Abstract

This thesis is a proof of concept for the potential of Variational Auto-Encoder (VAE) on representation learning of real-world Knowledge Graphs (KG). Inspired by successful approaches to the generation of molecular graphs, we experiment and evaluate the capabilities and limitations of our model, the Relational Graph Variational Auto-Encoder (RGVAE), characterized by its permutation invariant loss function. The impact of the modular hyperparameter choices, encoding through graph convolutions, graph matching and latent space prior, are analyzed and the added value is compared. The RGVAE is first evaluated on link prediction, a common experiment indicating its potential use for KG completion. The model is ranked by its ability to predict the correct entity for an incomplete triple. The mean reciprocal rank (MRR) scores on the two datasets FB15K-237 and WN18RR are compared between the RGVAE and the embedding-based model DistMult. To isolate the impact of each module, a variational DistMult and a RGVAE without latent space prior constraint are implemented. We conclude, that neither convolutions nor permutation invariance alter the scoring. The results show that between different settings, the RGVAE with relaxed latent space, scores highest on both datasets, yet does not outperform the DistMult.

Graph VAEs on molecular data are able to generate unseen and valid molecules as a result of latent space interpolation. We investigate, if the RGVAE can yield similar results on relational KG data. The experiment is twofold, first the distance between the latent representation of two triples is linearly interpolated, then each latent dimension is explored in a 95% confidence interval of its Normal distribution. The interpolations reveal, how successful the RGVAE is at disentangling the latent space and assigning each latent dimension to data characterizing features. Both interpolation experiments show that the RGVAE learns to reconstruct the adjacency matrix but fails to disentangle and assign the right node and edge attributes.

The assumption of an uninformative latent representation is confirmed in the last experiment of knowledge generation. For this experiment, we present a new validation method for generated triples from the FB15K-237 dataset. The relation type-constrains of generated triples are filtered and matched with entity types. The observed rate of valid generated triples is insignificantly higher than the random threshold. All generated and valid triples are unseen in both train and test set. A comparison between different latent space priors, using the $\delta$-VAE method, reveals insights in the behavior of the RGVAE’s parameter distribution and indicates a decoder collapse. Finally we analyze the limiting factors of our approach compared to molecule generation and propose solutions for the decoder collapse and successful representation learning of multi-relational KGs.
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1 Introduction

To begin with, we shall clarify the intended ambiguity of this work’s title. Before approaching the question behind this thesis in the context of Representational Learning (RL), we would like to present it from a philosophical point of view of AI safety and potential. One could argue, that we live in times of the fastest advances in science and technology in the history of humanity, therefore making us the Knowledge Generation. While we continuously keep researching and accumulating knowledge, we have historically not been willing to share our knowledge with any other species in this Universe. All scientific milestones are from us, for us.

The rise of AI marked a turning point of this tradition. For the first time we invest in sharing our knowledge and with systems which can act on superhuman dimension. While machine learning models might not be considered a species, they have learned to drive our cars and in fact beaten human intelligence in the game GO [1]. This is not only seen as progress but also as danger. The world’s richest man, Elon Musk, both build his fortune on AI and respects it as humanity’s biggest risk.

Figure 1: Elon Musk in a tweet on AI. Source [2]

Closing the circle to the ambiguity of the title and relating to the popular concern, that AI might reach a point where it does not need humans anymore to keep evolving, we ask the crucial question: Can AI generate knowledge?

1.1 Motivation

A key area of AI is RL, where the model learns to identify and disentangle characteristics and features of the data. Understanding the semantics of the data is specifically important for unsupervised learning of generative models. The task of generating data has been widely explored for images. Computer vision has reached a point where a simple image can be semantically segmented, where objects can be detected and classified and even relations between entities inferred [3].

Advances in the parallel field of graph generation have received less attention, yet showed promising results. Data stored as a graph has a high density of information and rich semantics, which makes it attractive for variational inference. The recent success of Simonovsky et al. [4] on the generation and completion of molecules represented in graph structure, initially inspired our research. Next to molecules, graphs can be used to store knowledge. While real world Knowledge Graphs (KG) have a far higher complexity than molecule graphs, the proposed generative model, Variational Auto-Encoder (VAE) also has proven its capacity to learn from huge datasets with high variance. Inspired by Simonovsky work and motivated by the vision of generating knowledge, we explore the possibilities and limitation of KG generation with VAEs.

1.2 Expected Contribution

The main contributions of this thesis are threefold. The main objective is to proof the hypothesis that a graph VAE can capture, disentangle and reproduce the underlying semantics of a real world KG, secondly. Further we contribute with a novel implementation of a graph matching algorithm and a validation method for generated triples.

In Simonovsky [4] work, small subgraphs with multiple edges are used to represent molecule graphs. In contrast we proof the hypothesis by generating the smallest possible graph of two nodes, also representable as single triple. The VAE is tested and evaluated in several experiments, including link prediction, latent space interpolation and accuracy of generating valid triples.

We compare our results to related KGs methods and investigate the impact of different hyperparameter. The main focus is be on the influence of the graph matching loss function, the encoding through graph convolutions and stochastic inference. Further, we aim to reproduce the success of molecule generation and point therefore continuously the similarities and differences to our work out.

On a lower level, we hope to contribute with our implementation of Cho’s et al. max-pooling graph matching algorithm for tensor batches. While the algorithm has been cited and implemented numerous times,
a working implementation, compatible with deep learning libraries, has to the best of our knowledge not yet been published.

Lastly we introduce a high-level method for evaluating the validity of generated data, which compares the type constraint of the generated triple's predicate with its entity types and reports accuracy. This is made possible by expanding the existing dataset FB15K-237 with entity types from its original KG Freebase. While this scoring method is error prone, it does give an insight into the level representational potential of the model and the syntax coherence of the generated triples. Future work can use this evaluation method to track progress and compare to the baseline.

1.3 Research Question

How successful is a graph VAE in representation learning of real world KG compared to molecule graph data and what is the impact of each major hyperparameter?

2 Related Work

This section presents previous work which inspired and laid the foundation for this thesis. Relevant publication on topics related to this thesis are presented in terms of methods and results. We focus on the fields of relational graph convolutions, graph encoders, and embedding-based link prediction.

2.1 Relational Graph Convolutions

We define a graph as $G = (\mathcal{V}, \mathcal{E})$ with a set of nodes $\mathcal{V}$ and a set of edges $\mathcal{E}$. The set of edges, with each edge connecting node $x$ and $y$, is defined by $\{ (x, y) \mid (x, y) \in \mathcal{V}^2 \land x \neq y \}$ while the constraint $x \neq y$ prohibits self-connections or self-loops, which is optional depending on the graphs function. Moreover, nodes and edges can have features, which contribute additional information about the nodes and their connection. In the literature, these features can be describing attributes and properties, in the context of this work we also use them as indicators to unique entities. Graph convolutions, make use of both these properties and the spectral information in a graph's adjacency matrix. In graph theory spectral properties are the characteristic polynomial, eigenvalues, and eigenvectors of the Laplacian and adjacency matrix $[5]$. Two popular tasks to evaluate the performance of a neural network on graphs, are node classification and link prediction. The first is a classification problem where the model predicts the class of a node. Link prediction is the task of completing a triple by correctly predicting the missing entity at either head or tail the triple. A in-depth explanation follows in section 4.4.

In Kipf et al. paper on graph convolutions $[6]$ a novel Graph Convolution Network (GCN) for semi-supervised classification is introduced. The model takes as input the adjacency matrix and optionally a feature matrix of the graph and predicts the classes of the nodes. Graph convolutions acts directly on the graph structure and are linearly scalable with the number of nodes. The GCN takes as input the adjacency matrix $A \in \mathbb{R}^{n \times n}$ with $n$ being the number of nodes in the graph. In the case of undirected graphs, the adjacency matrix is symmetric. The output is a matrix $H \in \mathbb{R}^{n \times dh}$ where $dh$ are the hidden dimensions or in case of the last layer, the number of classes to predict over. While the authors compare different propagation methods for the graph convolutions, their propagating rule using a first-order approximation of spectral graph convolutions, outperforms all other implementations. Propagation denotes the transformation of the input data between layers of a model. Kipf approximates the eigenvalues of the Laplacian with first order Chebyshev polynomials and circumvents the computationally expensive Eigendecomposition. The renormalization trick normalized the adjacency matrix and adds it to an identity matrix of same size. This keeps the eigenvalues in a range between $[0, 2]$ which again leads to a stable training, avoiding numerical instabilities and vanishing gradients during learning. Additionally the feature information of neighboring nodes is propagated in every layer what shows improvement in comparison to earlier methods, where only label information is aggregated. Kipf and Welling perform node classification on the three citation-network datasets, Citeseer, Cora and Pubmed as well as on the KG dataset NELL. In all classification tasks, their results outperform other recently proposed methods in this field and proves to be computationally more efficient than its competition. For more details on the implementation of graph convolutions we refer to section 3.2.3.

In their publication Modeling Relational Data with Graph Convolutional Networks Schlichtkrull et al. propose a relational graph convolutional network (RGCN) and evaluate it on link prediction on the FB15K-237 and WN18 dataset and node classification on the AIFB, MUTAG, BGS and AM datasets $[7]$. The RGCN, with its encoder properties, is used by itself as node classifier, yet for link prediction it is coupled with a DistMult model acting as decoder which scores triples encoded by the RGCN see figure 2. We go into details of the embedding-based DistMult model in section 2.3.
The RGCN works on dense graphs stored as triples, creating a hidden state for each node. A novel message passing network is layer-wise propagated with the hidden states of the entities. As regularization the authors propose a basis- and block-wise decomposition. While the first aims at an effective weight sharing between different relation types, the second can be seen as a sparsity constraint on the relation type’s weight. The model outperforms embedding based model on the link prediction task on the FB15K-237 dataset and scores competitive on the WN18 dataset. In the node classification task, the model sets state of the art results on the datasets AIFB and AM, while scoring competitive on the remaining. The authors conclude, that the model has difficulties encoding higher-degree hub nodes on datasets with many entities and low amount of classes. This is noticeable as it relates to the WN18RR [8], one of the two datasets used in this thesis.

2.2 Graph VAE

We have seen how graph convolutional neural networks can be combined in an encoder-decoder architecture, resulting in a generative model suitable for unsupervised learning. We present three recent publications with different methods and use cases of a graph generative model, in particular a VAE.

Kipf et al. introduce the the Variational Graph Autoencoder (VGAE), a framework for unsupervised learning on graph-structured data [9]. This generative model uses a GCN as encoder and a simple inner product module as decoder. Similar to the GCN, the VGAE incorporates node features, which significantly improves its performance on link prediction tasks compared to related models. The VGAE uses a two-layer GCN to encode the mean and the logvariance for the stochastic module to sample the latent space representation, more specifically, a latent vector per node. Referring to the above described GCN, the VGAE encoder outputs a latent matrix \( H \in \mathbb{R}^{n \times 2d_z} \) with \( d_z \) denoting the latent dimension. The activation of the inner product of this latent matrix yields the reconstruction of the adjacency matrix. Figure 3 shows how the model learns to cluster nodes according to their class, without these labels being provided to the model during training. This visualization shows that the VGAE learns successful learn an implicit representation of the data. The VGAE with added features outperforms state of the art methods (to the time of publication) Spectral Clustering [10] and Deepwalk [11] in the task of link prediction on the datasets Cora, Citeseer and Pubmed. The authors point out, that a Gaussian prior might be a poor choice combined with the inner-product decoder.

Simonovsky et al. introduce the GraphVAE, which generates a probabilistic fully-connected molecule graph of a predefined maximum size in a one-shot approach [4]. In this context fully-connected denotes that...
all nodes are connected within a graph, in contrast to citation networks where subgraphs can be disconnected from each other. While molecule graphs have a lower node and edge count than citation networks, their edges and nodes are attributed, which constrains each connection. The model includes a standard graph matching algorithm, which finds the optimal permutation between the predicted graph and the ground truth and. The reconstruction loss considers the permutation instead of the raw prediction. In contrast to the previously presented publications, the input to this model is a threefold and sparse graph, defined as $G = (A, E, F)$ with $A$ being the adjacency matrix, $E$ the edge attribute matrix and $F$ the node attribute matrix, with $E$ and $F$ being one-hot encoded. Considering that this method lays the foundation for this thesis, we adopt this notation for our own methods in section 4.5. Figure 4 shows the architecture of the GraphVAE. The encoder is a feed forward network with edge-conditioned graph convolutions [12], which takes as input the target graph $G$ with $n$ nodes. After the convolutions the result is flattened and conditioned on the node labels $y$. A fully-connected neural network encodes the stochastic latent representation, which is constrained by Standard Gaussian prior distribution. Note that in contrast to the GCN, which encodes one latent vector per node, the GraphVAE instead encodes a latent representation of the whole graph. This latent representation is again conditioned on the node labels $y$ and propagated through the decoder in form of a fully-connected neural network. The decoder reconstructs the latent representation to the graph prediction. The threefold decoder output is matched with the target using graph matching algorithm, which we discuss further in section 3.3. The matched and permuted graph is then used for the reconstruction term of the GraphVAE loss. It should be noted, that, while the size of the target and prediction graph are fixed, they do not necessarily have to match. While this approach seems promising, it is limited by the maximum graph size, which has been experimented with up to a node count of 40.

