeling [19] based continuous-bag-of-words (CBOW) and skip-gram (SG) models [130]. This is further extended by Pennington et al. [157] which learns embeddings by factorizing word co-occurrence matrix to leverage global statistical information. Other formulations for learning word embeddings include multi-task learning [42] and ranking frameworks [82].

**Syntax-based Embeddings:** Dependency parse context based word embeddings is first introduced by Levy and Goldberg [103]. They allow encoding syntactic relationships between words and show improvements on tasks where functional similarity is more relevant than topical similarity. The inclusion of syntactic context is further enhanced through second-order [93] and multi-order [104] dependencies. However, in all these existing approaches, the word vocabulary is severely expanded for incorporating syntactic relationships.

**Incorporating Semantic Knowledge Sources:** Semantic relationships such as *synonymy*, *antonymy*, *hyponymy*, etc. from several semantic sources have been utilized for improving the quality of word representations. Existing methods either exploit them jointly [215, 87, 3] or as a post-processing step [56, 140]. SynGCN falls under the latter category and is more effective at incorporating semantic constraints (Section 7.4.2.2 and 7.4.2.3).

**Graph Convolutional Networks:** In this chapter, we use the first-order formulation of GCNs via a layer-wise propagation rule as proposed by [90]. Recently, some variants of GCNs have also been proposed [218, 201]. A detailed description of GCNs and their applications can be found in Bronstein et al. [27]. In NLP, GCNs have been utilized for semantic role labeling [126], machine translation [15], and relation extraction [200]. Recently, Yao et al. [223] use GCNs for text classification by jointly embedding words and documents. However, their learned embeddings are task specific whereas in our work we aim to learn task agnostic word representations.

## 7.3 Proposed Methods: SynGCN and SemGCN

### 7.3.1 Overview

The task of learning word representations in an unsupervised setting can be formulated as follows: Given a text corpus, the aim is to learn a  $d$ -dimensional embedding for each word in the vocabulary. Most of the distributional hypothesis based approaches only utilize sequential context for each word in the corpus. However, this becomes suboptimal when the relevant context words lie beyond the window size. For instance in Figure 7.1, a relevant context word *discover* for *Mars* is missed if the chosen window size is less than 3. On the contrary, a large window size might allow irrelevant words to influence word embeddings negatively.

Using dependency based context helps to alleviate this problem. However, all existing syntactic context based methods [103, 93, 104] severely expand vocabulary size (as discussed in Section 7.1) which limits their scalability to a large corpus. To eliminate this drawback, we propose SynGCN which employs Graph Convolution Networks to better encode syntactic information in embeddings. We prefer GCNs over other graph encoding architectures such as

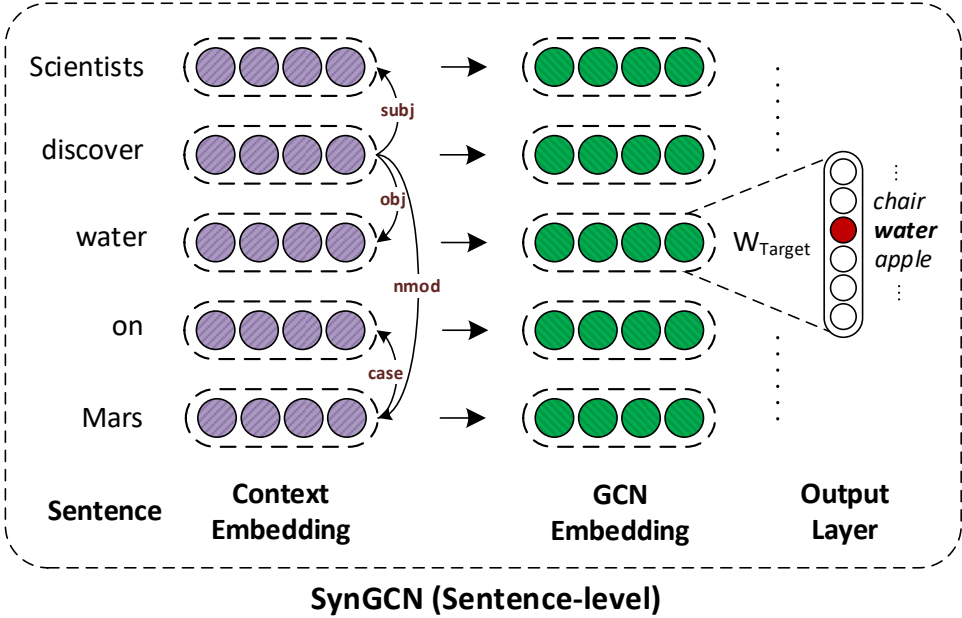


Figure 7.1: Overview of SynGCN: SynGCN employs Graph Convolution Network for utilizing dependency context for learning word embeddings. For each word in vocabulary, the model learns its representation by aiming to predict each word based on its dependency context encoded using GCNs. Please refer Section 7.3.2 for more details.

Tree LSTM [189] as GCNs do not restrict graphs to be trees and have been found to be more effective at capturing global information [233]. Moreover, they give substantial speedup as they do not involve recursive operations which are difficult to parallelize. The overall architecture is shown in Figure 7.1, for more details refer to Section 7.3.2.

Enriching word embeddings with semantic knowledge helps to improve their quality for several NLP tasks. Existing approaches are either incapable of utilizing these diverse relations or need to be explicitly modeled for exploiting them. In this chapter, we propose SemGCN which automatically learns to utilize multiple semantic constraints by modeling them as different edge types. It can be used as a post-processing method similar to Faruqui et al. [56], Mrkšić et al. [140]. We describe it in more detail in Section 7.3.3.

### 7.3.2 SynGCN

In this section, we provide a detailed description of our proposed method, SynGCN. Following Mikolov et al. [131], Levy and Goldberg [103], Komninos and Manandhar [93], we separately define target and context embeddings for each word in the vocabulary as parameters in the model. For a given sentence  $s = (w_1, w_2, \dots, w_n)$ , we first extract its dependency parse graph  $\mathcal{G}_s = (\mathcal{V}_s, \mathcal{E}_s)$  using Stanford CoreNLP parser [125]. Here,  $\mathcal{V}_s = \{w_1, w_2, \dots, w_n\}$  and  $\mathcal{E}_s$  denotes

the labeled directed dependency edges of the form  $(w_i, w_j, l_{ij})$ , where  $l_{ij}$  is the dependency relation of  $w_i$  to  $w_j$ .

Similar to Mikolov et al. [131]’s continuous-bag-of-words (CBOW) model, which defines the context of a word  $w_i$  as  $\mathcal{C}_{w_i} = \{w_{i+j} : -c \leq j \leq c, j \neq 0\}$  for a window of size  $c$ , we define the context as its neighbors in  $\mathcal{G}_s$ , i.e.,  $\mathcal{C}_{w_i} = \mathcal{N}(w_i)$ . Now, unlike CBOW which takes the sum of the context embedding of words in  $\mathcal{C}_{w_i}$  to predict  $w_i$ , we apply directed Graph Convolution Network (as defined in Section 2.1.5) on  $\mathcal{G}_s$  with context embeddings of words in  $s$  as input features. Thus, for each word  $w_i$  in  $s$ , we obtain a representation  $h_i^{k+1}$  after  $k$ -layers of GCN using Equation 2.9 which we reproduce below for ease of readability (with one exception as described below).

$$h_i^{k+1} = f \left( \sum_{j \in \mathcal{N}(i)} g_{l_{ij}}^k \times \left( W_{l_{ij}}^k h_j^k + b_{l_{ij}}^k \right) \right)$$

Please note that unlike in Equation 2.9, we use  $\mathcal{N}(i)$  instead of  $\mathcal{N}_+(i)$  in SynGCN, i.e., we do not include self-loops in  $\mathcal{G}_s$ . This helps to avoid overfitting to the initial embeddings, which is undesirable in the case of SynGCN as it uses random initialization. We note that similar strategy has been followed by Mikolov et al. [131]. Furthermore, to handle erroneous edges in automatically constructed dependency parse graph, we perform edge-wise gating (Section 2.1.5) to give importance to relevant edges and suppress the noisy ones. The embeddings obtained are then used to calculate the loss as described in Section 7.3.4.

SynGCN utilizes *syntactic context* to learn more meaningful word representations. We validate this in Section 7.4.2.1. Note that, the word vocabulary remains unchanged during the entire learning process, this makes SynGCN more scalable compared to the existing approaches.

Note that, SynGCN is a generalization of CBOW model, as shown below.

**Theorem 7.1** *SynGCN is a generalization of Continuous-bag-of-words (CBOW) model.*

**Proof:** The reduction can be obtained as follows. For a given sentence  $s$ , take the neighborhood of each word  $w_i$  in  $\mathcal{G}_s$  as its sequential context, i.e.,  $\mathcal{N}(w_i) = \{w_{i+j} : -c \leq j \leq c, j \neq 0\} \forall w_i \in s$ . Now, if the number of GCN layers are restricted to 1 and the activation function is taken as identity ( $f(x) = x$ ), then Equation 2.9 reduces to

$$h_i = \sum_{-c \leq j \leq c, j \neq 0} \left( g_{l_{ij}} \times \left( W_{l_{ij}} h_j + b_{l_{ij}}^k \right) \right).$$

Finally,  $W_{l_{ij}}^k$  and  $b_{l_{ij}}^k$  can be fixed to an identity matrix (**I**) and a zero vector (**0**), respectively,

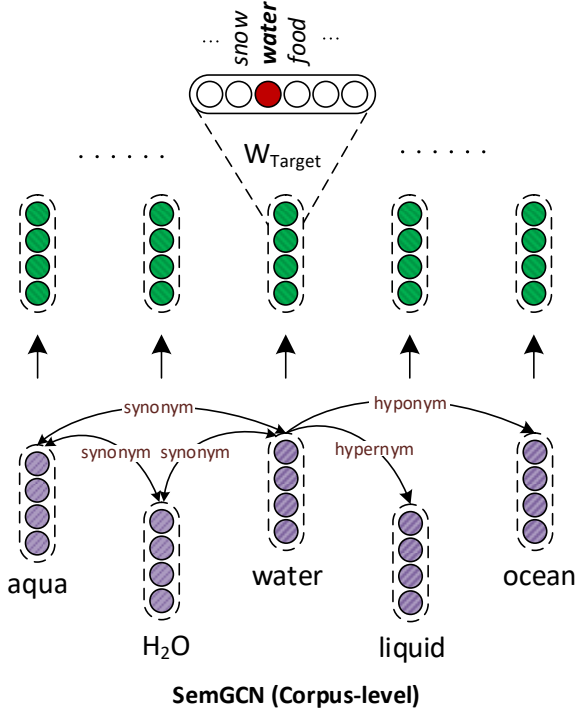


Figure 7.2: Overview of SemGCN, our proposed Graph Convolution based framework for incorporating diverse semantic information in learned embeddings. Double-headed edges denote two edges in both directions. Please refer to Section 7.3.3 for more details.

and edge-wise gating ( $g_{l_{ij}}$ ) can be set to 1. This gives

$$h_i = \sum_{-c \leq j \leq c, j \neq 0} (\mathbf{I} \cdot h_j + \mathbf{0}) = \sum_{-c \leq j \leq c, j \neq 0} h_j,$$

which is the hidden layer equation of CBOW model.  $\square$

### 7.3.3 SemGCN

In this section, we propose another Graph Convolution based framework, SemGCN, for incorporating semantic knowledge in pre-trained word embeddings. Most of the existing approaches like Faruqui et al. [56], Mrkšić et al. [140] are restricted to handling symmetric relations like *synonymy* and *antonymy*. On the other hand, although recently proposed [3] is capable of handling asymmetric information, it still requires manually defined relation strength function which can be labor intensive and suboptimal.

SemGCN is capable of incorporating both symmetric as well as asymmetric information jointly. Unlike SynGCN, SemGCN operates on a corpus-level directed labeled graph with words as nodes and edges representing semantic relationship among them from different sources.

For instance, in Figure 7.2, semantic relations such as *hyponymy*, *hypernymy* and *synonymy* are represented together in a single graph. Symmetric information is handled by including a directed edge in both directions. Given the corpus level graph  $\mathcal{G}$ , the training procedure is similar to that of SynGCN, i.e., predict the word  $w$  based on its neighbors in  $\mathcal{G}$ . Inspired by Faruqui et al. [56], we preserve the semantics encoded in pre-trained embeddings by initializing both target and context embeddings with given word representations and keeping target embeddings fixed during training. SemGCN uses Equation 2.9 to update node embeddings. Please note that in this case  $\mathcal{N}_+(v)$  is used as the neighborhood definition to preserve the initial learned representation of the words.

### 7.3.4 Training Details

Given the GCN representation ( $h_t$ ) of a word ( $w_t$ ), the training objective of SynGCN and SemGCN is to predict the target word given its neighbors in the graph. Formally, for each method we maximize the following objective<sup>1</sup>.

$$E = \sum_{t=1}^{|V|} \log P(w_t | w_1^t, w_2^t \dots w_{N_t}^t)$$

where,  $w_t$  is the target word and  $w_1^t, w_2^t \dots w_{N_t}^t$  are its neighbors in the graph. The probability  $P(w_t | w_1^t, w_2^t \dots w_{N_t}^t)$  is calculated using the softmax function, defined as

$$P(w_t | w_1^t, w_2^t \dots w_{N_t}^t) = \frac{\exp(v_{w_t}^T h_t)}{\sum_{i=1}^{|V|} \exp(v_{w_i}^T h_t)}.$$

Hence,  $E$  reduces to

$$E = \sum_{t=1}^{|V|} \left( v_{w_t}^T h_t - \log \sum_{i=1}^{|V|} \exp(v_{w_i}^T h_t) \right), \quad (7.1)$$

where,  $h_t$  is the GCN representation of the target word  $w_t$  and  $v_{w_t}$  is its target embedding.

The second term in Equation 7.1 is computationally expensive as the summation needs to be taken over the entire vocabulary. This can be overcome using several approximations like noise-contrastive estimation [71] and hierarchical softmax [139]. In our methods, we use negative sampling as used by Mikolov et al. [131].

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<sup>1</sup>We also experimented with joint SynGCN and SemGCN model but our preliminary experiments gave suboptimal performance as compared to the sequential model. This can be attributed to the fact that syntactic information is orders of magnitude greater than the semantic information available. Hence, the semantic constraints are not effectively utilized. We leave the analysis of the joint model as a future work.

## 7.4 Experiments

### 7.4.1 Experimental Setup

#### 7.4.1.1 Dataset and Training

In our experiments, we use Wikipedia<sup>1</sup> corpus for training the models. After discarding too long and too short sentences, we get an average sentence length of nearly 20 words. The corpus consists of 57 million sentences with 1.1 billion tokens and 1 billion syntactic dependencies.

#### 7.4.1.2 Baselines

For evaluating SynGCN (Section 7.3.2), we compare against the following baselines:

- **Word2vec** is continuous-bag-of-words model originally proposed by Mikolov et al. [131].
- **GloVe** [157], a log-bilinear regression model which leverages global co-occurrence statistics of corpus.
- **Deps** [103] is a modification of skip-gram model which uses dependency context in place of sequential context.
- **EXT** [93] is an extension of Deps which utilizes second-order dependency context features.

SemGCN (Section 7.3.3) model is evaluated against the following methods:

- **Retro-fit** [56] is a post-processing procedure which uses similarity constraints from semantic knowledge sources.
- **Counter-fit** [140], a method for injecting both antonym and synonym constraints into word embeddings.
- **JointReps** [3], a joint word representation learning method which simultaneously utilizes the corpus and KB.

#### 7.4.1.3 Evaluation method

To evaluate the effectiveness of our proposed methods, we compare them against the baselines on the following intrinsic and extrinsic tasks<sup>2</sup>:

- **Intrinsic Tasks:**

**Word Similarity** is the task of evaluating closeness between semantically similar words. Following Komninos and Manandhar [93], Pennington et al. [157], we evaluate on Simlex-999 [75], WS353 [59], and RW [120] datasets.

**Concept Categorization** involves grouping nominal concepts into natural categories.

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<sup>1</sup><https://dumps.wikimedia.org/enwiki/20180301/>

<sup>2</sup>Details of hyperparameters are in supplementary.

Method	Word Similarity				Concept Categorization				Word Analogy	
	WS353S	WS353R	SimLex999	RW	AP	Battig	BLESS	ESSLI	SemEval2012	MSR
Word2vec	71.4	52.6	38.0	30.0	63.2	43.3	77.8	63.0	18.9	44.0
GloVe	69.2	<b>53.4</b>	36.7	29.6	58.0	41.3	80.0	59.3	18.7	45.8
Deps	65.7	36.2	39.6	33.0	61.8	41.7	65.9	55.6	22.9	40.3
EXT	69.6	44.9	43.2	18.6	52.6	35.0	65.2	66.7	21.8	18.8
SynGCN	<b>73.2</b>	45.7	<b>45.5</b>	<b>33.7</b>	<b>69.3</b>	<b>45.2</b>	<b>85.2</b>	<b>70.4</b>	<b>23.4</b>	<b>52.8</b>

Table 7.1: **SynGCN Intrinsic Evaluation:** Performance on word similarity (Spearman correlation), concept categorization (cluster purity), and word analogy (Spearman correlation). Overall, SynGCN outperforms other existing approaches in 9 out of 10 settings. Please refer to Section 7.4.2.1 for more details.

For instance, *tiger* and *elephant* should belong to *mammal* class. In our experiments, we evaluate on AP [2], Battig [12], BLESS [13], ESSLI [14] datasets.

**Word Analogy** task is to predict word  $b_2$ , given three words  $a_1$ ,  $a_2$ , and  $b_1$ , such that the relation  $b_1 : b_2$  is same as the relation  $a_1 : a_2$ . We compare methods on MSR [131] and SemEval-2012 [85].

- **Extrinsic Tasks:**

**Named Entity Recognition (NER)** is the task of locating and classifying entity mentions into categories like *person*, *organization* etc. We use Lee et al. [101]’s model on CoNLL-2003 dataset [192] for evaluation.

**Question Answering** in Stanford Question Answering Dataset (**SQuAD**) [163] involves identifying answer to a question as a segment of text from a given passage. Following Peters et al. [159], we evaluate using Clark and Gardner [41]’s model.

**Part-of-speech (POS) tagging** aims at associating with each word, a unique tag describing its syntactic role. For evaluating word embeddings, we use Lee et al. [101]’s model on Penn Treebank POS dataset [127].

**Co-reference Resolution (Coref)** involves identifying all expressions that refer to the same entity in the text. To inspect the effect of embeddings, we use Lee et al. [101]’s model on CoNLL-2012 shared task dataset [160].

## 7.4.2 Results

In this section, we attempt to answer the following questions.

- Q1. Does SynGCN learn better word embeddings than existing approaches? (Section 7.4.2.1)
- Q2. Does SemGCN effectively handle diverse semantic information as compared to other methods? (Section 7.4.2.2)

Method	POS	SQuAD	NER	Coref
Word2vec	95.0±0.1	78.5±0.3	89.0±0.2	65.1±0.3
GloVe	94.6±0.1	78.2±0.2	89.1±0.1	64.9±0.2
Deps	95.0±0.1	77.8±0.3	88.6±0.3	64.8±0.1
EXT	94.9±0.2	<b>79.6±0.1</b>	88.0±0.1	64.8±0.1
SynGCN	<b>95.4±0.1</b>	<b>79.6±0.2</b>	<b>89.5±0.1</b>	<b>65.8±0.1</b>

Table 7.2: **SynGCN Extrinsic Evaluation:** Comparison on parts-of-speech tagging (POS), question answering (SQuAD), named entity recognition (NER), and co-reference resolution (Coref). SynGCN performs comparable or outperforms all existing approaches on all tasks. Refer Section 7.4.2.1 for details.

- Q3. How does SemGCN perform compared to other methods when provided with the same semantic constraints? (Section 7.4.2.3)
- Q4. Does dependency context based embedding encode complementary information compared to ELMo? (Section 7.4.2.4)

#### 7.4.2.1 SynGCN Evaluation

The evaluation results on intrinsic tasks – word similarity, concept categorization, and analogy – are summarized in Table 7.1. We report Spearman correlation for word similarity and analogy tasks and cluster purity for concept categorization task. Overall, we find that SynGCN, our proposed method, outperforms all the existing word embedding approaches in 9 out of 10 settings. The inferior performance of SynGCN and other dependency context based methods on WS353R dataset compared to sequential context based methods is consistent with the observation reported in Levy and Goldberg [103], Komninos and Manandhar [93]. This is because the syntactic context based embeddings capture functional similarity rather than topical similarity (as discussed in Section 7.1). On average, we obtain around 1.5%, 5.7% and 7.5% absolute increase in performance on word similarity, concept categorization and analogy tasks compared to the best performing baseline. The results demonstrate that the learned embeddings from SynGCN more effectively capture semantic and syntactic properties of words.

We also evaluate the performance of different word embedding approaches on the downstream tasks as defined in Section 7.4.1.3. The experimental results are summarized in Table 7.2. Overall, we find that SynGCN either outperforms or performs comparably to other methods on all four tasks. Compared to the sequential context based methods, dependency based methods perform superior at question answering task as they effectively encode syntactic information. This is consistent with the observation of Peters et al. [159].

Init Embeddings (=X)	Word2vec			GloVe			Deps			EXT			SynGCN		
	WS353	AP	MSR												
Datasets															
Performance of X	63.0	63.2	44.0	58.0	60.4	45.8	55.6	64.2	40.3	59.3	53.5	18.8	61.7	69.3	52.8
Retro-fit (X,1)	63.4	<b>67.8</b>	<b>46.7</b>	58.5	61.1	<b>47.2</b>	54.8	64.7	41.0	61.6	55.1	40.5	61.2	67.1	51.4
Counter-fit (X,2)	60.3	62.9	31.4	53.7	62.5	29.6	46.9	60.4	33.4	52.0	54.4	35.8	55.2	66.4	31.7
JointReps (X,4)	60.9	61.1	28.5	59.2	55.5	37.6	54.8	58.7	38.0	58.8	54.8	20.6	60.9	68.2	24.9
SemGCN (X,4)	<b>64.8</b>	<b>67.8</b>	36.8	<b>63.3</b>	<b>63.2</b>	44.1	<b>62.3</b>	<b>69.3</b>	<b>41.1</b>	<b>62.9</b>	<b>67.1</b>	<b>52.1</b>	<b>65.3</b>	<b>69.3</b>	<b>54.4</b>

Table 7.3: **SemGCN Intrinsic Evaluation:** Evaluation of different methods for incorporating diverse semantic constraints initialized using various pre-trained embeddings (X). M(X, R) denotes the fine-tuned embeddings using method M taking X as initialization embeddings. R denotes the type of semantic relations used as defined in Section 7.4.2.2. SemGCN outperforms other methods in 13 our of 15 settings. SemGCN with SynGCN gives the best performance across all tasks (highlighted using  $\boxed{\cdot}$ ). Please refer Section 7.4.2.2 for details.

Method	POS	SQuAD	NER	Coref
X = SynGCN	95.4±0.1	79.6±0.2	<b>89.5±0.1</b>	65.8±0.1
Retro-fit (X,1)	94.8±0.1	79.6±0.1	88.8±0.1	66.0±0.2
Counter-fit (X,2)	94.7±0.1	79.8±0.1	88.3±0.3	65.7±0.3
JointReps (X,4)	95.4±0.1	79.4±0.3	89.1±0.3	65.6±0.1
SemGCN (X,4)	<b>95.5±0.1</b>	<b>80.4±0.1</b>	<b>89.5±0.1</b>	<b>66.1±0.1</b>

Table 7.4: **SemGCN Extrinsic Evaluation:** Comparison of different methods for incorporating diverse semantic constraints in SynGCN embeddings on all extrinsic tasks. Refer Section 7.4.2.2 for details.

#### 7.4.2.2 Evaluation with Diverse Semantic Information

In this section, we compare SemGCN against the methods listed in Section 7.4.1.2 for incorporating diverse semantic information in pre-trained embeddings. We use *hypernym*, *hyponym*, and *antonym* relations from WordNet, and *synonym* relations from PPDB as semantic information. For each method, we provide the semantic information that it can utilize, e.g., Retro-fit can only make use of *synonym* relation<sup>1</sup>. In our results, M(X, R) denotes the fine-tuned embeddings obtained using method M while taking X as initialization embeddings. R denotes the types of semantic information used as defined below.