The model is trained on the QM9 dataset, containing the graph structure of 134k organic molecules with experiments on latent space dimension in the range of $[20, 80]$. On the free generation task, about 50% of the generated molecules are chemically valid and thereof remarkably 60% are not included in the training dataset. When testing the model for robustness, it showed little disturbance when adding Gaussian noise to the input graph $G$. The authors conclude that the problem of generating graphs from a continuous embedding was addressed successfully and that the GraphVAE performs better on small molecules, implying a low node count per graph.

Until here the presented models generate graphs in a single propagation through the model. For completeness we also present a successful approach for graph generation in autoregressive manner. Belli et al. introduce named approach for image-conditioned graph generation of road network graphs [13]. While we focus on the generative model, their contribution ranges wider, namely the introduction of the graph-based roadmap dataset Toulouse Road Network and the task specific distance metric StreetMover. The authors propose the Generative Graph Transformer (GGT) a deep autoregressive model that makes use of attention mechanisms on images, to tackle the challenging task of road network extraction from image data. The GGT has a encoder-decoder architecture, with a CNN as encoder, taking the grayscale image as input signal and predicting a conditioning vector. The decoder is a self-attentive transformer, which takes as input the encoded condition vector and a hidden representation of the adjacency matrix $A$ and feature vector $X$ of the previous step. The adjacency matrix here indicates the links between steps and the features are normalized coordinates. A multi-head operator outputs the hidden representation of $A$ and $X$ which finally are decoded by a MLP to the graph representation. For the first step, a empty hidden representation is fed into the decoder. The model terminates the recurrent graph generation by predicting a end-of-sequence token, which signalizes the end of the graph. During learning, the generated graphs are matched to the target graphs using the StreetMover metric, based on the Sinkhorn
distance. The authors attribute StreetMover as a scalable, efficient and permutation-invariant metric for graph comparison. The successful results of the experiments performed, show that this novel approach is suitable for the task of road network extraction and could yield similar success in graph generation task of different fields. While this publication does not directly align with the previously presented work, we find it of added value to present alternative approaches on our topic.

2.3 Embedding-Based Link Prediction

Finalizing this chapter, we look at embedding-based methods on KGs. Compared to the previously presented research, embedding models have a much simpler architecture and can be trained computationally very efficient on large graphs. Embedding-based models can only operate on triples, meaning a KG is represented as a set of triples with indices pointing to the unique entity and relation in the graph. Despite their simplicity, they achieve great results on node and link prediction tasks.

Already in 2013 Bordes et al. introduced in their paper Translating Embeddings for Modeling Multi-relational Data the low-dimensional embedding model TransE [14]. The core idea of this model is that relations can be represented as translations in the embedding space. Entities are encoded to a low-dimension embedding space and the relation is represented as vector between the head and tail entity. The assumption is that for correct triples the model learns to reduce the Euclidean distance between head and tail entity by placing them closer together in the embedding space. This results in correct triples having a lower norm of the relational vector than corrupted triples. Using this property, the model can predict the missing entity in link prediction.

The models loss function takes a set of corrupted triples for every triples in the training set and subtracts the translation vector of the corrupted triple in embedding space from the translation vector of the correct triple with added margin. To minimize the loss, the model has to place entities of correct triples closer together in embedding space. We think of a triple as \((s, r, o)\) and \((e_s, e_r, e_o)\) as its embedded representation, \(d()\) the Euclidean distance, \(\gamma\) the positive margin and \(S\) and \(S'\) as sets of correct and corrupt triples, the loss function of TransE is

\[
\mathcal{L} = \sum_{s} \sum_{s'} \left[ \gamma + d(s + r, o) - d(s' + r, o') \right].
\]

The model is trained on a subset of the KGs Freebase and Wordnet, which is also the source for the datasets used in this thesis. TransE’s link prediction results on both head and tail outperformed other competing methods of the time, such as RESCAL [15].

In 2015, Yang et al. proposed a similar, yet better performing KG embedding method [16]. Their model DistMult captures relational semantics by matrix multiplication of the embedded entity representation and uses a bilinear learning objective. The main difference to TransE is the bilinear scoring function \(d^b()\). Bilinear is indicated by the exponent \(b\) and connotes the functions score-invariance of swapping the triples head and tail entity. For the embedding space representation of subject and object \(e_s\) and \(e_o\) and a diagonal matrix \(\text{diag}(e_r)\) with the embedded relation \(e_r\) on the diagonal, the scoring function is

\[
d^b((e_s, e_r, e_o)) = e_s \text{diag}(e_r)e_o.
\]

The publication goes on to explore the options of embedding-based rule extraction from KGs. Concluding, the authors state that the prediction scores achieved with the embeddings learned from the bilinear objective not only outperform the state of the art in link prediction but can also capture compositional semantics of relations and extract Horn rules using compositional reasoning.

In a more recent publication, Ruffinelli et al. present a comprehensive review of KG embedding models such as TransE and DistMult, coupled with state of the art techniques in deep learning. The authors start by pointing out the similarities and differences of most models. While all methods share the same embedding approach, they differ in their scoring function and their original hyperparameter search. The authors perform a quasi-random hyperparameter search on the five models RESCAL, TransE, DistMult, ComplEx and ConvE, which each use a characteristically different loss function. They are compared by their MRR and Hits@10 scores on the two datasets FB15K-237 and WN18. Since these metrics and datasets are used later on in our research, they are explained in section 5.1(datasets) and 4.4(metrics). The tuned models report a higher MRR score of up to 24% compared to their first reported performance. The authors conclude that simple KG embedding methods can show strong performance when trained with state of the art techniques what indicates that higher complexity is not necessary. The optimal model configurations, which were found by a random search of the hyperparameter space, are included in this publication.
3 Background

This section derives and explains the techniques, which form the backbone of this research. For fundamental background on machine learning and probability theory we refer the reader to Bishops book \cite{17}. We begin with introducing the VAE and its differences to a normal autoencoder. Further we show how convolutional layers can act on graphs and how these layers can be used in an encoder model, the GCN. Building on these modules, we present the main model for this thesis, the Relational Graph VAE (RGVAE). Finally, we present a popular graph matching algorithm, which is intended to match prediction and target graph \cite{18}.

3.1 Knowledge Graph

'Knowledge graph' has become a popular term. Yet, the term is so broad, that its definitions varies depending on domain and use case \cite{19}. In this thesis we focus on KGs in the context of relational machine learning.

Both datasets of this thesis are derived from large-scale KG data in RDF format. The Resource Description Framework (RDF), originally introduced as infrastructure for structured metadata, is used as general description framework for the semantic-web applications \cite{20}. It involves a schema-based approach, meaning that every entity has a unique identifier and possible relations limited to a predefined set. The opposite schema-free approach is used in OpenIE models, such as AllenNLP \cite{21}, for information extraction. These models generate triples from text based on NLP parsing techniques which results in an open set of relations and entities. In this thesis a schema-based framework is used and triples are denote as \((s, r, o)\). A exemplar triple from the FB15K-237 dataset is

\[/m/02mjmr, /people/person/place_of_birth, /m/02hrh0_.\]

A human readable format of the entities is given by the \textit{id2text} translation of Wikidata \cite{22}.

- Subject \(s\): /m/02mjmr Barack Obama
- Relation/Predicate \(r\): /people/person/born-in
- Object \(o\): /m/02hrh0_ Honolulu

For all triples, \(s\) and \(o\) are part of a set of entities, while \(r\) is part of a set of relations. This is sufficient to define a basic KG \cite{23}.

Schema-based KG can include type hierarchies and type constraints. Classes group entities of the same type together, based on common criteria, e.g. all names of people can be grouped in the class 'person'. Hierarchies define the inheriting structure of classes and subclasses. Picking up our previous example, 'spouse' and 'person' would both be a subclass of 'people' and inherit its properties. At the same time the class of an entity can be the key to a relation with type constraint, since some relations can only be used in conjunction with entities fulfilling the constraining type criteria.

These schema based rules of a KG are defined in its ontology. Here properties of classes, subclasses and constraints for relations and many more are defined. Again, we have to differentiate between KGs with open-world or closed-world assumption. In a closed-world assumption all constraints must be sufficiently satisfied before a triple is accepted as valid. This leads to a huge ontology and makes it difficult to expand the KG. On the other hand open-world KGs such as Freebase, accept every triple as valid, as long as it does not violate a constrain. This again leads inevitably to inconsistencies within the KG, yet it is the preferred approach for large KGs. In context of this thesis we refer to the ontology as semantics of a KG, we research if our model can capture the implied closed-world semantics of an open-world KG \cite{23}.

3.2 Graph VAE

Since the graph VAE is a adaptation of the original VAE, we start by introducing the original version, which is unrelated to graph data. Furthermore we present each of the different modules, which compose the final model. This includes the different graph encoders as well as sparse graph loss functions. We define the notation upfront for this chapter, which touches upon three different fields. For the VAE and MLP we consider data in vector format. A bold variable denotes the full vector, e.g. \(\mathbf{x}\) and variable with index denotes the element at that index, e.g. for the vector element at index \(i\) we denote \(x_i\). Graphs are represented in matrices, which are denoted in capital letters. A typically denotes the adjacency matrix, from there on paths split and we use different notations for different methods. While \(X\) is described as feature matrix, we have to specify further when it comes to Simonovsky's GraphVAE, where \(E\) is the edge attribute and \(F\) the node attribute matrix. The reason we change from features to attributes, is due to the singularity, also one-hot encoding of attributes per node/edge, in contrast to features which can be numerous per node.
3.2.1 VAE

The VAE as first presented by [24] is an unsupervised generative model in the form of an autoencoder, consisting of an encoder and a decoder. Its architecture differs from a common autoencoder by having a stochastic module between encoder and decoder. The encoder can be represented as recognition model with the probability $p_{\theta}(z \mid x)$ with $x$ being the variable we want to inference and $z$ being the latent representation given an observed value of $x$. The encoder parameters are represented by $\theta$. Similarly, we denote the decoder as $p_{\theta}(x \mid z)$, which given a latent representation $z$ produces a probability distribution for the possible values, corresponding to the input of $x$. This is the base architecture of all our models in this thesis.

The main contribution of the VAE is to be a stochastic and fully backpropagatable generative model. This is possible due to the reparametrization trick. Sampling from the latent prior distribution, creates stochasticity inside the model, which can not be backpropagated and makes training of the encoder impossible. By placing the stochastic module outside the model, it becomes fully backpropagatable. We use the predicted encoding as mean and variance for a Gaussian normal distribution, from which we then sample $\epsilon$, which acts as external parameter and does not need to be backpropagated and updated.

![Figure 5: Representation of the VAE as Bayesian network, with solid lines denoting the generator $p_\theta(z)p_\theta(x \mid z)$ and the dashed lines the posterior approximation $q_\phi(z \mid x)$][22].

Figure 5 shows, that the true posterior $p_\theta(z \mid x)$ is intractable. To approximate the posterior, we assume a Standard Gaussian prior $p(z)$ with a diagonal covariance, which gives us the approximated posterior

$$\log q_\phi(z \mid x) = \log \mathcal{N}(z; \mu(x), \sigma(x)^2 I).$$

Now variational inference can be performed, which allows both $\theta$ the generative and $\phi$ the variational parameters to be learned jointly. Using Monte Carlo estimation of $q_\phi(z \mid x_1)$ we get the variational estimated lower bound (ELBO)

$$\mathcal{L}(\theta, \phi; x_1) = -D_{KL}(q_\phi(z \mid x_1) \parallel p_\theta(z)) + E_{q_\phi(z \mid x_1)}[\log p_\theta(x_1 \mid z)].$$

We call the first term the regularization term, as it encourages the approximate posterior to be close to the Gaussian normal prior. This term can be integrated analytically and does not require an estimation. The KL-divergence is a similarity measure between two distributions, resulting in zero for two equal distributions. Thus, the model gets penalized for learning a encoder distribution $p_\theta(z \mid x)$ which is not Standard Gaussian.

Higgins et al. present a constrained variational framework, the $\beta$-VAE [25]. This framework proposes an additional hyperparameter $\beta$ which acts as factor on the regularization term. The new ELBO including $\beta$ is

$$\mathcal{L}(\theta, \phi; x_1) = -\beta (D_{KL}(q_\phi(z \mid x_1) \parallel p_{theta}(z))) + E_{q_\phi(z \mid x_1)}[\log p_\theta(x_1 \mid z)].$$

For $\beta = 0$ the original VAE is restored and for $\beta > 0$ the influence of the regularization term on the ELBO is emphasized. Thus the model prioritizes to learn the approximate posterior $p_\theta(z \mid x)$ even closer to the Standard Gaussian prior. In the literature this results in a disentanglement of the latent space qualitatively outperforms the original VAE in representation learning of image data.

The second term represents the reconstruction error, which requires an estimation by sampling. This means using the decoder to generate samples from the latent distribution. In the context of the VAE, these probabilistic samples are the models output, thus, the reconstruction error the similarity between prediction and target [24].

Once the parameters $\phi$ and $\theta$ of the VAE are learned, the decoder can be used on its own to generate new samples from $p_\theta(x \mid z)$. Conventionally, a latent input signal is sampled from $\mathcal{N}$ with $d_z$ being the latent dimension. In the case of discrete binary data, each element of the generated sample is used as probability parameter $p$ for a Bernoulli $\mathcal{B}(1, p)$, from which the final output is sampled. In the case of categorical data, e.g. one-hot encoding, the final output is either sampled from a Categorical distribution with the prediction as probability parameters of each class, or simply sampled with the Argmax operator [26].
3.2.2 MLP

The Multi-Layer Perceptron (MLP) was the first of its kind, introducing a machine-learning model with a hidden layer between the input and the output. Its properties as universal approximator have been discovered and widely studied since 1989. While we presume that the reader interested in the topic of this thesis does not require a definition of the MLP, it is included for completeness, as we also define the GCN encoder, which both act as encoder and decoder for our final model, and contribute different hyperparameters.

The MLP takes a linear input vector of the form \(x_1, ..., x_D\) which is multiplied by the weight matrix \(W^{(1)}\) and then activated using a non-linear function \(h()\), which results in the hidden representation of \(x\). Due to its simple derivative, mostly the rectified linear unit (ReLU) function is used as activation. The hidden units get multiplied with the second weight matrix, denoted \(w^{(2)}\) and finally transformed by a sigmoid function \(\sigma()\), which produces the output. Grouping weight and bias parameter together we get the following equation for the MLP

\[
y_{k}(x, w) = \sigma \left( \sum_{j=0}^{M} \sum_{i=0}^{D} w^{(2)}_{kj} h \left( \sum_{i=0}^{D} w^{(1)}_{ji} x_{i} \right) \right)
\]

for \(j = 1, \ldots, M\) and \(k = 1, \ldots, K\), with \(M\) being the total number of hidden units and \(K\) of the output.

Since the sigmoid function returns a probability distribution for all classes, the MLP can have the function of a classifier. Instead of the initial sigmoid function, it was found to also produce good results for multi label classification activating the output through a Softmax function instead. Images or higher dimensional tensors can be processed by flattening them to a one dimensional tensor. This makes the MLP a flexible and easy to implement model \cite{17}.

3.2.3 Graph convolutions

Convolutional layers benefit from the symmetry of data, correlation between neighboring datapoints. Convolutional Neural Nets (CNN) are powerful at classification and object detection on image. Neighboring pixel in an image are not independent and identically distributed i.i.d.) but rather are highly correlated. Thus, patches of datapoints let the CNN infer and detecting local features. The model can further merged those to high-level features, e.g. a face in an image \cite{17}. Similar conditions hold for graphs. Nodes in a graph are not i.i.d. and allow inference of missing node or link labels.

Different approaches for graph convolutions have been published. Here we present the graph convolution network (GCN) of \cite{6}. We consider \(f(X, A)\) a GCN with an undirected graph input \(G = (V, E)\), where \(v_i \in V\) is a set of \(n\) nodes and \((v_i, v_j) \in E\) the set of edges. The input is a sparse graph representation, with \(X\) being a node feature matrix and \(A \in \mathbb{R}^{n \times n}\) being the adjacency matrix, defining the position of edges between nodes. In the initial case of no self-loops, the adjacency’s diagonal is filled resulting in \(A = A + I_n\). The graph forward pass through the convolutional layer \(l\) is then defined as

\[
H^{(l+1)} = \sigma \left( \sum_{i \in n} \frac{A_{i}^T i}{\|A_{i}||} H^{(l)} W^{(l)} \right).
\]

The adjacency is row-wise normalized for each node. \(W^{(l)}\) is the layer-specific weight matrix and contains the learnable parameters. \(H\) returns then the hidden representation of the input graph \cite{7}. The GCN was first introduced as node classifier, predicting a probability distribution over all classes for each node in the input graph. Thus, the output dimensions are \(Z \in \mathbb{R}^{n \times d_z}\) for the GCN prediction or latent representation matrix \(Z\). Let \(\tilde{A}\) be the normalized adjacency matrix, then the full equation for a two layer GCN is

\[
Z = f(X, A) = \text{softmax} \left( \tilde{A} \text{ReLU} \left( \tilde{A}XW^{(0)} \right) W^{(1)} \right).
\]

3.2.4 Graph VAE

We use the presented modules to compose the RGVAE. While the approaches from the literature for graph generative models differ in terms of the model and graph representation, we focus on the GraphVAE architecture presented by Simonovsky \cite{4}. A sparse graph model with graph convolutions.

Simonovsky’s GraphVAE follows the characterizing encoder decoder architecture. The encoder \(q_{\phi}(z | G)\) takes a graph \(G\) as input, on which graph convolutions are applied. After the convolutions the hidden representation is flattened and concatenated with the node label vector \(y\). A MLP encodes the mean \(\mu(z)\) and

\footnote{For symmetric normalization we use \(D^{-\frac{1}{2}} \tilde{A} D^{-\frac{1}{2}}\) with \(\tilde{D}_{ii} = \sum_{j} \tilde{A}_{ij}\).}
logvariance $\sigma^2(\mathbf{x})$ of the latent space distribution. Using the reparameterization trick the latent representation is sampled.

For the decoder $p_\theta(\mathbf{x} \mid G)$ the latent representation is again concatenated with the node labels. The decoder architecture for this model is a MLP with the same dimension as the encoder MLP but in inverted order, which outputs a flat prediction of $\hat{G}$, which is split and reshaped in to the sparse matrix representation.

Simonovsky’s GraphVAE [4] is optimized for molecular data. Our aim is to set a proof of concept with the RGVAE for multi-relational KGs, thus, the structure of the GraphVAE is adopted but reduced to a minimum viable product by dropping the conditioning on the node labels and instead using the node attributes as pointers towards the corresponding entity in $\mathcal{E}$. By using node attributes as unique pointers, we exclude any class or type information about the entity. Simplifying further we drop the convolutional layer and directly flatten $G$ as input for the MLP encoder. To isolate the impact of graph convolutions as encoder for the RGVAE, we make the choice between MLP or GCN as encoder a hyperparameter.

![Figure 6: Architecture of the RGVAE.](image)

In figure 6 the concept of the RGVAE is displayed. Each datapoint $G(A,E,F)$ is a subgraph from the KG dataset. Note that the model propagates batches instead of single datapoints. The RGVAE can generate graphs $\hat{G}(\hat{A},\hat{E},\hat{F})$ by sampling from the approximated posterior distribution $p_\theta(G \mid z)$. Since it predicts on closed sets of relations and entities, the generated subgraphs are either unseen and complement the KG or are already present in the dataset. The subgraph are sparse with $n$ nodes. A single triple being $n = 2$ and a subgraph representation $2 < n < 40$, where $n = 40$ was the explored maximum for the GraphVAE [4].

### 3.3 Graph Matching

In this subsection we explain the term permutation invariance and its impact on the RGVAE’s loss function. Further we present a $k$-factor graph matching algorithm for general graphs with edge and node attributes and the Hungarian algorithm as solution for the NP-hard problem of linear sum assignment. Finally we derive the full loss of the RGVAE when applying the calculated permutation matrix to the models prediction.

#### 3.3.1 Permutation Invariance

A visual example of permutation invariance is the image generation of numbers. If the loss function of the model would not be permutation invariant, the generated image could show a perfect replica of the input number, but being translated by one pixel the loss function would penalize the model. Geometrical permutations can be translation, scale or rotation around any axis.

In the context of sparse graphs the most common, and relevant permutation for this thesis, is the position of a link in the adjacency matrix. By altering its position through matrix-multiplication of the adjacency and the permutation matrix, the original link can change direction or turn into a self-loop. When matching graphs with more $n > 2$, permutation can change the nodes which the link connects. Further it is possible to match graphs only on parts of the graph, $k$-factor, instead of the full graph. In the case of different node counts between target and prediction graph, the target graph can be fully (1-factor) matched with the larger prediction graph.

In context of this thesis, a model or a function is called permutation invariant, if it can match any permutation of the original. This allows the model a wider spectrum of predictions instead of penalizing it on the element-wise correct prediction of the adjacency.
Graph matching of general (not bipartite) graphs is a nontrivial task. Inspired by Simonovsky’s approach \cite{Simonovsky2018}, the RGVAE uses the max-pool algorithm, which can be effectively integrated in its loss function. Presented in \cite{Borgwardt_2006} in the context of computer vision and successful in matching feature point graphs in an image. It first calculates the affinity between two graphs, considering node and edge attributes, then applies edge-wise maxpooling to reduce the affinity matrix to a similarity matrix. Cho et al. praise the max-pool graph matching algorithm as resilient to deformations and highly tolerant to outliers compared to the mean or sum alternatives.

The output is a normalized similarity matrix in continuous space of the same shape as the target adjacency. The similarity matrix is calculated as a linear assignment problem. Simonovsky improves the complexity of the update step from $O(n^3)$ to $O(n)$ and thus, reduces the total complexity to $O(n^2)$. This algorithm has a complexity of $O(n^3 \cdot k)$, thus is not applicable to complete KGs but only to subgraphs with limited number of nodes per graph \cite{Simonovsky2018}.

3.3.2 Max-Pool Graph matching algorithm

Graph matching of general (not bipartite) graphs is a nontrivial task. Inspired by Simonovsky’s approach \cite{Simonovsky2018}, the RGVAE uses the max-pool algorithm, which can be effectively integrated in its loss function. Presented in \cite{Borgwardt_2006} in the context of computer vision and successful in matching feature point graphs in an image. It first calculates the affinity between two graphs, considering node and edge attributes, then applies edge-wise maxpooling to reduce the affinity matrix to a similarity matrix. Cho et al. praise the max-pool graph matching algorithm as resilient to deformations and highly tolerant to outliers compared to the mean or sum alternatives.

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3.3.3 Hungarian algorithm

Starting with the normalized similarity matrix $X^*$, we reformulate the aim of finding the discrete permutation matrix as a linear assignment problem. Simonovsky et al. \cite{Simonovsky2018} use for this purpose an optimization algorithm, the so called Hungarian algorithm. It original objective is to optimally assign $n$ resources to $n$ tasks, thus $k - n$ rows of the permutation matrix are left empty. The cost of assigning task $i \in \mathbb{N}^{[0,n]}$ to $a \in \mathbb{N}^{[0,k]}$ is stored in $x_{ia}$ of the cost matrix $C \in \mathbb{N}^{n \times k}$. By assuming tasks and resources are nodes and taking $C = 1 - X^*$ we get the continuous cost matrix $C$. This algorithm has a complexity of $O(n^4)$, thus is not applicable to complete KGs but only to subgraphs with limited number of nodes per graph \cite{Simonovsky2018}.

The core of the Hungarian algorithm consist of four main steps, initial reduction, optimality check, augmented search and update. The presented algorithm is a popular variant of the original algorithm and improves the complexity of the update step from $O(n^2)$ to $O(n)$ and thus, reduces the total complexity to $O(n^3)$. Since throughout this thesis $n = k$ we imply the reduction step and continue with a quadratic cost matrix $C$. The following notation is solely to derive the Hungarian algorithm and does not apply to our graph
data. The algorithm takes as input a bipartite graph \( G = (V, U, E) \) and the cost matrix \( C \in \mathbb{R}^{n \times n} \). \( G \) is bipartite because it considers all possible edges in the cost matrix in one direction and no self-loops. \( V \in \mathbb{R}^n \) and \( U \in \mathbb{R}^n \) are the resulting sets of nodes and \( E \in \mathbb{R}^n \) the set of edges between the nodes. The algorithm’s output is a discrete matching matrix \( M \). To avoid two irrelevant pages of pseudocode, the steps of the algorithm are presented in the following short summary [30].

1. Initialization:
   
   (a) Initialize the empty matching matrix \( M_0 = \emptyset \).
   
   (b) Assign \( \alpha_i \) and \( \beta_i \) as follows:
       \[
       \forall v_i \in V, \quad \alpha_i = 0 \\
       \forall u_i \in U, \quad \beta_j = \min_i (c_{ij})
       \]

2. Loop \( n \) times over the different stages:
   
   (a) Each unmatched node in \( V \) is a root node for a Hungarian tree with completed results in an augmentation path.
   
   (b) Expand the Hungarian trees in the equality subgraph. Store the indices \( i \) of \( v_i \) encountered in the Hungarian tree in the set \( I^* \) and similar for \( j \) in \( u_j \) and the set \( J^* \). If an augmentation path is found, skip the next step.
   
   (c) Update \( \alpha \) and \( \beta \) to add new edges to the equality subgraph and redo the previous step.

   \[
   \theta = \frac{1}{2} \min_{i \in I^*, j \in J^*} (c_{ij} - \alpha_i - \beta_j) \\
   \alpha_i \left\{ \begin{array}{ll}
   l \alpha_i + \theta & i \in I^* \\
   \alpha_i - \theta & i \notin I^*
   \end{array} \right.
   \\
   \beta_j \left\{ \begin{array}{ll}
   l \beta_j - \theta & j \in J^* \\
   \beta_j + \theta & j \notin J^*
   \end{array} \right.
   
   (d) Augment \( M_{k-1} \) by flipping the unmatched with the matched edges on the selected augmentation path. Thus \( M_k \) is given by \((M_{k-1} - P) \cup (P - M_{k-1})\) and \( P \) is the set of edges of the current augmentation path.

3. Output \( M_n \) of the last and \( n^{th} \) stage.

### 3.3.4 Graph Matching VAE Loss

Coming back to our generative model, we now explain how the loss function needs to be adjusted to work with graphs and graph matching, which results in a permutation invariant graph VAE.