- **R=1:** Only synonym information.
- **R=2:** Synonym and antonym information.
- **R=4:** Synonym, antonym, hypernym and hyponym information.

For instance, Counter-fit (GloVe, 2) represents GloVe embeddings fine-tuned by Counter-fit using synonym and antonym information.

<sup>1</sup>Experimental results controlling for semantic information are provided in Section 7.4.2.3.

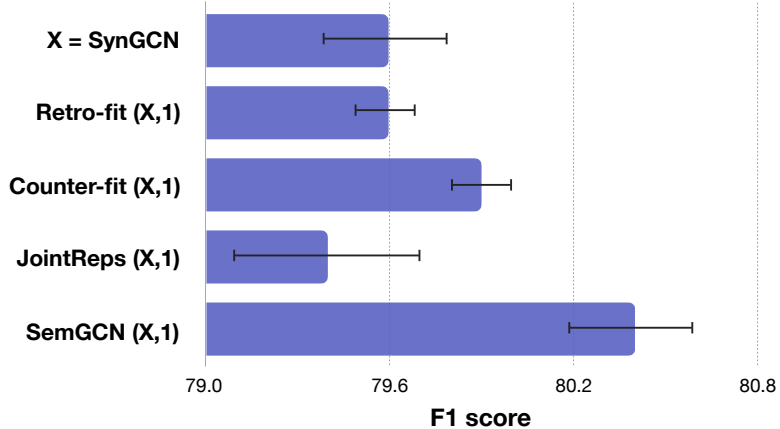


Figure 7.3: Comparison of different methods when provided with the same semantic information (synonym) for fine tuning SynGCN embeddings. Results denote the F1-score on SQuAD dataset. SemGCN gives considerable improvement in performance. Please refer Section 7.4.2.3 for details.

Similar to Section 7.4.2.1, the evaluation is performed on the three intrinsic tasks. Due to space limitations, we report results on one representative dataset per task. The results are summarized in Table 7.3. We find that in 13 out of 15 settings, SemGCN outperforms other methods. Overall, we observe that SemGCN, when initialized with SynGCN, gives the best performance on all the tasks (highlighted by  $\square$  in Table 7.3).

For comparing performance on the extrinsic tasks, we first fine-tune SynGCN embeddings using different methods for incorporating semantic information. The embeddings obtained by this process are then evaluated on extrinsic tasks, as in Section 7.4.2.1. The results are shown in Table 7.4. We observe that while the other methods do not always consistently give improvement over the baseline SynGCN, SemGCN is able to improve upon SynGCN in all settings (better or comparable). Overall, we observe that SynGCN along with SemGCN is the most suitable method for incorporating both syntactic and semantic information.

#### 7.4.2.3 Evaluation with Same Semantic Information

In this section, we compare SemGCN against other baselines when provided with the same semantic information: synonyms from PPDB. Similar to Section 7.4.2.2, we compare both on intrinsic and extrinsic tasks with different initializations. The evaluation results of fine-tuned SynGCN embeddings by different methods on SQuAD are shown in the Figure 7.3. The remaining results are included in the supplementary (Table S1 and S2). We observe that compared to other methods, SemGCN is most effective at incorporating semantic constraints across all the initializations and outperforms others at both intrinsic and extrinsic tasks.

Method	POS	SQuAD	NER	Coref
ELMo (E)	$96.1 \pm 0.1$	$81.8 \pm 0.2$	$90.3 \pm 0.3$	$67.8 \pm 0.1$
E+SemGCN(SynGCN, 4)	<b><math>96.2 \pm 0.1</math></b>	<b><math>82.4 \pm 0.1</math></b>	<b><math>90.9 \pm 0.1</math></b>	<b><math>68.3 \pm 0.1</math></b>

Table 7.5: Comparison of ELMo with SynGCN and SemGCN embeddings on multiple extrinsic tasks. For each task, models use a linear combination of the provided embeddings whose weights are learned. Results show that our proposed methods encode complementary information which is not captured by ELMo. Please refer Section 7.4.2.4 for more details.

#### 7.4.2.4 Comparison with ELMo

Recently, ELMo [159] has been proposed which fine-tunes word embedding based on sentential context. In this section, we evaluate SynGCN and SemGCN when given along with pre-trained ELMo embeddings. The results are reported in Table 7.5. The results show that dependency context based embeddings encode complementary information which is not captured by ELMo as it only relies on sequential context. Hence, our proposed methods serves as an effective combination with ELMo.

## 7.5 Conclusion

In this chapter, we have proposed SynGCN, a graph convolution based approach which utilizes syntactic context for learning word representations. SynGCN overcomes the problem of vocabulary explosion and outperforms state-of-the-art word embedding approaches on several intrinsic and extrinsic tasks. We also propose SemGCN, a framework for jointly incorporating diverse semantic information in pre-trained word embeddings. The combination of SynGCN and SemGCN gives the best overall performance. We make the source code of both models available to encourage reproducible research.

## **Part III**

# **Addressing Limitations in Existing GCN Architectures**

# Chapter 8

## Improving Semi-Supervised Learning through Confidence-based Graph Convolutional Networks

### 8.1 Introduction

Graphs are all around us, ranging from citation and social networks to knowledge graphs. Predicting properties of nodes in such graphs is often desirable. For example, given a citation network, we may want to predict the research area of an author. Making such predictions, especially in the semi-supervised setting, has been the focus of graph-based semi-supervised learning (SSL) [184]. In graph-based SSL, a small set of nodes are initially labeled. Starting with such supervision and while utilizing the rest of the graph structure, the initially unlabeled nodes are labeled. Conventionally, the graph structure has been incorporated as an explicit regularizer which enforces a smoothness constraint on the labels estimated on nodes [235, 17, 212]. Recently proposed Graph Convolutional Networks (GCN) [47, 90] provide a framework to apply deep neural networks to graph-structured data. GCNs have been employed successfully for improving performance on tasks such as semantic role labeling [126], machine translation [15], relation extraction [200, 233], document dating [198], shape segmentation [226], and action recognition [79]. GCN formulations for graph-based SSL have also attained state-of-the-art performance [90, 109, 202]. In this chapter, we also focus on the task of graph-based SSL using GCNs.

GCN iteratively estimates embedding of nodes in the graph by aggregating embeddings of neighborhood nodes, while backpropagating errors from a target loss function. Finally, the learned node embeddings are used to estimate label scores on the nodes. In addition to the label scores, it is desirable to also have confidence estimates associated with them. Such confidence

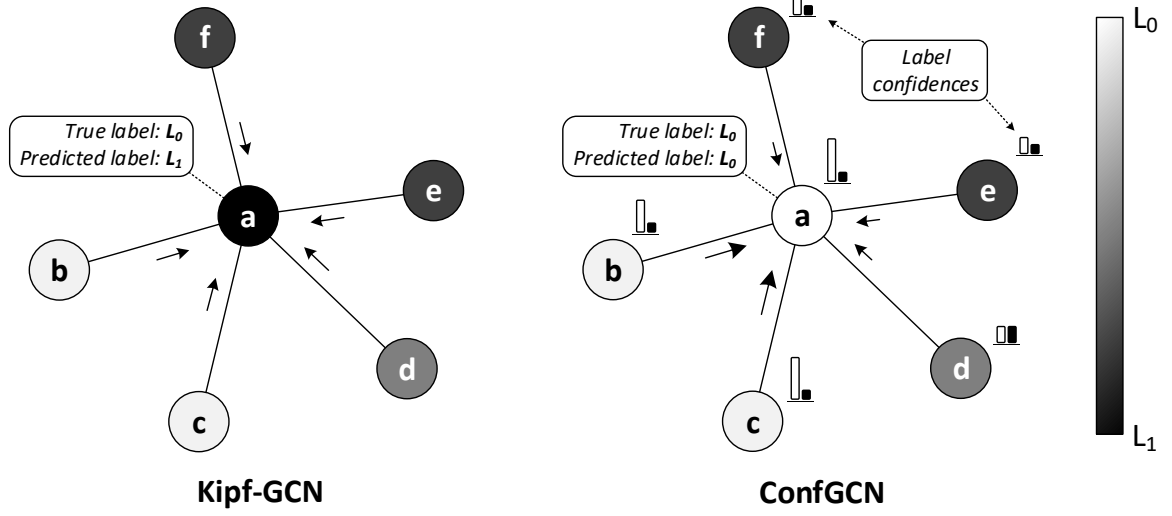


Figure 8.1: Label prediction on node  $a$  by Kipf-GCN and ConfGCN (this chapter).  $L_0$  is  $a$ 's true label. Shade intensity of a node reflects the estimated score of label  $L_1$  assigned to that node. Since Kipf-GCN is not capable of estimating influence of one node on another, it is misled by the dominant label  $L_1$  in node  $a$ 's neighborhood and thereby making the wrong assignment. ConfGCN, on the other hand, estimates confidences (shown by bars) over the label scores, and uses them to increase influence of nodes  $b$  and  $c$  to estimate the right label on  $a$ . Please see Section 8.1 for details.

scores may be used to determine how much to trust the label scores estimated on a given node. While methods to estimate label score confidence in non-deep graph-based SSL has been previously proposed [151], confidence-based GCN is still unexplored.

In order to fill this important gap, we propose ConfGCN, a GCN framework for graph-based SSL. ConfGCN jointly estimates label scores on nodes, along with confidences over them. One of the added benefits of confidence over node's label scores is that they may be used to subdue irrelevant nodes in a node's neighborhood, thereby controlling the number of effective neighbors for each node. In other words, this enables *anisotropic* behavior in GCNs. Let us explain this through the example shown in Figure 8.1. In this figure, while node  $a$  has true label  $L_0$  (white), it is incorrectly classified as  $L_1$  (black) by Kipf-GCN [90]<sup>1</sup>. This is because Kipf-GCN suffers from limitations of its neighborhood aggregation scheme [216]. For example, Kipf-GCN has no constraints on the number of nodes that can influence the representation of a given target node. In a  $k$ -layer Kipf-GCN model, each node is influenced by all the nodes in its  $k$ -hop neighborhood. However, in real world graphs, nodes are often present in *heterogeneous* neighborhoods, i.e., a node is often surrounded by nodes of other labels. For example, in Figure 8.1, node  $a$  is

<sup>1</sup>In this chapter, unless otherwise stated, we refer to Kipf-GCN whenever we mention GCN.

surrounded by three nodes ( $d$ ,  $e$ , and  $f$ ) which are predominantly labeled  $L_1$ , while two nodes ( $b$  and  $c$ ) are labeled  $L_0$ . Please note that all of these are estimated label scores during GCN learning. In this case, it is desirable that node  $a$  is more influenced by nodes  $b$  and  $c$  than the other three nodes. However, since Kipf-GCN doesn't discriminate among the neighboring nodes, it is swayed by the majority and thereby estimating the wrong label  $L_1$  for node  $a$ .

ConfGCN is able to overcome this problem by estimating confidences on each node's label scores. In Figure 8.1, such estimated confidences are shown by bars, with white and black bars denoting confidences in scores of labels  $L_0$  and  $L_1$ , respectively. ConfGCN uses these label confidences to subdue nodes  $d, e, f$  since they have low confidence for their label  $L_1$  (shorter black bars), whereas nodes  $b$  and  $c$  are highly confident about their labels being  $L_0$  (taller white bars). This leads to higher influence of  $b$  and  $c$  during aggregation, and thereby ConfGCN correctly predicting the true label of node  $a$  as  $L_0$  with high confidence. This clearly demonstrates the benefit of label confidences and their utility in estimating node influences. Graph Attention Networks (GAT) [202], a recently proposed method also provides a mechanism to estimate influences by allowing nodes to attend to their neighborhood. However, as we shall see in Section 8.5, ConfGCN, through its use of label confidences, is considerably more effective.

Our contributions in this chapter are as follows.

- We propose ConfGCN, a Graph Convolutional Network (GCN) framework for semi-supervised learning which models label distribution and their confidences for each node in the graph. To the best of our knowledge, this is the first confidence-enabled formulation of GCNs.
- ConfGCN utilize label confidences to estimate influence of one node on another in a label-specific manner during neighborhood aggregation of GCN learning.
- Through extensive evaluation on multiple real-world datasets, we demonstrate ConfGCN effectiveness over state-of-the-art baselines.