The normal VAE maximizes the evidence lower-bound or, in a practical implementation, minimizes the upper-bound on negative log-likelihood. Using the notation of section 3.3.1 the graph VAE loss is

\[
L(\phi, \theta; G) = \mathbb{E}_{q_\phi(z|G)}[-\log p_\theta(G \mid z)] + \beta (\text{KL} [q_\phi(z \mid G) \parallel p(z)]).
\]

The loss function \( L \) is a combination of reconstruction term and regularization term. The regularization term is the KL divergence between a standard normal distribution and the latent space distribution of \( z \). The change to graph data does not influence this term to graphs. The reconstruction term is the cross entropy between prediction and target, binary for the adjacency matrix and categorical for the edge and node attribute matrices.

The predicted output of the decoder is split in three parts and while \( \tilde{A} \) is activated through sigmoid, \( \tilde{E} \) and \( \tilde{F} \) are activated via edge- and nodewise Softmax. For the case of \( n < k \), the target adjacency is permuted \( A' = XAX^T \), so that the model can backpropagate over the full prediction. Since \( E \) and \( F \) are categorical, permuting prediction or target yields the same cross-entropy. Following Simonovsky’s approach [4] we permute.
the prediction, \( \tilde{F} = X^T \tilde{F} \) and \( \tilde{E}_{i, l} = X^T \tilde{E}_{i, l} X \). Let \( l \) be the one-hot encoded edge attribute vector which is permuted. These permuted subgraphs are then used to calculate the maximum log-likelihood estimate [4]:

\[
\log p(A' | z) = 1/k \sum_a A'_{a,a} \log \tilde{A}_{a,a} + (1 - A'_{a,a}) \log \left( 1 - \tilde{A}_{a,a} \right) + \\
+ 1/k(k - 1) \sum_{a \neq b} A'_{a,b} \log \tilde{A}_{a,b} + (1 - A'_{a,b}) \log \left( 1 - \tilde{A}_{a,b} \right)
\]

(13)

\[
\log p(F | z) = 1/n \sum_i \log F'T_i \tilde{F}'_i
\]

(14)

\[
\log p(E | z) = 1/(\|A\|_1 - n) \sum_{i \neq j} \log E'_{i,j} \tilde{E}'_{i,j}
\]

(15)

The normalizing constant \( 1/k(k - 1) \) takes into account the no self-loops restriction, thus an edge-less diagonal. In the case of self-loops this constant is \( 1/k^2 \). Similar \( 1/(\|A\|_1 - n) \) for \( \log p(E | z) \) where \(-n\) accounts for the edge-less diagonal and in case of self-loops is discarded, resulting in \( 1/(\|A\|_1) \).

### 3.4 Ranger Optimizer

Finalizing this chapter, we explain the novel deep learning optimizer Ranger. Ranger combines Rectified Adam (RAdam), lookahead and optionally gradient centralization. Let us briefly look into the different components. RAdam is based on the popular Adam optimizer. It improves the learning by dynamically rectifying Adam’s adaptive momentum. This is done by reducing the variance of the momentum, which is especially large at the beginning of the training. Thus, leading to a more stable and accelerated start [31]. The Lookahead optimizer was inspired by recent advances in the understanding of loss surfaces of deep neural networks, thus proposes an approach where, a second optimizer estimates the gradients behavior for the next steps on a set of parallel trained weights, while the number of ‘looks ahead’ steps is a hyperparameter. This improves learning and reduces the variance of the main optimizer [32]. The last and most novel optimization technique, Gradient Centralization, acts directly on the gradient by normalizing it to a zero mean. Especially on convolutional neural networks, this helps regularizing the gradient and boosts learning. This method can be added to existing optimizers and can be seen as constrain of the loss function [33]. Concluding we can say that Ranger is a state of the art deep learning optimizer with accelerating and stabilizing properties, incorporating three different optimization methods, which synergize with each other. Considering that generative models are especially unstable during training, we see Ranger as a good fit for this research.

### 4 Methods

This section describes the methods used for the implementation of the RGVAE and in the experiments of this thesis. We begin explaining the formatting and preprocessing of the data, and introduce the VanillaRGVAE, focusing on the encodervations and on the decoder as generator. Furthermore our implementation of the batch-wise max-pooling graph matching algorithm is presented as well as combined with the RGVAE’s loss function. Finally, we describe the link prediction pipeline which is the first experiment which the RGVAE is evaluated on. Our model implementation and experiments are written in Python using PyTorch, a high-performance deep-learning library [34]. All experiments are performed in a fully reproducible manner and the modular implementation of the RGVAE is meant to be reused and further developed in future work. The code is openly available on Github. [3]

### 4.1 Knowledge graph data

All our presented methods operate on KG data. While data from other graph domains is possible, this work focuses solely on datasets in triple format. We explain the sparse graph representation, which is the input format for our model and how to preprocess the original KG triples to match that format.

#### 4.1.1 Graph Representation

This work uses the graph representation \( G(A, E, F) \), where \( A \) denotes the adjacency matrix, \( E \) the edge feature matrix and \( F \) the node feature matrix. This input format for the model architectures as presented in section 3.2.4 The graph is binary and each matrix batch is stored as separate tensor.

[3]https://github.com/INDElab/rgvae
The adjacency matrix $A$ takes the shape $(n \times n)$ with $n$ being the number of nodes in our graph/subgraph. While most of previous work would only allow edges on the upper triangular adjacency matrix and fill the diagonal with ones, we chose a less constrained representation, which we assume is a better fit for representing KGs. In particular, we allow self-loops, meaning a triple where object and subject are the same entity and our relations are directed and can be inverted. Thus $A$ can have a positive signal at any position $A_{ij} \in \mathbb{R}^{n \times n}$, indicating a directed edge between node of index $i$ and node of index $j$, while $A_{ij}$ differs from $A_{ji}$.

The edge attribute matrix $E$ takes the shape $(n \times n \times d_e)$ with $d_e$ being the number of unique entities in our dataset. For each possible edge in the adjacency matrix we have a one hot encoded vector pointing to the unique relation in the dataset. Stacking these vectors leads to the three dimensional matrix $E$.

The shape of node attributes matrix $F$ is $(n \times d_e)$ with $d_e$ being the number of node attributes describing the nodes. Considering that we split the KG in subgraphs, we use the entity index as node attribute, making it possible to assign every node in a subgraph to the entity in the full KG. Thus, the number of node attributes $d_e$ equals the number unique entities in our dataset. Again the node attributes are one hot encoded vectors, which concatenated result in the two dimensional $F$ matrix.

### 4.1.2 Preprocessing

Our datasets consist of three tab separated value files full of triples for training, validation and testing. The preprocessing steps convert the triples to subgraphs $G(A,E,F)$ and during postprocessing back into triple format as well as a human readable translation. Best practices of research are followed by withholding the test set until the final run.

From all three sets, we create a set of all occurring entities and similar set for the relations. Now we can define our dimensions $d_e$ and $d_r$. For both sets we create two dictionaries index-2-entity and entity-2-index which map back and forth between numerical index and the string representation of the entity (similar for the relation set). These dictionaries are used to create a train and test set of triples with numeric indices. Depending if we are in the final testing stage or not, we include all triples from the training and evaluation file in the training set and use the triples in the testing file as test set, or we ignore the triples in the test file and use the evaluation file triples as test set.

Further we create two dictionaries, head and tail which for all occurring subject and relation combination, contain all entities which would complete it to a real triple in our dataset (similar for all relation and object combinations). This allows us to filter true triples, which reduces the score bias for link prediction and evaluates the ratio of unseen triples for graph generation.

The final step of preprocessing is a function, which takes a batch of numerical triples and converts them to a batch of binary, multidimensional tensors $A$, $E$ and $F$. While this might sound easy for only one triple per graph, it proves more complex for graphs with $n > 2$ facing exemption cases such as self loops or an entity occurring in two triples. We solve this by creating a separate set for head and tail entities, then storing the indices of both in a list, starting with the subject set and finally using this list as keys for a dictionary with values in the range to $n$. In both edge cases, this results in an adjacency matrix with a rank lower than $n$. A similar approach, with fewer edge cases to consider, is used to convert the tensor matrices back to triples.

### 4.2 RGVAE

The principle of a graph VAE has been explained in section 3.2.4, which also covers the foundation of our model, the RGVAE. Therefore we focus on the implementation as well as parameter and hyperparameter choice. Since this work is meant to be a proof of concept rather than aimed at outperforming the state of the art, our model is kept as simple as possible and only as complex as necessary. Our approach is modular for both experiment pipeline and model, meaning independence between sequential modules and compatibility with parallel modules. For the encoder we implemented two variations, a fully connected and a convolutional, while for the decoder we opted for a single fully connected network.

#### 4.2.1 Initialization

The RGVAE is initialized with a set of hyperparameters, which define the input shape. Table 1 shows a complete list of those parameters and their default values. It is left to mention that we use the Xavier uniform initialization with a gain of 0.01 to initialize the parameters.
| HYERP. | DEFAULT | DESCRIPTION |
|--------|---------|-------------|
| \(n\)  | 2       | Number of nodes |
| \(d_e\) | -       | Total number of entities |
| \(d_r\) | -       | Total number of relations |
| \(d_z\) | 100     | Latent space dimension |
| \(d_h\) | 512     | Hidden dimension |
| dropout | 0.2     | Dropout |
| \(\beta\) | 1       | \(\beta\) value for regularization |
| perminv | True    | Permutation invariant loss function |
| clipgrad | True    | Learning w/ gradient clipping |
| encoder | MLP     | Choice of encoder architecture |

Table 1: The initial hyperparameters of the RGVAE with default value and description.

4.2.2 Encoder

The proof-of-concept encoder is a MLP as described in section 3.2.2, which takes the flattened concatenated threefold graph \(x = G(A, E, F)\) as batch input. We use the initial parameters to calculate the input

\[
d_{input} = n^2 + n^2 \ast d_e + n \ast d_e
\]  

The main encoder architecture is a 3 layer fully connected network, with both layers using ReLU as activation function. The choice for two hidden layers is based on the huge difference between \(d_{input}\) and \(d_z\).

The first layer has a dimension of \(2 \ast d_h\) and the option to use dropout, which by default is set to 0.2. The second (hidden) layer has the dimension \(d_h\) which is by default set to 1024. After the second ReLU activation, the encoder linearly transforms the hidden state to an output vector of \(2 \ast d_z\). This vector is split and makes the mean and log-variance of size \(d_z\) for the reparametrization trick. Sampling \(\epsilon\) from an autonomous module, we get the latent representation \(z\).

The second option for our RGVAE encoder is a GCN as described in section 3.2.3. We adopt the architecture from [6] namely two layers of graph convolutions with dropout in between. To match the encoder output to the base model, we then add a flattening and a final linear transformation layer. To substitute the feature matrix used in Kipf’s work, we reduce the edge attribute matrix \(E\) by one dimension and concatenate it with \(F\) resulting in \(X_{GCN} \in \mathbb{R}^{n \times (d_e + nd_r)}\). The forward pass of the adjacency matrix \(A\) and \(X_{GCN}\) through the first GCN layer with a hidden dimension of \(d_h\) and ReLU as activation function is followed by a dropout layer. It should be mentioned that dropout is only applied during learning, not on evaluation. The second GCN layer takes the output from the previous layer, the two dimensional hidden state and again \(A\) as input. Now, instead of having the GCN predict on a number of classes, we use it to output a logits vector of dimension \(2d_z\). Therefore we pass the GCN output through a flattening and a linear transformation layer. Similar to the above described encoder we use the reparametrization trick to output the latent reparametrization \(z\). Table 2 shows the two encoder architectures side by side.

| MLP | Graph Conv. |
|-----------------|-----------------|
| Flatten\(d_{input}\) | Concatenate |
| Linear\(d_{input}, 2 \ast d_h\) | Convolution(A, X) |
| ReLu() | ReLU() |
| Dropout(0.2) | Dropout(0.2) |
| Linear\(2 \ast d_h, d_h\) | Convolution\(H^{(1)}, X\) |
| ReLu() | Flatten |
| Linear\(d_h, 2 \ast d_z\) | Linear\(d_H^{(1)}, 2 \ast d_z\) |

Table 2: Comparison of the two variants for the encoder of the RGVAE.

4.2.3 Decoder

The RGVAE decoder is a MLP with architecture and dimensions similar but in inverse order to the encoder MLP described in table 2. Since we are decoding the latent space, the input dimension is \(d_z\) and the output dimension is \(d_{input}\) as calculated in equation 16. The flat logits output tensor is split threefold and reshaped to the original input shape of \(G(A, E, F)\).

To sample from the generated graph we apply the Sigmoid activation function to the logits prediction for the adjacency matrix and use the normalized output as weights for binomial distributions, from which we can sample the discrete \(\tilde{A}\). For \(\tilde{E}\) and \(\tilde{F}\) we take the argmax on the last dimension of both matrices. Each node
and edge can have only one attribute, referring to its index in $E$ and $V$, thus only the highest predicted value is relevant. The generated sample is a discrete graph $\tilde{G}(\tilde{A}, \tilde{E}, \tilde{F})$.

### 4.2.4 Limitations

The main limitation of the RGVAE is the quadratic increase of model parameters with the increase of nodes per input graph $O(n^2)$. The number of parameters to train is directly linked with the GPU memory requirements. Even more computationally expensive is the use of permutation invariant graph matching, with a complexity of $O(n^3)$. This sets an empirical limitation for the model to small graphs with $2 < n < 40$.

### 4.3 RGVAE learning

In this section we present our implementation of how to fit the model to the data. Learning a model on data is a mostly standardized procedure, which includes training and evaluation per epoch. During training, the model forward passes the data, computes the loss, then does a backwards pass and updates its parameters. During evaluation, it is presented a split of the dataset unseen during training. Only the forward pass is done and the loss tracked during evaluation. Up to this point the RGVAE does not differ from the vanilla VAE training. Special is the graph matching function which is applied to the predicted graph and the loss function which takes into account the permutation. Thus, we look deeper into graph matching and derive RGVAE loss. The training and all experiments are performed on the GPU cluster LISA of the supercomputer SURFsara on the Dutch national e-infrastructure with the support of SURF Cooperative. Each GPU node is powered by a Nvidia titan RX 25GB. We log our experiments and results using Weights & Biases, a cloud-based experiment tracking tool.

#### 4.3.1 Max pooling graph matching

While the pseudocode presented in [27] is simple and straightforward, it proves complicated to implement this in algorithms for batches and thus, without looping over the indices. Yet, our batch implementation solves these challenges and is more efficient than the direct implementation, which we use for validating our results. Given the target graph $G$ and the predicted graph $\tilde{G}$, the algorithm can be divided in three steps, calculating the five dimensional affinity matrix (the first being the batch dimension), max-pooling the continuous similarity matrix $X^*$ and discretizing $X^*$ to our final permutation matrix $X$.

We use equation 9 for the first step but instead of adding the two terms to a single output, we return $S$ twofold as $S_r$, five dimensional holding the information of edge affinity and $S_e$, three dimensional with the affinity information of the nodes. In a preprocessing step we zero out the diagonal of $A$, $\tilde{A}$ and for $E$ and $\tilde{E}$ the diagonal of the second and third dimension, to compile with the constrain $[i \neq j \land a \neq b]$ of the first term. For the second term we only take into account the diagonal of $\tilde{A}$ to compile with the constrain $[i = j \land a = b]$. Pseudocode [1] shows the implementation, here $\text{diag}()$ stands for a vector with only the diagonal entries. For the dot product of $E$ and $\tilde{E}$ over the last dimension we implement our own version of $\text{torch.matmul}()$ to cope with higher dimensions. The operator $\odot$ denotes element-wise matrix multiplication.

#### Algorithm 1 Batch implementation for the affinity between two graphs

| Input: $G(A, E, F)$ and $\tilde{G}(\tilde{A}, \tilde{E}, \tilde{F})$ |
|---------------------------------------------------------------|
| **First term:** $[i \neq j \land a \neq b]$ |
| 1: $E_{\text{term1}} = E^T \tilde{E}$ ○ Dot product over the last dimension |
| 2: $A_{\text{term1}} = A$, unsqueeze($-1)^T(\tilde{A} \odot (\tilde{A}^T))$, unsqueeze($-1)$ ○ Dot product over the last (empty) dimension |
| 3: $S_r = E_{\text{term1}} \odot A_{\text{term1}}$ |
| **Second term:** $[i = j \land a = b]$ |
| 4: $A_{\text{term2}} = \text{ones_like(diag}(\tilde{A}))^T \text{diag}(\tilde{A})$ ○ Dot product over the last dimension |
| 5: $F_{\text{term2}} = F^T \tilde{F}$ |
| 6: $S_e = F_{\text{term2}} \odot A_{\text{term2}}$ |
| 7: return $(S_r, S_e)$ |

The next step is the graph matching algorithm is the max-pool loop presented in [27]. We initialize the similarity matrix as ones $X^* \in 1^{bs \times n \times n}$ with $bs$ denoting the batch size. For a certain number of iterations, Cho et al. proposes 40 but the number should be adjusted to the number of nodes in the graph, we multiply $X^*$ with a reduced version of $S$ and use its Frobenius norm as normalizer. The algorithm [2] shows our implementation for batches.

To the best of our knowledge, this is the first time this algorithm is implemented in batch style. Thus, we would like to believe that laying out the implementation in detail contribute to the academic value of this thesis.
Algorithm 2 Max-pool graph matching for batches

Input: \((S_i, S_j)\)

1. Init \(X^* \in 1^{b \times n \times n}\)
2. for iteration = 1, 2, \ldots do
3. \(S_{\text{max}} = \text{sum}(\text{max}(S_i \odot X^*, \text{unsqueeze}([1, 1]))) \odot \text{Sum and max over the last dimension. Unsqueeze two times on the second dimension}
4. \(X^* = X^* \odot S + S_{\text{max}}\)
5. \(X^* = X^*/\text{frobenius_norm}(X^*)\)
6. end for
7. return \(X^*\)

The last step in the graph matching pipeline is the discretization of \(X^*\). We adopt Simonovsky’s \([4]\) approach which for this purpose uses the Hungarian algorithm as presented in section 3.3.3. To our disappointment and resulting in a bottleneck, no batch nor tensor implementation of the named algorithm has been published so far. Thus, the discrete permutation matrix \(X\) is obtained by iterating over the batch dimension\(^3\). If no permutation is needed, \(X\) is the identity matrix of \(A\).

4.3.2 Loss function

The RGVAE uses the ELBO loss from equation \([1]\) consisting of two terms, the regularization loss and the reconstruction loss. We present our implementation of both loss terms with graph matching and an alternative loss without graph matching for comparative evaluation of our results.

We implement \(\log p(A' | z)\) from equation \([13]\) with the second normalizing constant as \(1/k \ast K\) since we allow self-loops. The permutation matrix \(X\) is applied to the target adjacency, resulting in \(A'\). For \(\log p(E' | z)\) and \(\log p(F | z)\) the permutation is applied to the prediction, which in the case of \(E'\) requires our own implementation of matrix multiplication of \(d > 2\). Taking into account self-loops we change the normalization constant of \(\log p(E' | z)\) to \(1/\|A\|_1\). It is left to mention that when implementing \(\sum_{i,j} \log E'_{i,j} = E'_{i,j}\) in matrix multiplication style, we have to account for the zero values before taking the logarithm. We implement \(\text{torch.sum} (\text{torch.sum} (\text{torch.log} (\text{no_zero}(E * E'))), \text{dim}=1)\) with \(\text{no_zero()}\) being a function which replaces 0 values with 1. This implementation of the loss function can be backpropagated with exception of the graph matching part, where the \texttt{numpy} implementation of the Hungarian algorithm prevents backpropagation.

The regularization loss is given by the KL divergence between the approximated posterior \(q_{\phi}(z | x)\) and the Gaussian prior \(p(z)\). The only modification we make to the original loss, is adding a \(\beta\) parameter which in values \(100 < \beta < 500\) has shown great results in factorizing the latent space \([25]\). By setting \(\beta = 1\) we return to the original loss function. This hyperparameter is explored in the experiments.

Alternatively and as ground truth we implement the VAE loss from equation \([4]\) for graphs without graph matching. We use binary cross entropy (BCE) as reconstruction loss of the adjacency, and categorical cross entropy (CE) for the attribute matrices. The regularization loss is similar to the above presented and also includes the hyperparameter \(\beta\). The equation for the ELBO with \(\sigma()\) indicating Sigmoid activation is

\[
\mathcal{L}(\phi, \theta : G) = \text{BCE}(A, \sigma(\tilde{A})) + \text{CC}(E, \tilde{E}) + \text{CC}(F, \tilde{F}) - D_{KL}(q_{\phi}(z | G) \parallel p_{\theta}(z))
\]

We train the model on the negative ELBO. Further we use \textit{Ranger} presented in section 3.4 as optimizer combining three learning optimization methods. Out of the various optimization parameters, we achieved good performance with the default values and only adjusted the learning rate and the number of lookahead steps.

Missing a publication to cite, we refer to the source code of \textit{Ranger}\(^4\).

4.4 Link prediction and Metrics

Our first experiment is link prediction, which we give an insight in the models capacity to reproduce the data. It is intended as proof of concept rather than an attempt to set the state of the art. The results let us draw conclusions on the impact and function of different parameters. Besides the final link prediction experiment, we perform link prediction on a randomly drawn small subset of the test set during training. This gives us a broader view on the models performance, which otherwise would only be evaluated by the ELBO loss.

Link prediction on multi-relational KGs is the task of completing unobserved triples, based on the information gained from the trainings data. To evaluate a model on this task, the most common method is entity ranking in the form of triple completion of unseen triples from the test set. Given a KG \(G(E, V)\) we want

\(^3\text{We convert \(X^*\) to \texttt{numpy.array()}, create the cost matrix \(X_{\text{cost}} = 1 - X^*\) and use the \texttt{Scipy} package \([38]\) \texttt{scipy.optimize.linear_sum_assignment} to iteratively calculate the permutation matrix \(X\).}

\(^4\text{https://github.com/lessw2020/Ranger-Deep-Learning-Optimizer}\)
our model find the right entity out of $E$ which completes the unseen triple $(s, r, ?)$ or $(?, r, o)$ for heads or tail prediction. Thus, the model scores the triple for all possible combination with the entities from $E$. The rank of the true triple, in descending order, defines the performance of the model \[39\].

In the preprocessing step we create a dictionary with all occurring combinations for all possible triples with missing head or tail. These dictionaries we use to filter out real triples from the scoring. Unfiltered scores are referred to as raw scores. Per link prediction run, the model has to score the number of triples in the test set times the $d_e$ the number of entities in $E$ times two for head and tail, which mostly results in a number much larger than the size of the actual dataset.

Finally, the metrics for link prediction are the mean reciprocal rank (MRR) of the score for the true triple and the average HITS@$k$ with $k \in \{1, 3, 10\}$. We denote $K_{\text{test}}$ the unseen test set and $|K_{\text{test}}|$ the number of triples in the test set. The operator $\text{rank}()$ returns in descending order the by the model scored position of the true triple and the average HITS@$k$. The Iverson brackets $[\text{rank}(s | r, o) \leq k]$ return 1 if the scored rank is equal or lower than $k$, else 0.

\[
\text{MRR} = \frac{1}{2|K_{\text{test}}|} \sum_{(s, r, o) \in K_{\text{test}}} \left( \frac{1}{\text{rank}(s | r, o)} + \frac{1}{\text{rank}(o | s, r)} \right) \\
\text{Hits@ } k = \frac{1}{2|K_{\text{test}}|} \sum_{(s, r, o) \in K_{\text{test}}} ([\text{rank}(s | r, o) \leq k] + [\text{rank}(o | s, r) \leq k]) 
\]

\[18\]

4.5 Variational DistMult

In the context of the link prediction experiment, we implement a control model. This allows a better interpretation of the results and understanding of the impact of the isolated model parameters. From the wide range of embedding-based models, DistMult reports both good results as well as an efficient architecture. Its bilinear property aligns with the permutation invariance of the RGVAE. Based the original model we implement a variational version of DistMult to isolate the effect of variational inference on multi-relational link prediction.

The DistMult encoder has a linear embedding layer for both sets of entities and relations. The embedded or latent representation is passed through the bilinear scoring function, which is equation$^2$. During training the model scores the triples in the training set among a number of corrupted triples. Loss function and optimizer are tunable hyperparameters. The optimal hyperparameter settings for the FB15k-237 dataset are published in Ruffellini et al. \[39\] and are used for the embedding-based link prediction experiment.

The model implementation is adopted from Peter Bloem’s work.\[4\] Additionally a variational module is implemented, which uses the embedding vector representation as mean and logvariance for a latent distribution from where the latent representation is sampled using the reparametrization trick. When this method is not selected, the stochastic coefficient $\epsilon$ is set to 1 rather than a random sample from the standard normal distribution. Lastly we implement the ELBO loss, which adds a regularization term, including $\beta$ parameters to the BCE loss, thus we can only use the variational DistMult (VDistMult) in combination with BCE loss. The scoring function of the VDistMult stays identical to the original. We use the parameter settings for both the original and the variational DistMult.

5 Experiments & Results

This section presents the experiments and results aimed at evaluating our proposed graph generative model. First we run a grid-search on the hyperparameter space to find the optimal configuration of the RGVAE. We use the ELBO and MRR as evaluation metric. The best configurations are used to perform link prediction. Here we compare the model performance with MLP versus GCN encoder. We use the VDistMult as control model for link prediction. Finally we run two proof-of-concept experiments. The first generating triples and filter on a entity class constraining relation, thus we get an insight of how much percent of the generated triples are valid. For the experiments we use two multi-relational KG datasets.

5.1 Data

For this sake of comparison with state of the art results, we chose the two dataset used in this field of KG link prediction, FB15k237 and WN18rr.

FB15K-237 is a successor of the FB15K dataset, first introduced by \[14\], which suffered of major test leakage, meaning that triples from test set could be inferred by inverting triples from the train set. In FB15K-237, introduced in \[40\], these triples were removed. The data was scraped from Freebase, while only

\[https://github.com/pbloem/emb\]
the most frequent entities and relations were considered. The huge open-world KG Freebase \cite{41}, which before its discontinuation had around 1.2 billion triples and 80 million entities, was structured by assigning types and classes to entities and type constraints to relations. Thus, a triple can only be formed if the relation constraint matches the entity’s type. Freebase was free for everyone to access and expand. This led to inconsistencies, duplicates and highly inconsistent notation, which might have been the reason for its discontinuation. Data dumps of the latest version are still available. Since the launch date of Freebase, advanced systems for storage and query of large-scale maintenance KGs have evolved \cite{42}.