ConfGCN's source code and datasets used in the chapter are available at <http://github.com/mallabiisc/ConfGCN>.

## 8.2 Related Work

**Semi-Supervised learning (SSL) on graphs:** SSL on graphs is the problem of classifying nodes in a graph, where labels are available only for a small fraction of nodes. Conventionally, the graph structure is imposed by adding an explicit graph-based regularization term in the loss function [235, 212, 17]. Recently, implicit graph regularization via learned node representation

has proven to be more effective. This can be done either sequentially or in an end to end fashion. Methods like DeepWalk [158], node2vec [70], and LINE [191] first learn graph representations via sampled random walk on the graph or breadth first search traversal and then use the learned representation for node classification. On the contrary, Planetoid [222] learns node embedding by jointly predicting the class labels and the neighborhood context in the graph. Recently, Kipf and Welling [90] employs Graph Convolutional Networks (GCNs) to learn node representations.

**Graph Convolutional Networks (GCNs):** The generalization of Convolutional Neural Networks to non-euclidean domains is proposed by Bruna et al. [28] which formulates the spectral and spatial construction of GCNs. This is later improved through an efficient localized filter approximation [47]. Kipf and Welling [90] provide a first-order formulation of GCNs and show its effectiveness for SSL on graphs. Marcheggiani and Titov [126] propose GCNs for directed graphs and provide a mechanism for edge-wise gating to discard noisy edges during aggregation. This is further improved by Veličković et al. [202] which allows nodes to attend to their neighboring nodes, implicitly providing different weights to different nodes. Liao et al. [109] propose Graph Partition Neural Network (GPNN), an extension of GCNs to learn node representations on large graphs. GPNN first partitions the graph into subgraphs and then alternates between locally and globally propagating information across subgraphs. Recently, Lovasz Convolutional Networks Yadav et al. [218] is proposed for incorporating global graph properties in GCNs. An extensive survey of GCNs and their applications can be found in Bronstein et al. [27].

**Confidence Based Methods:** The natural idea of incorporating confidence in predictions has been explored by Li and Sethi [105] for the task of active learning. Lei [102] proposes a confidence based framework for classification problems, where the classifier consists of two regions in the predictor space, one for confident classifications and other for ambiguous ones. In representation learning, uncertainty (inverse of confidence) is first utilized for word embeddings by Vilnis and McCallum [205]. Athiwaratkun and Wilson [5] further extend this idea to learn hierarchical word representation through encapsulation of probability distributions. Orbach and Crammer [151] propose TACO (Transduction Algorithm with COnfidence), the first graph based method which learns label distribution along with its uncertainty for semi-supervised node classification. Bojchevski and Gnnemann [20] embeds graph nodes as Gaussian distribution using ranking based framework which allows to capture uncertainty of representation. They update node embeddings to maintain neighborhood ordering, i.e. 1-hop neighbors are more similar to 2-hop neighbors and so on. Gaussian embeddings have been used for collaborative filtering [53] and topic modelling [44] as well.

## 8.3 Notation & Problem Statement

Let  $\mathcal{G} = (\mathcal{V}, \mathcal{E}, \mathcal{X})$  be an undirected graph, where  $\mathcal{V} = \mathcal{V}_l \cup \mathcal{V}_u$  is the union of labeled ( $\mathcal{V}_l$ ) and unlabeled ( $\mathcal{V}_u$ ) nodes in the graph with cardinalities  $n_l$  and  $n_u$ ,  $\mathcal{E}$  is the set of edges and  $\mathcal{X} \in \mathbb{R}^{(n_l+n_u) \times d}$  is the input node features. The actual label of a node  $v$  is denoted by a one-hot vector  $Y_v \in \mathbb{R}^m$ , where  $m$  is the number of classes. Given  $\mathcal{G}$  and seed labels  $Y \in \mathbb{R}^{n_l \times m}$ , the goal is to predict the labels of the unlabeled nodes. To incorporate confidence, we additionally estimate label distribution  $\boldsymbol{\mu}_v \in \mathbb{R}^m$  and a diagonal co-variance matrix  $\boldsymbol{\Sigma}_v \in \mathbb{R}^{m \times m}$ ,  $\forall v \in \mathcal{V}$ . Here,  $\boldsymbol{\mu}_{v,i}$  denotes the score of label  $i$  on node  $v$ , while  $(\boldsymbol{\Sigma}_v)_{ii}$  denotes the variance in the estimation of  $\boldsymbol{\mu}_{v,i}$ . In other words,  $(\boldsymbol{\Sigma}_v^{-1})_{ii}$  is ConfGCN's confidence in  $\boldsymbol{\mu}_{v,i}$ .

## 8.4 Proposed Method: Confidence Based Graph Convolutional Networks (ConfGCN)

Following [151], ConfGCN uses co-variance matrix based symmetric Mahalanobis distance for defining distance between two nodes in the graph. Formally, for any two given nodes  $u$  and  $v$ , with label distributions  $\boldsymbol{\mu}_u$  and  $\boldsymbol{\mu}_v$  and co-variance matrices  $\boldsymbol{\Sigma}_u$  and  $\boldsymbol{\Sigma}_v$ , distance between them is defined as follows.

$$d_M(u, v) = (\boldsymbol{\mu}_u - \boldsymbol{\mu}_v)^T (\boldsymbol{\Sigma}_u^{-1} + \boldsymbol{\Sigma}_v^{-1}) (\boldsymbol{\mu}_u - \boldsymbol{\mu}_v).$$

Characteristic of the above distance metric is that if either of  $\boldsymbol{\Sigma}_u$  or  $\boldsymbol{\Sigma}_v$  has large eigenvalues, then the distance will be low irrespective of the closeness of  $\boldsymbol{\mu}_u$  and  $\boldsymbol{\mu}_v$ . On the other hand, if  $\boldsymbol{\Sigma}_u$  and  $\boldsymbol{\Sigma}_v$  both have low eigenvalues, then it requires  $\boldsymbol{\mu}_u$  and  $\boldsymbol{\mu}_v$  to be close for their distance to be low. Given the above properties, we define  $r_{uv}$ , the influence score of node  $u$  on its neighboring node  $v$  during GCN aggregation, as follows.

$$r_{uv} = \frac{1}{d_M(u, v)}.$$

This influence score gives more relevance to neighboring nodes with highly confident similar label, while reducing importance of nodes with low confident label scores. This results in ConfGCN acquiring anisotropic capability during neighborhood aggregation. For a node  $v$ , ConfGCN's equation for updating embedding at the  $k$ -th layer is thus defined as follows.

$$\mathbf{h}_v^{k+1} = f \left( \sum_{u \in \mathcal{N}(v)} r_{uv} \times (\mathbf{W}^k \mathbf{h}_u^k + \mathbf{b}^k) \right), \forall v \in \mathcal{V}. \quad (8.1)$$

Dataset	Nodes	Edges	Classes	Features	Label Mismatch	$\frac{ \mathcal{V}_l }{ \mathcal{V} }$
Cora	2,708	5,429	7	1,433	0.002	0.052
Cora-ML	2,995	8,416	7	2,879	0.018	0.166
Citeseer	3,327	4,372	6	3,703	0.003	0.036
Pubmed	19,717	44,338	3	500	0.0	0.003

Table 8.1: Details of the datasets used in the chapter. Please refer Section 8.5.1.1 for more details.

The final node representation obtained from ConfGCN is used for predicting labels of the nodes in the graph as follows.

$$\hat{\mathbf{Y}}_v = \text{softmax}(\mathbf{W}^K \mathbf{h}_v^K + \mathbf{b}^K), \quad \forall v \in \mathcal{V}$$

where,  $K$  denotes the number of ConfGCN’s layers. Finally, in order to learn label scores  $\{\boldsymbol{\mu}_v\}$  and co-variance matrices  $\{\boldsymbol{\Sigma}_v\}$  jointly with other parameters  $\{\mathbf{W}^k, \mathbf{b}^k\}$ , following Orbach and Crammer [151], we include the following two terms in ConfGCN’s objective function.

For enforcing neighboring nodes to be close to each other, we include  $L_{\text{smooth}}$  defined as

$$L_{\text{smooth}} = \sum_{(u,v) \in \mathcal{E}} (\boldsymbol{\mu}_u - \boldsymbol{\mu}_v)^T (\boldsymbol{\Sigma}_u^{-1} + \boldsymbol{\Sigma}_v^{-1}) (\boldsymbol{\mu}_u - \boldsymbol{\mu}_v).$$

To impose the desirable property that the label distribution of nodes in  $\mathcal{V}_l$  should be close to their input label distribution, we incorporate  $L_{\text{label}}$  defined as

$$L_{\text{label}} = \sum_{v \in \mathcal{V}_l} (\boldsymbol{\mu}_v - \mathbf{Y}_v)^T (\boldsymbol{\Sigma}_v^{-1} + \frac{1}{\gamma} \mathbf{I}) (\boldsymbol{\mu}_v - \mathbf{Y}_v).$$

Here, for input labels, we assume a fixed uncertainty of  $\frac{1}{\gamma} \mathbf{I} \in \mathbb{R}^{L \times L}$ , where  $\gamma > 0$ . We also include the following regularization term,  $L_{\text{reg}}$  to constraint the co-variance matrix to be finite and positive.

$$L_{\text{reg}} = \sum_{v \in \mathcal{V}} \text{Tr} \max(-\boldsymbol{\Sigma}_v, 0),$$

This regularization term enforces soft positivity constraint on co-variance matrix. Additionally in ConfGCN, we include the  $L_{\text{const}}$  in the objective, to push the label distribution ( $\boldsymbol{\mu}$ ) close to the final model prediction ( $\hat{\mathbf{Y}}$ ).

$$L_{\text{const}} = \sum_{v \in \mathcal{V}} (\boldsymbol{\mu}_v - \hat{\mathbf{Y}}_v)^T (\boldsymbol{\mu}_v - \hat{\mathbf{Y}}_v).$$

Finally, we include the standard cross-entropy loss for semi-supervised multi-class classification over all the labeled nodes ( $\mathcal{V}_l$ ).

$$L_{\text{cross}} = - \sum_{v \in \mathcal{V}_l} \sum_{j=1}^m \mathbf{Y}_{vj} \log(\hat{\mathbf{Y}}_{vj}).$$

The final objective for optimization is the linear combination of the above defined terms.

$$\begin{aligned} L(\theta) = & - \sum_{i \in \mathcal{V}_l} \sum_{j=1}^L \mathbf{Y}_{ij} \log(\hat{\mathbf{Y}}_{ij}) \\ & + \lambda_1 \sum_{(u,v) \in \mathcal{E}} (\boldsymbol{\mu}_u - \boldsymbol{\mu}_v)^T (\boldsymbol{\Sigma}_u^{-1} + \boldsymbol{\Sigma}_v^{-1})(\boldsymbol{\mu}_u - \boldsymbol{\mu}_v) \\ & + \lambda_2 \sum_{u \in \mathcal{V}_l} (\boldsymbol{\mu}_u - \mathbf{Y}_u)^T (\boldsymbol{\Sigma}_u^{-1} + \frac{1}{\gamma} \mathbf{I})(\boldsymbol{\mu}_u - \mathbf{Y}_u) \\ & + \lambda_3 \sum_{v \in \mathcal{V}} (\boldsymbol{\mu}_v - \hat{\mathbf{Y}}_v)^T (\boldsymbol{\mu}_v - \hat{\mathbf{Y}}_v) \\ & + \lambda_4 \sum_{v \in \mathcal{V}} \text{Tr} \max(-\boldsymbol{\Sigma}_v, 0) \end{aligned}$$

where,  $\theta = \{\mathbf{W}^k, \mathbf{b}^k, \boldsymbol{\mu}_v, \boldsymbol{\Sigma}_v\}$  and  $\lambda_i \in \mathbb{R}$ , are the weights of the terms in the objective. We optimize  $L(\theta)$  using stochastic gradient descent. We hypothesize that all the terms help in improving ConfGCN's performance and we validate this in Section 8.5.2.4.