WN18RR, a dataset of synonyms and hypernyms, is a successor of yet another dataset WN18 introduced by \cite{14}. Similar to the above the original dataset suffered from test leakage, thus, an updated version without reciprocal triples was introduced by \cite{43}. This dataset is characterized by its few relations and large corpus of entities. In contrary to FB15K-237, its triples require a higher level of grammatical understanding and prior knowledge of the entities.

| Dataset   | Entities | Relations | Triples  |
|-----------|----------|-----------|----------|
| FB15K-237 | 14,951   | 237       | 310,116  |
| WN18RR    | 40,943   | 11        | 93,003   |

Table 3: Statistics of the FB15K-237 \cite{40} and WN18RR \cite{43} datasets.

5.2 Hyperparameter Tuning

In this section the hyperparameters for the VanillaRGVAE are tuned. The VanillaRGVAE has a MLP encoder, a permutation invariant loss function and a Standard Gaussian latent space prior. We run a grid search for the three hyperparameter $\beta$, $d_z$ and $d_h$ for a set of contrastive values. To reduce the computational expenses we train each model for 60 epochs and evaluate link prediction on a subset of 50 triples.

We set the learning rate to $3 \times 10^{-5}$ and the maximum batch size fitting on the GPU memory. For $d_h$ we did not see any significant changes for higher values, thus we choose a lower number to reduce the total model parameters. The remaining hyperparameter did influence and the optimal setting vary for each dataset, table \ref{table:hyperparams} shows the results of our hyperparameter tuning.

For the hyperparameter tuning of $\beta \in [0, 1, 10, 100]$ we chose significant values. With $\beta = 0$ we do not constrain our model on the Gaussian prior, thus the latent distribution can take the form of any distribution. This reduces the influence of the variational module and the model becomes closer to an autoencoder. For $\beta = 1$ we get our base model, and for $\beta \in [10, 100]$ the $\beta$-VAE version.

![Validation ELBO for the different $\beta$ values and for both datasets.](image)

Figure 7: Validation ELBO for the different $\beta$ values and for both datasets. We notice two interesting outcomes.

- For $\beta = 0$ converges further than the rest.
- The remaining values behave identical with $\beta = 100$ performing slightly better.

Since setting $\beta = 0$ yields a VAE with a latent space which is conditionally independent from the data, this setting is not considered valuable. Thus, we chose $\beta = 100$ as default for the following experiments. This also compares with the $\beta$ values proposed by Higgins in \cite{25} to achieve a factorization of the latent space.
Especially on the FB15k-237 dataset the $\beta = 0$ configuration converges to a much lower ELBO. Thus, we have the trained models perform link-prediction on a 1% subset of the validation set. Figure 8 indicates an inverse correlation between the ELBO and the MRR score.

![Figure 8: MRR scores during training for different $\beta$ values on 1% of the validation set.](image)

Experiments on the impact of $d_z$ on the ELBO show little improvement for $10 < d_z < 100$ and from $100 < d_z < 1000$ insignificant to no improvement. Thus, we chose $d_z = 100$ as default for our experiments.

Lastly, we evaluate the models hidden dimensions $d_h$ and its influence on the ELBO and the (subset)MRR. We compare between $d_h \in [256, 512, 1024, 2048]$, while the lowest configuration performs slightly worse on the ELBO, there is no significant difference between the remaining three configurations. Considering the models parameter count we chose the $d_h = 512$ as default.

### 5.3 Link Prediction

We now get to the most extensive experiment of this thesis. The results of this experiment show, whether the RGVAE is suitable for link prediction. Specifically, if it is able to grasp the underlying semantics of the KG data at significantly better than the baseline of random predictions. For this experiment the RGVAE is trained until convergence of the ELBO. The setup for the link prediction experiment consists of two iterations for completing each head and tail entity. For each triple in the test set, the RGVAE scores all possible combinations with all entities occurring in the dataset. Since the ranking is in increasing order, we use the negative ELBO to score each triple. This allows the triple with the lowest ELBO to rank the highest. For all test triples and both iterations the scores and the index of the true triple are stored. In a post processing step the raw results are filtered and the scores for real triples, which are not the target are excluded. Further the rank of the correct triple is determined by counting all scores which are higher than the target. For scoring ties with the correct triple, the rank is settled halfway. The rank is determined for every test triple and for missing head and tail entity. These ranks are used to calculate the MRR and Hits@k results.

In this experiment, we compare the RGVAE with MLP and GCN encoder. Further we investigate the influence of the variational inference by comparing the variational and original versions of DistMult on link prediction.

#### 5.3.1 RGVAE

At this point we compare the convolutional encoder variation of our model, which we denote as cRGVAE. The experiments reveal if the convolutional architecture holds an advantage compared to the simple MLP baseline. Further a randomly initiated and untrained RGVAE is used as control model.

Due to its sparse graph computation, the RGVAE takes about 7 days to evaluate link prediction on the full test set and even 3 days when prediction tasks run parallel on a node with 4 GPUs. Since the exemplary link prediction during experimenting with different hyperparameter already gave us an idea of the mediocre performance of our model, we chose to spare computation time and power by running link prediction on a randomly drawn one-third of the complete test set. Each run is repeated three times using a different random seed.

The results are visualized in figure 9. We chose a visualization over a table, to emphasize the observed differences between encoders and datasets. These results should not be directly compared to other experiments since they were performed, as mentioned above, one-third of the test set. Note that the figures are scales to a range $[0, 0.1]$ while all metrics have a maximum of 1. The dotted line represents the baseline score of an untrained RGVAE with MLP encoder.
Comparing the RGVAE MRR score 0.08 to the DistMult score 0.3 on the FB15K-237 dataset our model does not perform competitively on link prediction tasks. The DistMult scores are reproduced within the scope of the embedding-based link prediction experiment, solely on the FB15K-237 dataset, using the by Ruffinelli et al. published hyperparameter settings. For the WN18RR dataset the MRR score from [39] is referenced. The DistMult scores a MRR 0.45 compared to the RGVAE score 0.015 on WN18RR. The baseline MRR score of the untrained RGVAE is 0.006 for FB15K-237 and 0.0015 for WN18RR.

In this experiment, graph convolutions do not yield an advantage over the MLP encoder. In fact, the RGVAE with GCN encoder even scores slightly worse. All versions of the RGVAE score better than the baseline, proving that the RGVAE learns a representation the data.

The model scores about three times better on the FB15K-237 dataset than on WN18RR. FB15k-237 is a richer dataset with more triples and a more balanced ratio of entities to relations. WN18RR operates on only 18 relations, what makes the relation most crucial when completing a triple. The architecture of the RGVAE puts twice the emphasis entities, described by the adjacency and the node feature matrices, while the relation is only represented by the overly sparse edge attribute matrix. Thus, we could conclude that our model learns to predict based on the hidden types and topics of the entities. All possible conclusions for this are discussed in chapter 6.5. Relevant for this section is solely, that we chose the FB15K-237 dataset to investigate further, how well the RGVAE’s grasps the underlying entity types and triple topics.

![Figure 9: Link prediction scores of the RGVAE. Dotted line represents MRR baseline of an untrained model.](image)

5.3.2 Impact of Variational Inference and Gaussian prior

In order to explain the poor performance of the RGVAE on the task of link prediction, we investigate the impact of the variational inference. Since the RGVAE with relaxed latent space, meaning less variance, indicated higher scores than the version with Gaussian prior, we examine the two variants by means of embedding models. The original DistMult model with optimized parameter serves as control model, while we compare it to the VDistMult, described in section 4.5, learning the full ELBO versus learning only on the reconstruction loss. By not including the regularization term in the loss the model is no longer bound to the Gaussian prior, which results in a relaxation of the latent space.

We train the three models for 300 epochs solely on the FB15k-237 dataset and evaluate MRR, Hits@1, Hits@3 and Hits@10. Table 4 shows the mean scores with $\mu \pm \sigma^2$ of three runs per model. Note that the exponent on $\sigma^2$ holds for the whole term. We can clearly see that both variational versions of the DistMult perform significantly worse than the original model. Learning on the full ELBO or only the reconstruction loss does not seem to influence the scores in this setting. This indicates, that the models performance on link prediction suffers from using variational inference.

In the last row we show the results of the RGVAE with relaxed latent space. This model was trained with $\beta = 0$ thus not constraining the latent space on a Gaussian prior. The model outperforms the versions with hyperparameter choice $\beta > 0$ and scores the closest to the DistMult model. The impact of $\beta$ shows in the regularization, which we tracked separately. The maximum values of $D_{KL}$ during the experiment are

$$D_{reg} = \beta D_{KL} (q_\phi(z | G) \parallel p_\theta(z))$$

$$\max_{\beta=0} D_{KL} = 3506$$

$$\max_{\beta=100} D_{KL} = 0.0154$$
While the results of the relaxed RGVAE might seem promising, DistMult is a much simpler and faster link predictor, what leads us to the conclusion that the RGVAE in none of the chosen setup scores competitive with embedding-based models on this task. Note that due to the high computation cost of the RGVAE we only run the experiment once on the full dataset, thus no variance is reported.

| Model             | MRR      | Hits@1    | Hits@3    | Hits@3    |
|-------------------|----------|-----------|-----------|-----------|
| DistMult          | 0.2854 ± 0.0025 | 0.2 ± 0.001 | 0.3149 ± 0.0038 | 0.4512 ± 0.0053 |
| VDistMult         | (0.517 ± 0.0197)e^{-3} | (0.2442 ± 0.1994)e^{-4} | (0.8145 ± 0.3049)e^{-4} | (0.399 ± 0.0576)e^{-3} |
| VDistMult w/ ELBO | (0.6397 ± 0.0357)e^{-3} | (0.57 ± 0.3046)e^{-4} | (0.1547 ± 0.1023)e^{-3} | (0.6351 ± 0.1992)e^{-4} |
| RGVAE w/o ELBO    | 0.1412   | 0.0981    | 0.1494    | 0.2275    |

Table 4: Link prediction scores of DistMult and RGVAE versions on the FB15k-237 dataset.

### 5.4 Impact of permutation

Furthermore we examine the influence of the permutation invariant loss function described in 4.3.2. During training and subset link prediction, no significant difference was observed between the RGVAE with versus without matching target and prediction graph. Yet, two observations draw our attention, namely:

- The amount of nodes permuted per batch converges during training from 100% to exactly 80
- The RGVAE with permutation invariant loss function learns to predict many variations of adjacency matrix while the standard model predicts similar to the target.

The first point indicates that the model learns a set of adjacency matrices, which can be permuted to match the target while optimizations the loss. Note that the generated matrix representation of the triples either has only one edge on the right upper index $A_{0,n}$ or, in the rare case of self-loops in $A_{0,0}$. The number of nodes per graph for these observations is set to $n = 2$. Curiosity remains why the model converges to steadily permute $\frac{3}{5}$ of the prediction. Secondly, we see that even when converged, the model predicts variations od the adjacency matrix very different to the target. The most common is a single edge on $A_{n,0}$ and on $A_{n,n}$. Less common and with lack of explanation are the predictions of an empty, or multi edge adjacency matrix. In contrast to this and as expected, the RGVAE with the standard loss function learns to solely predict edges on $A_{0,0}$. Finally we analyze the impact of permutation invariance on the experiment of generating valid triples section 5.6.

### 5.5 Interpolate Latent Space

Inspired by the popular results of Higgins, who disentangled the latent space in a way that each dimension mapped to a specific facial feature from data of the FACES dataset [46]. The VAE generates faces controlling feature such as age, gender and emotions by manipulating single latent dimensions [25]. We run this experiment with the RGVAE on the FB15k-237 dataset, using two different interpolation methods. The latent dimension for this experiment is set to $d_z = 10$ in order to analyze each dimension separately and the interpolation is linear with a step count of 10.

The first experiment is linear interpolating between two triples. Therefor two valid triples from the train set are encoded into their latent representation. We chose the two semantically related triples to analyze if the linear movement in latent space correlates with an obvious semantic feature.
RGVAE PERMUTATION

[France] [/base/petbreeds/city_with_dogs/top_breeds] [Imperial Japanese Army]
[Cree Summer] [/tv/tv_program/program_creator] [David Chase]
[Michelangelo] [/people/deceased_person/place_of_death] [Rome]

Table 5: Latent space interpolation between two triples in 10 steps.

For the second experiment, we interpolate each latent dimension isolated in a 95% confidence interval of the Standard Gaussian distribution. Starting with the encoded representation of the Obama triple, we incrementally add $z_i = -1.96 + j \times s$ with $s = \frac{1.96}{n_s}$ for the number of steps $n_s = 10$. Due to the size of the tables representing the interpolations and the low value they add to the presentation of this work, the result of this experiments can be found in the annex B.2.

5.6 Generator Validation

On closed-world schema-based KGs the approach for testing the validity of a new triple is to add it to the existing KG and run a ontology reasoner on it. A inconsistency in the KG appear as null-Class, but only if an axiom is violated. This approach works only for fully constrained KGs and is not scalable, since the reasoner recursively checks every triple for every axiom. Thus, we present an alternative and improvised way to estimate the validity of generated triples.