## 8.5 Experiments

### 8.5.1 Experimental Setup

#### 8.5.1.1 Datasets

For evaluating the effectiveness of ConfGCN, we evaluate on several semi-supervised classification benchmarks. Following the experimental setup of [90, 109], we evaluate on Cora, Citeseer, and Pubmed datasets [172]. The dataset statistics is summarized in Table 8.1. Label mismatch denotes the fraction of edges between nodes with different labels in the training data. The benchmark datasets commonly used for semi-supervised classification task have substantially low label mismatch rate. In order to examine models on datasets with more heterogeneous neighborhoods, we also evaluate on Cora-ML dataset [20].

Method	Citeseer	Cora	Pubmed	Cora ML
LP [235]	45.3	68.0	63.0	-
ManiReg [17]	60.1	59.5	70.7	-
SemiEmb [212]	59.6	59.0	71.1	-
Feat [222]	57.2	57.4	69.8	-
DeepWalk [158]	43.2	67.2	65.3	-
GGNN [107]	68.1	77.9	77.2	-
Planetoid [222]	64.9	75.7	75.7	-
Kipf-GCN [90]	$69.4 \pm 0.4$	$80.9 \pm 0.4$	$76.8 \pm 0.2$	$85.7 \pm 0.3$
G-GCN [126]	$69.6 \pm 0.5$	$81.2 \pm 0.4$	$77.0 \pm 0.3$	$86.0 \pm 0.2$
GPNN [109]	$68.1 \pm 1.8$	$79.0 \pm 1.7$	$73.6 \pm 0.5$	$69.4 \pm 2.3$
GAT [202]	$72.5 \pm 0.7$	$83.0 \pm 0.7$	$79.0 \pm 0.3$	$83.0 \pm 0.8$
ConfGCN (this work)	<b><math>72.7 \pm 0.8</math></b>	$82.0 \pm 0.3$	<b><math>79.5 \pm 0.5</math></b>	<b><math>86.5 \pm 0.3</math></b>

Table 8.2: Performance comparison of several methods for semi-supervised node classification on multiple benchmark datasets. ConfGCN performs consistently better across all the datasets. Baseline method performances on Citeseer, Cora and Pubmed datasets are taken from Liao et al. [109], Veličković et al. [202]. We consider only the top performing baseline methods on these datasets for evaluation on the Cora-ML dataset. Please refer Section 8.5.2.1 for details.

All the four datasets are citation networks, where each document is represented using bag-of-words features in the graph with undirected citation links between documents. The goal is to classify documents into one of the predefined classes. We use the data splits used by [222] and follow similar setup for Cora-ML dataset. Following [90], additional 500 labeled nodes are used for hyperparameter tuning.

**Hyperparameters:** We use the same data splits as described in [222], with a test set of 1000 labeled nodes for testing the prediction accuracy of ConfGCN and a validation set of 500 labeled nodes for optimizing the hyperparameters. The ranges of hyperparameters were adapted from previous literature [151, 90]. The model is trained using Adam [89] with a learning rate of 0.01. The weight matrices along with  $\mu$  are initialized using Xavier initialization [67] and  $\Sigma$  matrix is initialized with identity. To avoid numerical instability we model  $\Sigma^{-1}$  directly and compute  $\Sigma$  wherever required. Following Kipf and Welling [90], we use two layers of GCN ( $K$ ) for all the experiments in this chapter.

### 8.5.1.2 Baselines

For evaluating ConfGCN, we compare against the following baselines:

- **Feat** [222] takes only node features as input and ignores the graph structure.

- **ManiReg** [17] is a framework for providing data-dependent geometric regularization.
- **SemiEmb** [212] augments deep architectures with semi-supervised regularizers to improve their training.
- **LP** [235] is an iterative iterative label propagation algorithm which propagates a node's labels to its neighboring unlabeled nodes according to their proximity.
- **DeepWalk** [158] learns node features by treating random walks in a graph as the equivalent of sentences.
- **Planetoid** [222] provides a transductive and inductive framework for jointly predicting class label and neighborhood context of a node in the graph.
- **GCN** [90] is a variant of convolutional neural networks used for semi-supervised learning on graph-structured data.
- **G-GCN** [126] is a variant of GCN with edge-wise gating to discard noisy edges during aggregation.
- **GGNN** [107] is a generalization of RNN framework which can be used for graph-structured data.
- **GPNN** [109] is a graph partition based algorithm which propagates information after partitioning large graphs into smaller subgraphs.
- **GAT** [202] is a graph attention based method which provides different weights to different nodes by allowing nodes to attend to their neighborhood.

### 8.5.2 Results

In this section, we attempt to answer the following questions:

- Q1. How does ConfGCN compare against existing methods for the semi-supervised node classification task? (Section 8.5.2.1)
- Q2. How do the performance of methods vary with increasing node degree and neighborhood label mismatch? (Section 8.5.2.2)
- Q3. How does increasing the number of layers effect ConfGCN's performance? (Section 8.5.2.3)
- Q4. What is the effect of ablating different terms in ConfGCN's loss function? (Section 8.5.2.4)

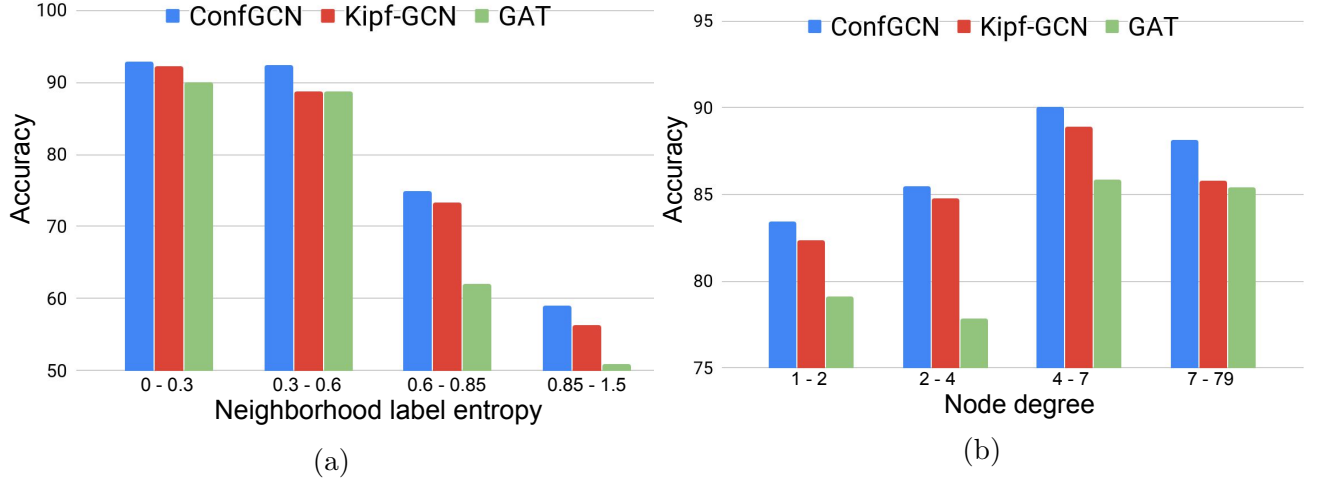


Figure 8.2: Plots of node classification accuracy vs. (a) neighborhood label entropy and (b) node degree. On  $x$ -axis, we plot quartiles of (a) neighborhood label entropy and (b) degree, i.e., each bin has 25% of the samples in sorted order. Overall, we observe that ConfGCN performs better than Kipf-GCN and GAT at all levels of node entropy and degree. Please see Section 8.5.2.2 for details.

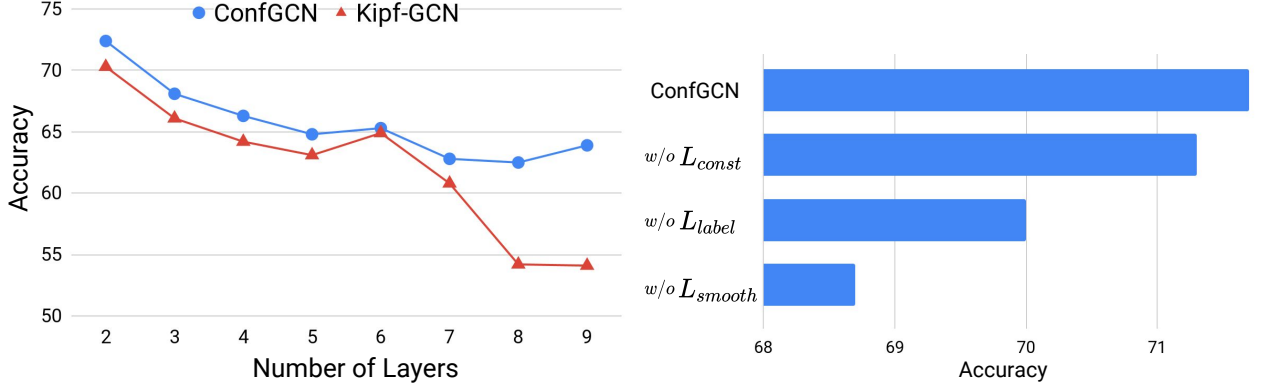
### 8.5.2.1 Node Classification

The evaluation results for semi-supervised node classification are summarized in Table 8.2. Results of all other baseline methods on Cora, Citeseer and Pubmed datasets are taken from [109, 202] directly. For evaluation on the Cora-ML dataset, only top performing baselines from the other three datasets are considered. Overall, we find that ConfGCN outperforms all existing approaches consistently across all the datasets.

This may be attributed to ConfGCN’s ability to model nodes’ label distribution along with the confidence scores which subdues the effect of noisy nodes during neighborhood aggregation. The lower performance of GAT [202] compared to Kipf-GCN on Cora-ML shows that computing attention based on the hidden representation of nodes is not much helpful in suppressing noisy neighborhood nodes. We also observe that the performance of GPNN [109] suffers on the Cora-ML dataset. This is due to the fact that while propagating information between small subgraphs, the high label mismatch rate in Cora-ML (please see Table 8.1) leads to wrong information propagation. Hence, during the global propagation step, this error is further magnified.

### 8.5.2.2 Effect of Node Entropy and Degree on Performance

In this section, we provide an analysis of the performance of Kipf-GCN, GAT and ConfGCN for node classification on the Cora-ML dataset which has higher label mismatch rate. We use



(a) Evaluation of Kipf-GCN and ConfGCN on the citeseer dataset with increasing number of layers. Overall, ConfGCN outperforms Kipf-GCN, and while both methods' performance degrades with increasing layers, ConfGCN's degradation is more gradual than Kipf-GCN's abrupt drop. Please see Section 8.5.2.3 for details.

(b) Performance comparison of different ablated version of ConfGCN on the citeseer dataset. These results justify inclusion of the different terms in ConfGCN's objective function. Please see Section 8.5.2.4 for details.

neighborhood label entropy to quantify label mismatch, which for a node  $u$  is defined as follows.

$$\text{NeighborLabelEntropy}(u) = - \sum_{l=1}^L p_{ul} \log p_{ul}$$

$$\text{where, } p_{ul} = \frac{|\{v \in \mathcal{N}(u) \mid \text{label}(v) = l\}|}{|\mathcal{N}(u)|}.$$

Here,  $\text{label}(v)$  is the true label of node  $v$ . The results for neighborhood label entropy and node degree are summarized in Figures 8.2a and 8.2b, respectively. On the x-axis of these figures, we plot quartiles of label entropy and degree, i.e., each bin has 25% of the instances in sorted order. With increasing neighborhood label entropy, the node classification task is expected to become more challenging. We indeed see this trend in Figure 8.2a where performances of all the methods degrade with increasing neighborhood label entropy. However, ConfGCN performs comparatively better than the existing state-of-art approaches at all levels of node entropy.

In case of node degree also (Figure 8.2b), we find that ConfGCN performs better than Kipf-GCN and GAT at all quartiles of node degrees. Classifying sparsely connected nodes (first and second bins) is challenging as very little information is present in the node neighborhood. Performance improves with availability of moderate number of neighboring nodes (third bin), but further increase in degree (fourth bin) results in introduction of many potentially noisy

neighbors, thereby affecting performance of all the methods. For higher degree nodes, ConfGCN gives an improvement of around 3% over GAT and Kipf-GCN. This shows that ConfGCN, through its use of label confidences, is able to give higher influence score to relevant nodes in the neighborhood during aggregation while reducing importance of the noisy ones.