The FB15K-237 is a subset from the FreeBase KG [41], thus, even thought they are not part of the dataset, 14541 entities have type properties in their original Freebase representation. Querying the last official Freebase dump, as proposed by Xie et al. [47], we get the types for each entity in the FB15K-237 dataset, With exception of 8 entities, which could not be found in the query. Our approach is to randomly generate triples, from signals randomly drawn from a Standard Gaussian distribution. Then to filter those triples on predicates which contain the type people, which is within the top 10 most common Freebase types. We differentiate between base-class types subclass types, both can contain the word people. The entity [/m/02mjmr Barack Obama] has between many others the type [/people/measured_person]. Here the base class is people and the subclass measured_person. We could filter directly on subclasses, but we chose to give our model more creative freedom and filter for people in the full set of types. Using this choice, the generated triples are scored on logic rather than facts. E.g. any person can hypothetically be a measured_person, understanding this implies semantical reasoning, while differentiating between which person is and is not a measured_person implies contextual knowledge. Thus, we use the base type people to validate triples. Furthermore the relations are inconsistent in their notation, partly not only having a head type constrain. Thus, we check only the head entity for the key type.

- From 14541 entities, 5283 contain the keyword people, or 36.332%.
- From 237 predicates, 25 contain the keyword people, or 10.549%.
- From 310116 triples, 47354 contain the predicate keyword people, or 15.269%.

Considering these facts, we calculate the marginal probability of guessing a head entity $s$ of type people, given a triple which contains the type people. Without prior knowledge $p(s_p)$ and $p(r_p)$ are conditionally independent. The probability is calculated as:

$$p(s_p | r_p) = p(s_p) = 0.3633$$  \hspace{1cm} (19)

For this experiment we generate triples until $10^5$ contain the key type. Those filtered triples are validated on the type of the head entity and compared to the full dataset for novelty. Here we again compare the performance between the RGVAE with the two different encoder architectures. Furthermore the models are trained both with regular and permutation invariant loss function. Lastly, the experiments are repeated for
sampling the latent signal from \( \mathcal{N}(0, 1) \) and for sampling from \( \mathcal{N}(0, 2) \). We average the accuracy of three runs of generating a valid triple and of this triple being unseen in the dataset. The results for valid triples are shown in figure 11. The dotted horizontal line indicates the random probability of generating a valid triple, calculated in equation 19 and \( \sigma_1^2 \) and \( \sigma_2^2 \) denote the variance for the different latent space distributions.

To our disappointment the model does not perform significantly better than the baseline of random predictions. Neither the choice loss function nor the doubled variance show a correlation with the accuracy. The only configuration standing out is the RGVAE with convolutional encoder, standard loss function and \( \sigma^2 = 2 \), scoring 4% higher than the baseline. From all valid generated triples 100 \( \pm \) 0.001% are new and unseen in the dataset. Coming back to ‘/m/02mjmr Barack Obama’, we filter the unseen triples for the first three appearances of this entity. These are displayed in table 6 for every variation of the RGVAE and in the same order as in figure 11. Note that in order to fit the triples to this thesis’ text width, some relations with object type specification have been cropped at ‘/’ as well as entity names with overlength.

The generated triples confirm the accuracy results. With exception of the triple

Barack Obama, /people/person/places.lived./people/place.lived/location, Casablanca

all remaining triples violate common sense logic. The RGVAE does not differentiate between the types gender, location, movie, person or medicine. It even goes so far to state that Obama was born in ‘Multiple sclerosis’. While this might sound funny it also clearly indicates, that our model did not learn the underlying semantics of this real world KG.

While investigating the model and the generated triple set, we notice two outcomes. Neither the augmentation of the variance nor the enabling of the permutation invariance has an impact on either of the model with two encoder versions. The regularization loss converges during training to zero, meaning that the model learns a latent representation of the dataset as nearly perfect Standard Gaussian distribution. Yet, even when sampling latent signal from the exact same distribution, we notice that each model repeatedly predicts combinations of a small subset of entities and relations, e.g. the RGVAE version which did not predict the Obama entity once in a total of 111583 valid triples. This also aligns with the interpolation results, where we observed a static relation for the full grid search of the latent space. If we look at the gradient and parameter values \( \phi \) and \( \theta \) of the MLP encoder and decoder, we see a much higher variance and gradients for \( \phi \). The decoder shows higher values and variance for a small subset of neighboring parameter, while the remaining parameters converge to a very similar and low value. This indicates that the encoder learns very well to represent each different triple as Standard Gaussian latent representation. The decoder MLP on the opposite seems to ignore most of this representation by assigning vanishing values to the connected parameters. Intuitively it seems that the decoder learns to interpret the part of the latent representation corresponding to the adjacency matrix and minimizes as well as stabilizes the loss of the edge and node attribute matrix by uniformly distributing their probability. This leads to the decoder reconstructing edge and node attribute randomly. Further, depending on the values of \( \theta \) when the model finishes learning, the decoder repeatedly predicts the same subset of entities and relation independent of the latent signal \( z \). If we look at the flattened representation of our input graph, we see that the part representing the adjacency matrix is way shorter and has a tractable mean of \( \frac{1}{4} \) while the mean for the edge and node attribute matrices are \( \frac{1}{1345} \) and \( \frac{1}{11599} \). The problem of the RGVAE decoder partly ignoring the latent input and potential solutions are discussed further in section 6.3.
5.7 Delta Correction

A solution to the observed phenomenon, which is known as decoder collapse, is the use of a $\delta$ parameter, presented by Razavi et al. in [48]. The regularization term is expanded by subtracting the $\delta$ parameter and by taking the absolute value. This forces the latent space into a truncated Gaussian distribution. The simple nature of this solution allows us to integrate it in the RGVAE setup and evaluate its impact on the previous experiments.

In a final experiment the RGVAE is trained in 4 different modes, with $\delta \in [0, 0.6]$ and using the standard versus the graph matching loss function. Since we do not expect this new parameter to have a significant impact on the link prediction results, the most informative and comparable experiments are latent space interpolation and the generation of valid triples.

Table 7 shows the results for $\delta = 0.6$ and graph matching. The model shows the same behavior as the for $\delta = 0$. The start and end triple do not match the encoded target triples and 9 out of 10 interpolation steps predict the same relation. A difference can be seen in the traversing of each latent dimension. Here minimal semantic coherence can be observed (Table in Appendix). The new parameter $\delta$ did not raise the ratio of valid generated triples above the baseline. A difference between standard, appended in the annex 9, and graph matching loss could also not be observed. The results of the full latent space interpolation for both variations of loss functions are appended in the annex B.2.
Table 7: RGVAE latent space interpolation with $\delta = 0.6$ and graph matching.

Finally, we plot the parameter values, both weight and bias to visualize the collapsing decoder. The values are logscaled and plotter per layer number of encoder and decoder as described in table 2. The parameters of the decoder behave equally for both $\delta$ values. All values of the 3rd layer conglomerate around zero revealing a bottleneck of the information stream in the decoder. This indicates that the problem of collapsing decoder was not solved by the addition of the new parameter. The parameter distribution for the RGVAE with standard loss are appended in the annex 14 and 15. The impact of the standard loss function is a more narrow distributed decoder. Since the GCN encoder has different layers, a direct parameter comparison to the MLP encoder is not possible, yet we observe the same parameter distribution for the decoder. Note in figure 12 the different scales of the value axis, which are $10^2$ larger for $\delta = 0.6$ than for $\delta = 0$.

![Parameter values per layer of the RGVAE encoder and decoder with $\delta = 0.6$.](image-url)
Discussion & Future Work

In this final chapter, we discuss our results from different experiments, emphasizing the comparison to Simonovsky’s work on molecule generation. The fact that we did not achieve similarly successful results does not hinder us from analyzing the responsible factors and causalities. Despite the nature of our results we are happy to propose possible solutions and while being of the opinion that asking the right question is more valuable than providing the correct answer, we question the value of these solutions for further research.

6.1 Experiment Take-away

The first relevant observation made is the lower ELBO score to which the RGVAE converges when using $\beta = 0$, meaning it is only trained on the reconstruction loss. We tested the assumption that the model performs better without regularization term. This leads to the assumption that the RGVAE performs better without prior constrain, yet a fully unconstrained VAE, defies its purpose when it comes to disentangling the latent space and sampling. The tuned RGVAE shows poor results on the task of link prediction, with slightly worse scores for the convolutional encoder. In order to find the isolate the causalities, we experiment further on link prediction with two variational version of DistMult learning on the full ELBO and only on the reconstruction loss. Both models scored equally bad compared to the original model, indicating that DistMult and embedding-based link predictors in general are not complex enough for variational inference. Following our prior assumption we added the unconstrained RGVAE to this comparison, which led to significantly higher scored than the all versions of the tuned RGVAE and the variation DistMult. Yet, the scores were not competitive with state of the art link predictors of with much lower parameter count, thus we do not recommend further researching in the direction of VAE as link predictor.

The graph matching loss function does not seem to have any impact on the link prediction results, yet we cannot draw a conclusion here since the poor performance is linked to a different causality which might impede the graph matching from adding value. To measure the percentage of permutation we adopt Simonovsky’s method of averaging the identity of permutation matrix $X$, which is invariant for permutation equivalent nodes. We see that during training the permutation rate converges to 60%.

Comparison of the two datasets, showed a better performance on the FB15K-237 dataset. While both datasets are real-world KGs, WN18RR is composed of synsets with vast discrepancy between the relation and entity count, resulting in a lower chance of generating a valid triple. Thus, we decided to continue the interpolation and graph generation experiments only on the FB15k-237 dataset.

For the last two experiments we trained the RGVAE with $d_z = 10$ for better disentaglement and analysis of each latent dimension. The interpolation experiment between two triples reveals that the RGVAE in all versions fails to reconstruct the start and end triple. We notice an influence of the graph matching loss, namely the model predicting constantly the same relation versus the standard loss, where the model predicts...
randomly from a subset of triples. Same can be said for the predicted entities, which form a small subset of what seems to be the most frequent occurrences in the dataset. Empirically we can say that the prediction has little prior conditioning on the latent signal.

These last experiment is designed and evaluated in a similar fashion as Simonovsky did to generate valid molecular graphs. We interpolate the full latent space for $-\sigma^2 < z < \sigma$ and $-2\sigma^2 < z < 2\sigma$. Based on our proposed validation criteria, the RGVAE generated the same rate of valid triples as random. We observe further independence between the decoder and the latent signal. This slightly compares with Simonovsky’s result where the generated valid triples had a variance of only 10%.

6.2 Research Answer

To answer the initial research question we first summarize the influence of the main hyperparameter. A convolutional encoder does not seem to contribute to the performance of the RGVAE. This could be caused by the simple implementation of using node and edge attributes both as feature matrix, or be due to the decoder issue, explained in section 6.3 that the prediction is independent of the encoding. Graph matching does have a noticeable impact. The model expresses more creative freedom when predicting the adjacency. Especially for experiments with larger subgraphs we recommend to keep investing in this method.

Variational inference is part of the VAE and allows disentanglement of the latent space, thus, even considering that the unconstrained model scored better, we recommend keeping the VAE structure. Instead of the full stochastic module, we assume the Standard Gaussian prior to be the reason behind the poor representation and recommend comparing alternative prior constrains. Overall we can answer the research question:

*The RGVAE failed in representation learning of real-world KG due to a decoder collapse.*

Yet, metaphorically, this is not the end of the book, but rather the beginning of a new chapter. In the remaining part of this discussion we analyze the underlying problem of the RGVAE and propose a variety of solutions.

6.3 Decoder Collapse

To introduce the main part of our conclusion we explain a phenomenon observed when training GANs. A GAN is yet another generative model which based on its potential to generate high resolution images, has drawn much attention in recent years. It consists of a generator, whose task it is to decode a latent signal to an image, and a discriminator, which given the fake image and the real data has to distinguish between them. Training of GANs is unstable and one of the major problem which occur is *mode collapse*. The generator learns to fool the discriminator by generating only a single mode with high precision such that the discriminator classifies it as real image [49].

While VAEs cannot suffer from mode collapse, because they backpropagate over the predicted distributions of all modes, it has a related phenomenon. *Decoder collapse* occurs when the decoder has enough capacity to choose not to consider the latent signal and instead stores the information to reconstruct the data’s distribution partly or solely in its parameters. This means that the generated data is not conditioned on the latent signal anymore. The cause of this problem is found in the regularization term $D_{KL}(q_\phi(z \mid x) \| p_\theta(z))$, more specifically in the constrain on a standard Gaussian prior. The multidimensional encoding of $d_z > 1$ the approximated posterior is a mixture of Gaussians, which can only match the multivariate Normal distribution, in the case of all $\mu_z = 0$ and $\Sigma_z = I$, what also implies that no information is encoded. Thus, the VAE has to decide if storing information in the latent vector is necessary to model the dataset distribution $p(x)$. If the penalty for altering the latent Normal distribution outweighs the benefits of the additional information towards the reconstruction loss, the VAE choses for *decoder collapse*.