#### 8.5.2.3 Effect of Increasing Convolutional Layers

Recently, Xu et al. [216] highlighted an unusual behavior of Kipf-GCN where its performance degrades significantly with increasing number of layers. This is because of increase in the number of influencing nodes with increasing layers, resulting in “averaging out” of information during aggregation. For comparison, we evaluate the performance of Kipf-GCN and ConfGCN on citeseer dataset with increasing number of convolutional layers. The results are summarized in Figure 8.3a. We observe that Kipf-GCN’s performance degrades drastically with increasing number of layers, whereas ConfGCN’s decrease in performance is more gradual. This shows that confidence based GCN helps in alleviating this problem. We also note that ConfGCN outperforms Kipf-GCN at all layer levels.

#### 8.5.2.4 Ablation Results

In this section, we evaluate the different ablated version of ConfGCN by cumulatively eliminating terms from its objective function as defined in Section 8.4. The results on citeseer dataset are summarized in Figure 8.3b. Overall, we find that each term ConfGCN’s loss function (Equation 8.4) helps in improving its performance and the method performs best when all the terms are included.

## 8.6 Conclusion

In this chapter, we present ConfGCN, a confidence based Graph Convolutional Network which estimates label scores along with their confidences jointly in a GCN-based setting. In ConfGCN, the influence of one node on another during aggregation is determined using the estimated confidences and label scores, thus inducing anisotropic behavior to GCN. We demonstrate the effectiveness of ConfGCN against state-of-the-art methods for the semi-supervised node classification task and analyze its performance in different settings. We make ConfGCN’s source code available.

# Chapter 9

## Composition-based Multi-Relational Graph Convolutional Networks for Relational Graphs

### 9.1 Introduction

In this chapter, we address another important limitation of Graph Convolutional models. Most of the existing research on GCNs [90, 72, 202] have focused on learning representations of nodes in simple undirected graphs. A more general and pervasive class of graphs are multi-relational graphs<sup>1</sup>. A notable example of such graphs is knowledge graphs. Most of the existing GCN based approaches for handling relational graphs [126, 171] suffer from over-parameterization and are limited to learning only node representations. Hence, such methods are not directly applicable for tasks that require relation embedding vectors such as link prediction. Initial attempts at learning representations for relations in graphs [138, 16] have shown some performance gains on tasks like node classification and neural machine translation.

There has been extensive research on embedding Knowledge Graphs (KG) [145, 208] where representations of both nodes and relations are jointly learned. These methods are restricted to learning embeddings using link prediction objective. Even though GCNs can learn from task-specific objectives such as classification, their application has been largely restricted to non-relational graph setting. Thus, there is a need for a framework which can utilize KG embedding techniques for learning task-specific node and relation embeddings. In this chapter, we propose COMPGCN, a novel GCN framework for multi-relational graphs which systematically

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<sup>1</sup>In this chapter, multi-relational graphs refer to graphs with edges that have labels and directions.

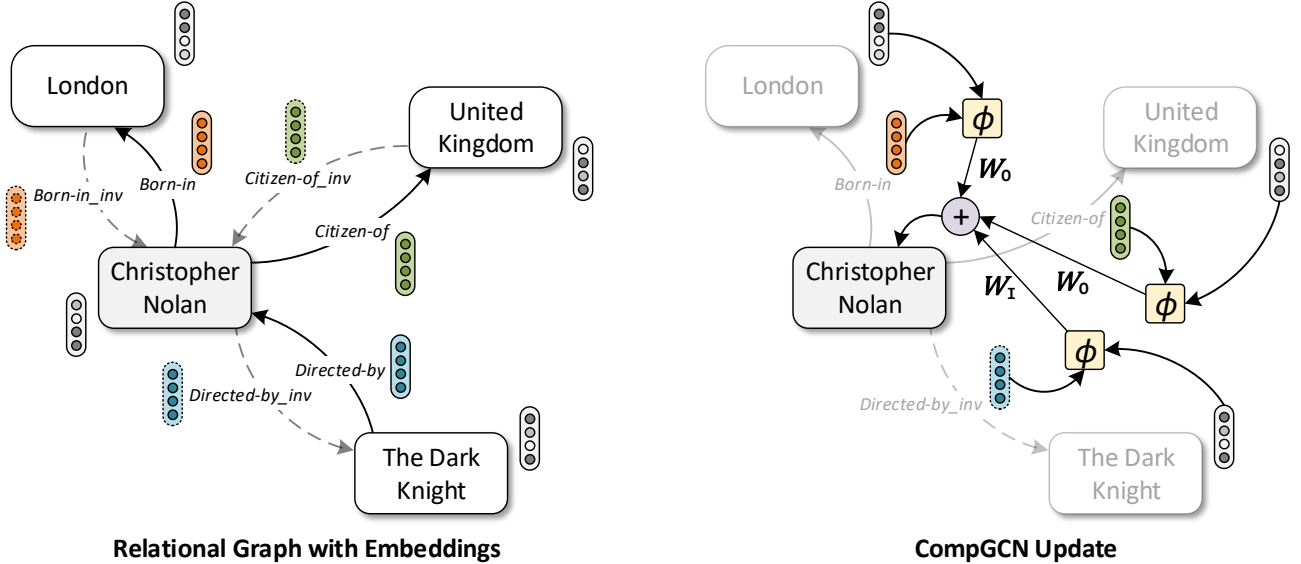


Figure 9.1: Overview of COMPGCN. Given node and relation embeddings, COMPGCN performs a composition operation  $\phi(\cdot)$  over each edge in the neighborhood of a central node (e.g. *Christopher Nolan* above). The composed embeddings are then convolved with specific filters  $\mathbf{W}_O$  and  $\mathbf{W}_I$  for original and inverse relations respectively. We omit self-loop in the diagram for clarity. The message from all the neighbors are then aggregated to get an updated embedding of the central node. Also, the relation embeddings are transformed using a separate weight matrix. Please refer to Section 9.3 for details.

leverages entity-relation composition operations from knowledge graph embedding techniques. COMPGCN addresses the shortcomings of previously proposed GCN models by jointly learning vector representations for both nodes and relations in the graph. An overview of COMPGCN is presented in Figure 9.1. The contributions of our work can be summarized as follows:

1. We propose COMPGCN, a novel framework for incorporating multi-relational information in Graph Convolutional Networks which leverages a variety of composition operations from knowledge graph embedding techniques to jointly embed both nodes and relations in a graph.
2. We demonstrate that COMPGCN framework generalizes several existing multi-relational GCN methods (Proposition 9.1) and also scales with the increase in number of relations in the graph (Section 9.4.2.3).
3. Through extensive experiments on tasks such as node classification, link prediction, and graph classification, we demonstrate the effectiveness of our proposed method.

## 9.2 Related Work

**Graph Convolutional Networks:** GCNs generalize Convolutional Neural Networks (CNNs) to non-Euclidean data. GCNs were first introduced by [28] and later made scalable through efficient localized filters in the spectral domain [47]. A first-order approximation of GCNs using Chebyshev polynomials has been proposed by [90]. Recently, several of its extensions have also been formulated [72, 202, 217]. Most of the existing GCN methods follow *Message Passing Neural Networks* (MPNN) framework [66] for node aggregation. Our proposed method can be seen as an instantiation of the MPNN framework. However, it is specialized for relational graphs.

**GCNs for Multi-Relational Graph:** An extension of GCNs for relational graphs is proposed by [126]. However, they only consider direction-specific filters and ignore relations due to over-parameterization. [171] address this shortcoming by proposing basis and block-diagonal decomposition of relation specific filters. *Weighted Graph Convolutional Network* [173] utilizes learnable relational specific scalar weights during GCN aggregation. While these methods show performance gains on node classification and link prediction, they are limited to embedding only the nodes of the graph. Contemporary to our work, [225] have also proposed an extension of GCNs for embedding both nodes and relations in multi-relational graphs. However, our proposed method is a more generic framework which can leverage any KG composition operator. We compare against their method in Section 9.4.2.1.

**Knowledge Graph Embedding:** Knowledge graph (KG) embedding is a widely studied field [145, 208] with application in tasks like link prediction and question answering [23]. Most of KG embedding approaches define a score function and train node and relation embeddings such that valid triples are assigned a higher score than the invalid ones. Based on the type of score function, KG embedding method are classified as translational [22, 211], semantic matching based [221, 147] and neural network based [178, 50]. In our work, we evaluate the performance of COMPGCN on link prediction with methods of all three types.

## 9.3 CompGCN Details

In this section, we provide a detailed description of our proposed method, COMPGCN. The overall architecture is shown in Figure 9.1. We represent a multi-relational graph by  $\mathcal{G} = (\mathcal{V}, \mathcal{R}, \mathcal{E}, \mathcal{X}, \mathcal{Z})$  as defined in Section 2.1.5 where  $\mathcal{Z} \in \mathbb{R}^{|\mathcal{R}| \times d_0}$  denotes the initial relation features. Our model is motivated by the first-order approximation of GCNs using Chebyshev polynomials [90]. Following Marcheggiani and Titov [126], we also allow the information in a directed edge to flow along both directions. Hence, we extend  $\mathcal{E}$  and  $\mathcal{R}$  with corresponding

inverse edges and relations, i.e.,

$$\mathcal{E}' = \mathcal{E} \cup \{(v, u, r^{-1}) \mid (u, v, r) \in \mathcal{E}\} \cup \{(u, u, \top) \mid u \in \mathcal{V}\},$$

and  $\mathcal{R}' = \mathcal{R} \cup \mathcal{R}_{inv} \cup \{\top\}$ , where  $\mathcal{R}_{inv} = \{r^{-1} \mid r \in \mathcal{R}\}$  denotes the inverse relations and  $\top$  indicates the self loop.

### 9.3.1 Relation-based Composition

Unlike most of the existing methods which embed only nodes in the graph, COMPGCN learns a  $d$ -dimensional representation  $\mathbf{h}_r \in \mathbb{R}^d, \forall r \in \mathcal{R}$  along with node embeddings  $\mathbf{h}_v \in \mathbb{R}^d, \forall v \in \mathcal{V}$ . Representing relations as vectors alleviates the problem of over-parameterization while applying GCNs on relational graphs. Further, it allows COMPGCN to exploit any available relation features ( $\mathcal{Z}$ ) as initial representations. To incorporate relation embeddings into the GCN formulation, we leverage the entity-relation composition operations used in Knowledge Graph embedding approaches [22, 145], which are of the form

$$\mathbf{e}_o = \phi(\mathbf{e}_s, \mathbf{e}_r).$$

Here,  $\phi : \mathbb{R}^d \times \mathbb{R}^d \rightarrow \mathbb{R}^d$  is a composition operator,  $s$ ,  $r$ , and  $o$  denote subject, relation and object in the knowledge graph and  $\mathbf{e}_{(.)} \in \mathbb{R}^d$  denotes their corresponding embeddings. In this chapter, we restrict ourselves to non-parameterized operations like subtraction [22], multiplication [221] and circular-correlation [147]. However, COMPGCN can be extended to parameterized operations like Neural Tensor Networks (NTN) [178] and ConvE [50]. We defer their analysis as future work.

As we show in Section 9.4.2, the choice of composition operation is important in deciding the quality of the learned embeddings. Hence, superior composition operations for Knowledge Graphs developed in future can be adopted to improve COMPGCN’s performance further.

### 9.3.2 CompGCN Update Equation

The GCN update equation (Eq. 2.5)) defined in Section 2.1.4 can be re-written as

$$\mathbf{h}_v = f \left( \sum_{(u,r) \in \mathcal{N}(v)} \mathbf{W}_r \mathbf{h}_u \right),$$

where  $\mathcal{N}(v)$  is a set of immediate neighbors of  $v$  for its outgoing edges. Since this formulation suffers from over-parameterization, in COMPGCN we perform composition ( $\phi$ ) of a neighboring

node  $u$  with respect to its relation  $r$  as defined above. This allows our model to be relation aware while being linear ( $\mathcal{O}(|\mathcal{R}|d)$ ) in the number of feature dimensions. Moreover, for treating original, inverse, and self edges differently, we define separate filters for each of them. The update equation of COMPGCN is given as:

$$\mathbf{h}_v = f \left( \sum_{(u,r) \in \mathcal{N}(v)} \mathbf{W}_{\lambda(r)} \phi(\mathbf{x}_u, \mathbf{z}_r) \right), \quad (9.1)$$

where  $\mathbf{x}_u, \mathbf{z}_r$  denotes initial features for node  $u$  and relation  $r$  respectively,  $\mathbf{h}_v$  denotes the updated representation of node  $v$ , and  $\mathbf{W}_{\lambda(r)} \in \mathbb{R}^{d_1 \times d_0}$  is a relation-type specific parameter. In COMPGCN, we use direction specific weights, i.e.,  $\lambda(r) = \text{dir}(r)$ , given as:

$$\mathbf{W}_{\text{dir}(r)} = \begin{cases} \mathbf{W}_O, & r \in \mathcal{R} \\ \mathbf{W}_I, & r \in \mathcal{R}_{\text{inv}} \\ \mathbf{W}_S, & r = \top \text{ (self-loop)} \end{cases} \quad (9.2)$$

Further, in COMPGCN, after the node embedding update defined in Eq. 9.1, the relation embeddings are also transformed as follows:

$$\mathbf{h}_r = \mathbf{W}_{\text{rel}} \mathbf{z}_r, \quad (9.3)$$

where  $\mathbf{W}_{\text{rel}} \in \mathbb{R}^{d_1 \times d_0}$  is a learnable transformation matrix which projects the relations to the same embedding space as nodes and allows them to be utilized in the next COMPGCN layer.

To ensure that COMPGCN scales with the increasing number of relations, we use a variant of the basis formulations proposed in Schlichtkrull et al. [171]. Instead of independently defining an embedding for each relation, they are expressed as a linear combination of a set of basis vectors. Formally, let  $\{\mathbf{v}_1, \mathbf{v}_2, \dots, \mathbf{v}_{\mathcal{B}}\}$  be a set of learnable basis vectors. Then, initial relation representation is given as:

$$\mathbf{z}_r = \sum_{b=1}^{\mathcal{B}} \alpha_{br} \mathbf{v}_b.$$

Here,  $\alpha_{br} \in \mathbb{R}$  is relation and basis specific learnable scalar weight. Note that this is different from the formulation in Schlichtkrull et al. [171], where a separate set of basis matrices is defined for each GCN layer. In COMPGCN, basis vectors are defined only for the first layer, and the later layers share the relations through transformations according to Equation 9.3.

We can extend the formulation of Equation 9.1 to the case where we have  $k$ -stacked

Methods	$\mathbf{W}_{\lambda(r)}^k$	$\phi(\mathbf{h}_u^k, \mathbf{h}_r^k)$
Kipf-GCN [90]	$\mathbf{W}^k$	$\mathbf{h}_u^k$
Relational-GCN [171]	$\mathbf{W}_r^k$	$\mathbf{h}_u^k$
Directed-GCN [126]	$\mathbf{W}_{\text{dir}(r)}^k$	$\mathbf{h}_u^k$
Weighted-GCN [173]	$\mathbf{W}^k$	$\alpha_r^k \mathbf{h}_u^k$

Table 9.1: Reduction of COMPGCN to several existing Graph Convolutional methods. Here,  $\alpha_r^k$  is a relation specific scalar,  $\mathbf{W}_r^k$  denotes a separate weight for each relation, and  $\mathbf{W}_{\text{dir}(r)}^k$  is as defined in Equation 9.2. Please refer to Proposition 9.1 for more details.

COMPGCN layers. Let  $\mathbf{h}_v^{k+1}$  denote the representation of a node  $v$  obtained after  $k$  layers which is defined as

$$\mathbf{h}_v^{k+1} = f\left(\sum_{(u,r) \in \mathcal{N}(v)} \mathbf{W}_{\lambda(r)}^k \phi(\mathbf{h}_u^k, \mathbf{h}_r^k)\right). \quad (9.4)$$

Similarly, let  $\mathbf{h}_r^{k+1}$  denote the representation of a relation  $r$  after  $k$  layers. Then,

$$\mathbf{h}_r^{k+1} = \mathbf{W}_{\text{rel}}^k \mathbf{h}_r^k.$$

Here,  $\mathbf{h}_v^0$  and  $\mathbf{h}_r^0$  are the initial node ( $\mathbf{x}_v$ ) and relation ( $\mathbf{z}_r$ ) features respectively.

**Proposition 9.1** COMPGCN generalizes the following Graph Convolutional based methods: **Kipf-GCN** [90], **Relational GCN** [171], **Directed GCN** [126], and **Weighted GCN** [173].

**Proof:** For Kipf-GCN, this can be trivially obtained by making weights ( $\mathbf{W}_{\lambda(r)}$ ) and composition function ( $\phi$ ) relation agnostic in Equation 9.4, i.e.,  $\mathbf{W}_{\lambda(r)} = \mathbf{W}$  and  $\phi(\mathbf{h}_u, \mathbf{h}_r) = \mathbf{h}_u$ . Similar reductions can be obtained for other methods as shown in Table 9.1.  $\square$

## 9.4 Experiments

### 9.4.1 Experimental Setup

#### 9.4.1.1 Evaluation tasks

In our experiments, we evaluate COMPGCN on the below-mentioned tasks.

- **Link Prediction** is the task of inferring missing facts based on the known facts in Knowledge Graphs. In our experiments, we utilize FB15k-237 [193] and WN18RR [50] datasets for evaluation. Following [22], we use filtered setting for evaluation and report Mean Reciprocal Rank (MRR), Mean Rank (MR) and Hits@N.

	Link Prediction		Node Classification		Graph Classification	
	FB15k-237	WN18RR	MUTAG (Node)	AM	MUTAG (Graph)	PTC
Graphs	1	1	1	1	188	344
Entities	14,541	40,943	23,644	1,666,764	17.9 (Avg)	25.5 (Avg)
Edges	310,116	93,003	74,227	5,988,321	39.6 (Avg)	29.5 (Avg)
Relations	237	11	23	133	4	4
Classes	-	-	2	11	2	2

Table 9.2: The details of the datasets used for node classification, link prediction, and graph classification tasks. Please refer to Section 9.4.1.1 for more details.

- **Node Classification** is the task of predicting the labels of nodes in a graph-based on node features and their connections. Similar to [171], we evaluate COMP-GCN on MUTAG (Node) and AM [168] datasets.
- **Graph Classification**, where, given a set of graphs and their corresponding labels, the goal is to learn a representation for each graph which is fed to a classifier for prediction. We evaluate on 2 bioinformatics dataset: MUTAG (Graph) and PTC [220].

A summary statistics of the datasets used is provided in Table 9.2.

#### 9.4.1.2 Baselines

Across all tasks, we compare against the following GCN methods for relational graphs: (1) Relational-GCN (**R-GCN**) [171] which uses relation-specific weight matrices that are defined as a linear combinations of a set of basis matrices. (2) Directed-GCN (**D-GCN**) [126] has separate weight matrices for incoming edges, outgoing edges, and self-loops. It also has relation-specific biases. (3) Weighted-GCN (**W-GCN**) [173] assigns a learnable scalar weight to each relation and multiplies an incoming "message" by this weight. Apart from this, we also compare with several task-specific baselines mentioned below.

**Link prediction:** For evaluating COMP-GCN, we compare against several non-neural and neural baselines: TransE [22], DistMult [221], ComplEx [195], R-GCN [171], KBGAN [30], ConvE [50], ConvKB [143], SACN [173], HypER [9], RotatE [186], ConvR [83], and VR-GCN [225].

**Node and Graph Classification:** For node classification, following [171], we compare with Feat [154], WL [174], and RDF2Vec [168]. Finally, for graph classification, we evaluate against PACHYSAN [148], Deep Graph CNN (DGCNN) [232], and Graph Isomorphism Network (GIN) [217].

	FB15k-237					WN18RR				
	MRR	MR	H@10	H@3	H@1	MRR	MR	H@10	H@3	H@1
TransE [22]	.294	357	.465	-	-	.226	3384	.501	-	-
DistMult [221]	.241	254	.419	.263	.155	.43	5110	.49	.44	.39
ComplEx [195]	.247	339	.428	.275	.158	.44	5261	.51	.46	.41
R-GCN [171]	.248	-	.417	-	.151	-	-	-	-	-
KBGAN [30]	.278	-	.458	-	-	.214	-	.472	-	-
ConvE [50]	.325	244	.501	.356	.237	.43	4187	.52	.44	.40
ConvKB [143]	.243	311	.421	.371	.155	.249	3324	.524	.417	.057
SACN [173]	.35	-	.54	.39	.26	.47	-	.54	.48	.43
HypER [9]	.341	250	.520	.376	.252	.465	5798	.522	.477	.436
RotatE [186]	.338	<b>177</b>	.533	.375	.241	.476	<b>3340</b>	<b>.571</b>	.492	.428
ConvR [83]	.350	-	.528	.385	.261	.475	-	.537	.489	<b>.443</b>
VR-GCN [225]	.248	-	.432	.272	.159	-	-	-	-	-
COMPGCN (Proposed Method)	<b>.355</b>	197	<b>.535</b>	<b>.390</b>	<b>.264</b>	<b>.479</b>	3533	.546	<b>.494</b>	<b>.443</b>

Table 9.3: Link prediction performance of COMPGCN and several recent models on FB15k-237 and WN18RR datasets. The results of all the baseline methods are taken directly from the previous papers. We find that COMPGCN outperforms all the existing methods on 4 out of 5 metrics on FB15k-237 and 3 out of 5 metrics on WN18RR. Please refer to Section 9.4.2.1 for more details.

## 9.4.2 Results

In this section, we attempt to answer the following questions.

- Q1. How does COMPGCN perform on link prediction compared to existing methods? (9.4.2.1)
- Q2. What is the effect of using different GCN encoders and choice of the compositional operator in COMPGCN on link prediction performance? (9.4.2.1)
- Q3. Does COMPGCN scale with the number of relations in the graph? (9.4.2.3)
- Q4. How does COMPGCN perform on node and graph classification tasks? (9.4.2.4)

### 9.4.2.1 Performance Comparison on Link Prediction

In this section, we evaluate the performance of COMPGCN and the baseline methods listed in Section 9.4.1.2 on link prediction task. The results on FB15k-237 and WN18RR datasets are presented in Table 9.3. The scores of baseline methods are taken directly from the previous papers [186, 30, 173, 9, 83, 225]. However, for ConvKB, we generate the results using the corrected evaluation code<sup>1</sup>. Overall, we find that COMPGCN outperforms all the existing methods in 4 out of 5 metrics on FB15k-237 and in 3 out of 5 metrics on WN18RR dataset. We note that the best performing baseline RotatE uses rotation operation in complex domain. The same operation can be utilized in a complex variant of our proposed method to improve

<sup>1</sup><https://github.com/KnowledgeBaseCompleter/eval-ConvKB>

Scoring Function (=X) →	TransE			DistMult			ConvE		
Methods ↓	MRR	MR	H@10	MRR	MR	H@10	MRR	MR	H@10
X	0.294	357	0.465	0.241	354	0.419	0.325	244	0.501
X + D-GCN	0.299	351	0.469	0.321	225	0.497	0.344	200	0.524
X + R-GCN	0.281	325	0.443	0.324	230	0.499	0.342	197	0.524
X + W-GCN	0.267	1520	0.444	0.324	229	0.504	0.344	201	0.525
X + COMPGCN (Sub)	0.335	<b>194</b>	0.514	0.336	231	0.513	0.352	199	0.530
X + COMPGCN (Mult)	<b>0.337</b>	233	0.515	<b>0.338</b>	<b>200</b>	<b>0.518</b>	0.353	216	0.532
X + COMPGCN (Corr)	0.336	214	<b>0.518</b>	0.335	227	0.514	<b>0.355</b>	197	<b>0.535</b>
X + COMPGCN ( $\mathcal{B} = 50$ )	0.330	203	0.502	0.333	210	0.512	0.350	<b>193</b>	0.530

Table 9.4: Performance on link prediction task evaluated on FB15k-237 dataset. X + M (Y) denotes that method M is used for obtaining entity (and relation) embeddings with X as the scoring function. In the case of COMPGCN, Y denotes the composition operator used.  $\mathcal{B}$  indicates the number of relational basis vectors used. Overall, we find that COMPGCN outperforms all the existing methods across different scoring functions. ConvE + COMPGCN (Corr) gives the best performance across all settings (highlighted using  $\boxed{\cdot}$ ). Please refer to Section 9.4.2.1 for more details.

its performance further. We defer this as future work.

#### 9.4.2.2 Comparison of Different GCN Encoders on Link Prediction Performance

Next, we evaluate the effect of using different GCN methods as an encoder along with a representative score function (shown in Figure 9.2) from each category: TransE (translational), DistMult (semantic-based), and ConvE (neural network-based). In our results, **X + M (Y)** denotes that method **M** is used for obtaining entity embeddings (and relation embeddings in the case of COMPGCN) with **X** as the score function as depicted in Figure 9.2. **Y** denotes the composition operator in the case of COMPGCN. We evaluate COMPGCN on three non-parametric composition operators inspired from TransE [22], DistMult [221], and HolE [147] defined as

- **Subtraction (Sub):**  $\phi(\mathbf{e}_s, \mathbf{e}_r) = \mathbf{e}_s - \mathbf{e}_r$ .
- **Multiplication (Mult):**  $\phi(\mathbf{e}_s, \mathbf{e}_r) = \mathbf{e}_s * \mathbf{e}_r$ .
- **Circular-correlation (Corr):**  $\phi(\mathbf{e}_s, \mathbf{e}_r) = \mathbf{e}_s * \mathbf{e}_r$

The overall results are summarized in Table 9.4. Similar to Schlichtkrull et al. [171], we find that utilizing Graph Convolutional based method as encoder gives a substantial improvement in performance for most types of score functions. We observe that although all the baseline GCN methods lead to some degradation with TransE score function, no such behavior is observed for COMPGCN. On average, COMPGCN obtains around 6%, 4% and 3% relative increase in MRR with TransE, DistMult, and ConvE objective respectively compared to the best performing

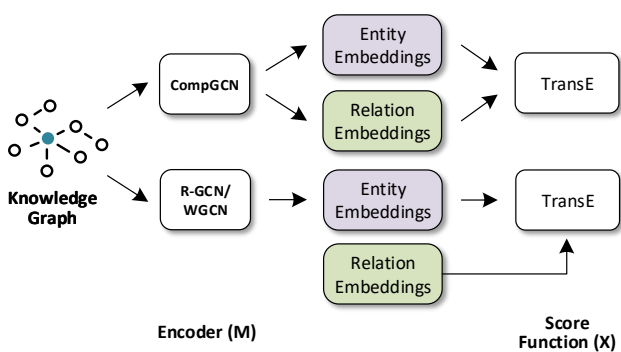


Figure 9.2: Knowledge Graph link prediction with COMPGCN and other methods. COMPGCN generates both entity and relation embedding as opposed to just entity embeddings for other models. For more details, please refer to Section 9.4.2.2

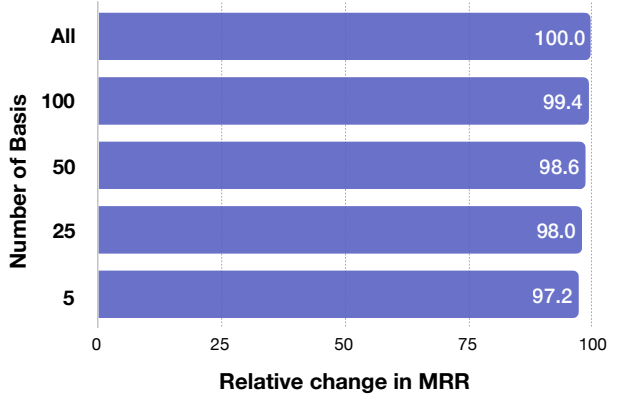


Figure 9.3: Performance of COMPGCN with different number of relation basis vectors on link prediction task. We report the relative change in MRR on FB15k-237 dataset. Overall, COMPGCN gives comparable performance even with limited parameters. Refer to Section 9.4.2.3 for details.

baseline. The superior performance of COMPGCN can be attributed to the fact that it learns both entity and relation embeddings jointly thus providing more expressive power in learned representations. Overall, we find that COMPGCN with ConvE (highlighted using  $\square$ ) is the best performing method for link prediction.

**Effect of composition Operator:** The results on link prediction with different composition operators are presented in Table 9.4. We find that with DistMult score function, multiplication operator (Mult) gives the best performance while with ConvE, circular-correlation surpasses all other operators. Overall, we observe that more complex operators like circular-correlation outperform or perform comparably to simpler operators such as subtraction.

#### 9.4.2.3 Parameter Efficiency of CompGCN

In this section, we analyze the performance of COMPGCN on changing the number of relation basis vectors ( $\mathcal{B}$ ) as defined in Section 9.3. For this, we evaluate the best performing model for link prediction (ConvE + COMPGCN (Corr)) with a variable number of basis vectors. The results are summarized in Figure 9.3. We find that our model performance improves with the increasing number of basis vectors. We note that with  $\mathcal{B} = 100$ , the performance of the model becomes comparable to the case where all relations have their individual embeddings. In Table 9.4, we report the results for the best performing model across all score function with  $\mathcal{B}$  set to 50. We note that the parameter-efficient variant also gives a comparable performance and outperforms the baselines in all settings. This demonstrates that COMPGCN is scalable with

	MUTAG (Node)	AM		MUTAG (Graph)	PCT
Feat*	77.9	66.7		PACHYSAN <sup>†</sup>	<b>92.6 ± 4.2</b>
WL*	80.9	87.4		DGCNN <sup>†</sup>	85.8
RDF2Vec*	67.2	88.3		GIN <sup>†</sup>	89.4 ± 4.7
R-GCN*	73.2	89.3		R-GCN	82.3 ± 9.2
SynGCN	74.8 ± 5.5	86.2 ± 1.9		SynGCN	79.3 ± 10.3
WGCN	77.9 ± 3.2	90.2 ± 0.9		WGCN	78.9 ± 12.0
COMPGCN	<b>85.3 ± 1.2</b>	<b>90.6 ± 0.2</b>		COMPGCN	89.0 ± 11.1
					<b>71.6 ± 12.0</b>

Table 9.5: Performance comparison on node classification (**Left**) and graph classification (**Right**) tasks. \* and † indicate that results are directly taken from [171] and [217] respectively. Overall, we find that COMPGCN either outperforms or performs comparably compared to the existing methods. Please refer to Section 9.4.2.4 for more details.

the increasing number of relations and thus can be utilized for larger graphs effectively.

#### 9.4.2.4 Evaluation on Node and Graph Classification

In this section, we evaluate COMPGCN on node and graph classification tasks on datasets as described in Section 9.4.1.1. The experimental results are presented in Table 9.5. For node classification task, we report accuracy on test split provided by [169], whereas for graph classification, following [220] and [217], we report the average and standard deviation of validation accuracies across the 10 folds cross-validation. Overall, we find that COMPGCN outperforms all the baseline methods on node classification and gives a comparable performance on graph classification task. This demonstrates the effectiveness of incorporating relations using COMPGCN over the existing GCN based models. On node classification, compared to the best performing baseline, we obtain an average improvement of 3% across both datasets while on graph classification, we obtain an improvement of 3% on PCT dataset.

## 9.5 Conclusion

In this chapter, we proposed COMPGCN, a novel Graph Convolutional based framework for multi-relational graphs which leverages a variety of composition operators from Knowledge Graph embedding techniques to jointly embed nodes and relations in a graph. Our method generalizes several existing multi-relational GCN methods. Moreover, our method alleviates the problem of over-parameterization by sharing relation embeddings across layers and using basis decomposition. Through extensive experiments on knowledge graph link prediction, node classification, and graph classification tasks, we showed the effectiveness of COMPGCN over existing GCN methods and demonstrated its scalability with increasing number of relations.

# Chapter 10

## Conclusion and Future Work

The first part of the thesis explored three different ways of addressing the sparsity problem in Knowledge Graphs (KG). We begin with alleviating it through canonicalization, which involves identifying and merging identical nodes in a given KG. For this, we proposed CESI (Canonicalization using Embeddings and Side Information), a novel method for canonicalizing Open KBs using learned embeddings and side information. CESI solves a joint objective to learn noun and relation phrase embeddings while utilizing relevant side information in a principled manner. These learned embeddings are then clustered together to obtain canonicalized noun and relation phrase clusters. The second approach is Relation Extraction which involves using unstructured text for extracting facts to densify KGs. We propose RESIDE, a novel neural network-based model which makes principled use of relevant side information, such as entity type and relation alias, from Knowledge Base, for improving distantly supervised relation extraction. RESIDE employs Graph Convolution Networks for encoding syntactic information of sentences and is robust to limited side information. Finally, the third solution is to use the existing facts to infer new facts in the KG; the task is termed as link prediction. For this, we propose InteractE, a novel knowledge graph embedding method which alleviates the limitations of ConvE by capturing additional heterogeneous feature interactions. InteractE achieves this by utilizing three central ideas, namely feature permutation, checkered feature reshaping, and circular convolution. Through experimental results, we demonstrated the effectiveness of all the proposed methods on several benchmark datasets.

In the second part of the thesis, we showed the effectiveness of utilizing recently proposed Graph Convolutional Networks (GCNs) for exploiting several graph structures in NLP. We demonstrated their effectiveness on two important problems: Document Timestamping and learning word embeddings. For the first problem, we proposed NeuralDater, a GCN based method for document dating which exploits syntactic and temporal structures in the document

in a principled way. To the best of our knowledge, this is the first application of deep learning techniques for the problem of document dating. For the second problem, we proposed SynGCN, a graph convolution-based approach which utilizes syntactic context for learning word representations. SynGCN overcomes the problem of vocabulary explosion and outperforms state-of-the-art word embedding approaches on several intrinsic and extrinsic tasks. We also propose SemGCN, a framework for jointly incorporating diverse semantic information in pre-trained word embeddings. The combination of SynGCN and SemGCN gives the best overall performance.

Finally, in the third part of the thesis, we addressed two significant limitations of existing Graph Convolutional Network-based methods. First, we address the issue of noisy representation of hub nodes in GCNs because of neighborhood aggregation scheme which puts no constraint on the influence neighborhood of a node. For this, we present ConfGCN, confidence based Graph Convolutional Network, which estimates label scores along with their confidences jointly in a GCN based setting. In ConfGCN, the influence of one node on another during aggregation is determined using the estimated confidences and label scores, thus inducing anisotropic behavior to GCN. Apart from this also extend existing GCN models for multi-relational graphs, which are a more pervasive class of graphs for modeling data. We propose CompGCN, a novel Graph Convolutional based framework for multi-relational graphs which leverages a variety of composition operators from Knowledge Graph embedding techniques to embed nodes and relations in a graph jointly. Our method generalizes several existing multi-relational GCN methods. Moreover, our method alleviates the problem of over-parameterization by sharing relation embeddings across layers and using basis decomposition. Through extensive experiments on several tasks, we demonstrated the effectiveness of our proposed solutions.

**Future Works:** An exciting future direction for addressing sparsity in Knowledge Graphs is to utilize contextualized embedding methods such as ELMo [159] and BERT [51] instead of GloVe for obtaining the representation of noun and relation phrases in Open KG canonicalization. Contextualized embedding approaches have been shown to give superior performance than standard word2vec based embeddings for a variety of tasks. However, utilizing them for canonicalization has not been explored so far. Another future work includes extending our proposed model RESIDE to utilize more types of side information. KGs are a vast storehouse of facts, among which a lot of them can be utilized for further improving RE. In Chapter 5, we explored how increasing heterogeneous interaction can help to improve link prediction performance for ConvE method. The same idea can be extended for further improving other state-of-the-art link prediction methods.

For further enhancing performance on document timestamping problem, one can explore

utilizing knowledge graphs which contain information about world events. Exploiting world knowledge for the task is also more close to how a human would approach the problem. In Chapter 7, we explored extending word2vec using Graph Convolutional Networks. However, recently, contextualized embedding methods have been shown to be much more effective. Thus, one can use similar ideas to extend models such as BERT using GCNs for utilizing syntactic information for learning better representation. Finally, existing GCN methods suffer from several other limitations, which have been highlighted by Xu et al. [216] in his work.

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