Exactly this phenomenon can be observed in the triple interpolation and generation experiments. The RGVAE converges rapidly during training, minimizing the regularization loss close to zero and stabilizing the reconstruction loss by learning or matching the adjacency and predicting the most frequent node and edge attributes. This falsely indicates that the model has learned to reproduce the data. Yet, when sampling using only the decoder, the generated triples repeat combinations of a subset of entities and relations which are more frequent in the dataset. During interpolation of the latent space, the model predicts steadily the same relation. Subject and object seem to a certain degree be conditioned on the latent signal, altering while traversing the latent space, yet no clear disentanglement is observed. The final observation on the node attributes is, that the model shows a preference to predicts people’s name and film’s name entities as subject and location or music genre entities as object.

An interesting question, when observing the RGVAE generate all triples with the same single relation is, if the reason for this behavior can still be attributed to decoder collapse, as it rather collapses only on one mode?
A possible explanation is that the decoder learns to ignore the latent signal only for the relation prediction, still updating the full decoder during training and thus predicting different values after every update. Yet, when done training and used solely as generator, the decoder’s parameters stay constant and predicting the relation uninformed by any latent signal, which results in the collapse on the same relation prediction.

The results of the last experiment are surprising. While the RGVAE performed significantly better than the untrained baseline on link prediction, the ratio of valid generated triples does not differ from random sampling for any of the different model variations. Further we again notice the predominance of the more frequent data entities and relations.

Why does the RGVAE perform better on link prediction than on generating valid triples, if both tasks require representational understanding of the data?

Crucial in this comparison could be that link prediction is scored by the ELBO, including both encoder and decoder, while triples are generated only by the decoder.

Projecting this finding on the link prediction experiment, we question, why the model scored better than the random baseline. Possible reasons are for once, that the encoder learned to encode the dataset close to a Normal distribution, but when encountering unseen triples, the mean and variance of the encoding vary. A second reason is, that every triple in the test set is corrupted in all possible combinations, including the less frequent entities. The collapsed decoder subsequently keeps generating frequent triples, thus the reconstruction loss between a less frequent combination of entities and the collapsed prediction is be higher than for a frequent combination. Therefore the model learns score the real triple is higher but for the wrong reason. The RGVAE with \( \beta = 0 \) and therefore unconstrained on the Standard Gaussian prior shows the best results between the different model settings. We can assume that the model learns a latent representation for subject and object. Despite this improvement, it is probable that the decoder still collapses on the reconstruction of the relation index, which explains the remaining scoring gap to the much simpler DistMult model.

### 6.4 VAE surgery

Similar problems in different fields have been encountered for generative VAE applications. While the author of this thesis wishes to have drawn these parallels earlier, all the proposed solutions imply a significant modification of the VanillaVAE, thus would not align with this work’s research question. We present three approaches from different literature which tackle the VAE mode collapse in the filed of image and voice generation.

In a publication of Adobe Research and Google Brain, Hoffman et al. propose an elegant modification of the variational evidence lower bound \[50\]. More specifically they change the VanillaVAE’s regularization term, where instead of imposing a Standard Gaussian prior on the full latent distribution, the prior is imposed on the mixture of all single latent dimension, giving space for more expressive probability distributions such as truncated Gaussians. Further, a index-code mutual information term is added, intended to maximize the mutual information between every index of the observation and \( z \). While the reconstruction term enforces to encode every feature of \( x \) in a corresponding latent dimension, the information term opposes this by maximizing the mutual information between all \( x_i \) and \( z_i \). The information term compares compact to the reconstruction loss, yet it is enough to prevent decoder collapse.

Kingma et al. turn towards a autoregressive solution in their publication \[51\]. Their VAE model generates images recursive per pixel, each conditioned on previous point, using both RNN and RCN as decoder. Their solution to the decoder collapse is a normalizing flow, which predicts the encoder posterior \( qφ(z \mid x) \). Besides not suffering from decoder collapse, this model can be set to discard irrelevant information in the data. As downside, the autoregressive nature causes slow image generation.

The last approach we present closes the circle to the introduction of this section. The paper Wasserstein Auto-Encoders \[52\] propose a combination between GAN and VAE. A model which uses the VAE’s encoder-decoder architecture but instead of the normal regularization term, the posterior distribution is learned and penalized by its Wasserstein distance to the data distribution. This is in fact a generalization of the adversarial loss of generator and discriminator. Since our task of generating triples would benefit from a precise prediction, such as GANs achieve on image generation, we question, if employing adversarial loss also on the reconstruction term would yield better generation results in this field than the exact index-wise loss?

Besides these three, numerous other approaches have been proposed. Worth mentioning because of their originality are the idea of adding skip-connections between latent space and the decoder’s hidden layer \[53\] as well as regularizing the amortized inference \[54\].

### 6.5 Future Work

There are many avenues to follow for future work. Obviously the first step is to fix the collapsing decoder. The suggested solutions should be compared before making a choice, since none has yet been shown to work on KG VAEs.
Once the decoder reaches the point of predicting the right graph based on its latent representation, the RGVAE should perform significantly better on both link prediction and generating valid triples. Comparing the RGVAE’s efficiency at both tasks, generating triples is efficient and fast whereas link prediction inefficient and therefore disproportionately slow. Optimizing this could yield a bigger challenge than improving the scores.

Besides the model we should also look at the data, and question the representation. The adjacency matrix can be inferred from the edge attribute matrix and thus, is obsolete. Since it does not contain additional information, does the model perform worse without it? Approaching this thought from a different angle, we could leave the adjacency as it is and represent the node and edge indices as continuous number, as it is done for color scales of images. This would greatly reduce the number of parameters, but certainly also imply new challenges. Important in this context is that the adjacency is represented in sparse format since this the experiments on triples is but a proof of concept for larger subgraphs.

The model’s mayor hurdle seems to be predicting the correct relation index. While interpolating the entire latent space, less than 10 different relations were generated. A further challenge is the varying count and position of the edges, in contrast to the number of nodes which is constant. Thus, we wonder if a setup of two VAEs, where the first one predicts the adjacency and node attributes and the second recurrent one, while conditioned on the predicted edges of the first VAE, predicts the edge attribute.

Finally and making use of the batch wise implementation of the max-pooling graph matching algorithm presented in this work, we suggest to explore a more efficient and intelligent graph matching approach. Would it be possible drastically reduce complexity by training a neural network to predict the permutation matrix based on target and prediction graph?

The author hopes that his questions spark creative ideas for further research on generative models for KGs and looks with excitement towards future findings.
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Annex

A Parameter distributions

Figure 14: Parameter values per layer of the RGVAE with standard loss and \( \delta = 0.6 \).

Figure 15: Parameter values per layer of the RGVAE with standard loss and \( \delta = 0 \).
B Interpolation

B.1 Interpolation between two

| RGVAE STANDARD                                                                 |
|--------------------------------------------------------------------------------|
| [[Jennifer Jason Leigh] [/film/film/release_date_s] [William Wyler]]           |
| [[Keyboard] [/award/award_category/nominees] [Jessica Biel]]                   |
| [[Die Another Day] [/language/human_language/countries_spoken_in] [Stockholm University]] |
| [[Mission: Impossible II] [/education/educational_institution] [Intel Corporation]] |
| [[Techno] [/organization/role/leaders] [Makeup Artist-GB]]                   |
| [[The Pianist] [/soccer/football_team/current_roster] [Female]]               |
| [[Paul Winchell] [/award/award_nominee] [Buddhism]]                          |
| [[California] [/business/business_operation/industry] [Marriage]]             |
| [[PFC Beroe Stara Zagora] [/people/person/profession] [National Society of Film Critics Award]] |
| [[Knight and Day] [/award/award_category/winners_winner] [Fox Searchlight Pictures]] |

Table 8: Latent space interpolation between two triples for RGVAE with standard loss.

| δ-RGVAE STANDARD                                                                 |
|--------------------------------------------------------------------------------|
| [[Willem Dafoe] [/music/instrument/instrumentalists] [Peritonitis]]             |
| [[Tuscaloosa] [/influence/influence_node/influenced_by] [Alex Rodriguez]]      |
| [[BAFTA Award for Best Sound] [/people/person/gender] [Fat Possum Records]]   |
| [["Conan O'Brien"] [/film/film/country] [World]]                             |
| [["They Shoot Horses Don't They?"] [/base/popstra/location/vacationers] [Buzz Aldrin]] |
| [[The Lady] [/award/award_winning_work/awards_won] [Faith Hill]]               |
| [[Laura] [/film/film/country] [Talking Heads]]                                |
| [[Peter Morgan] [/film/film/release_date_s] [John Amos]]                      |
| [[The Butterfly Effect] [/award/award_nominee] [This Must Be the Place]]      |
| [[Aliens] [/award/award_winner/awards_won] [Isle of Man]]                     |

Table 9: RGVAE latent space interpolation between two triples with $\delta = 0.6$ and standard loss.

B.2 Interpolation per latent dimension

| RGVAE PERMUTE                                                                 |
|--------------------------------------------------------------------------------|
| $i_d = 1$                                                                      |
| [[Estudiantes Tecos] [/film/film/costume_design_by] [Sylvester Stallone]]     |
| [[The Man with the Golden Gun] [/film/film/costume_design_by] [Bill Fagerbakke]] |
| [[BAFTA Award for Best Film Music] [/film/film/costume_design_by] [Writer-GB]] |
| [[The Shipping News] [/film/film/costume_design_by] [Teri Garr]]              |
| [[Robert De Niro] [/film/film/costume_design_by] [Cesar Romero]]              |
| [[Inception] [/film/film/costume_design_by] [Hertha BSC Berlin]]              |
| [[Walter F. Parkes] [/film/film/costume_design_by] [Mary Lynn Rajskub]]       |
| [[Lorraine Bracco] [/film/film/costume_design_by] [Electronic keyboard]]      |
| [[Ben Folds] [/film/film/costume_design_by] [1956 Summer Olympics]]           |
| [[Eurasia] [/film/film/costume_design_by] [Video]]                            |

| $i_d = 2$                                                                      |
| [[Piano] [/film/film/costume_design_by] [Janes Addiction]]                    |
| [[Razzie Award for Worst Director] [/film/film/costume_design_by] [Visual Effects Director]] |
| [[Gone Baby Gone] [/film/film/costume_design_by] [Josh Sussman]]              |
| [[Ellen Pompeo] [/film/film/costume_design_by] [Joss Whedon]]                 |
| [[Bachelor of Arts] [/film/film/costume_design_by] [Chicago]]                 |
| [[News Corporation] [/film/film/costume_design_by] [2012 Summer Olympics]]    |
| [[Ukulele] [/film/film/costume_design_by] [85th Academy Awards]]             |
| [[Columbia Pictures] [/film/film/costume_design_by] [The Last Picture Show]]  |
| [[Catherine Keener] [/film/film/costume_design_by] [Gladyss Knight]]         |
| [[Amelia] [/film/film/costume_design_by] [Hikaru Utada]]                      |

| $i_d = 3$                                                                      |

33
[[Steve Carell] [film/film/costume_design_by] [Australia]]
[[About a Boy] [film/film/costume_design_by] [United States Dollar]]
[[Academy Award for Best Cinematography] [film/film/costume_design_by] [Defender]]
[[Jayma Mays] [film/film/costume_design_by] [Broadcast Film Critics Association Award]]
[[Arthur C. Clarke] [film/film/costume_design_by] [Comedy-GB]]
[[1968 Summer Olympics] [film/film/costume_design_by] [Quarterback]]
[[Tim Rice] [film/film/costume_design_by] [Electric guitar]]
[[The Faculty] [film/film/costume_design_by] [Linebacker]]
[[Academy Award for Best Screenplay] [film/film/costume_design_by] [Singer-songwriter-GB]]
[[Raoul Walsh] [film/film/costume_design_by] [Julianna Margulies]]

i_d = 4

[[Trumpet] [film/film/costume_design_by] [Argentina]]
[[Costa Mesa] [film/film/costume_design_by] [51st Annual Grammy Awards-US]]
[[Amanda Lear] [education/educational_degree/people_with_this_degree] [New Jersey]]
[[Jean-Paul Sartre] [film/film/costume_design_by] [Hong Kong]]
[[Charles LeMaire] [film/film/costume_design_by] [United States of America]]
[[13 Assassins] [film/film/costume_design_by] [V for Vendetta]]
[[Academy Award for Best Original Song] [film/film/costume_design_by] [Million Dollar Baby]]
[[Alan Parsons] [film/film/costume_design_by] [Mark Kirkland]]
[[Africa] [film/film/costume_design_by] [General-GB]]
[[Joanna Lumley] [film/film/costume_design_by] [Melanie Griffith]]

i_d = 5

[[Brighton Rock] [film/film/costume_design_by] [Brown University]]
[[Claude Rains] [film/film/costume_design_by] [Dhaka]]
[[Drums] [film/film/costume_design_by] [ Marty Stuart]]
[[Alameda County] [film/film/costume_design_by] [Striptease]]
[[Bachelor of Arts] [film/film/costume_design_by] [G-Force]]
[[Owen Wilson] [film/film/costume_design_by] [University of London]]
[[Being There] [film/film/costume_design_by] [14th Screen Actors Guild Awards]]
[[Tim McGraw] [film/film/costume_design_by] [Japanese Language]]
[[Brandenburg] [film/film/costume_design_by] [Honolulu]]
[[Rekha] [film/film/costume_design_by] [Defender]]

i_d = 6

[[Catherine Deneuve] [film/film/costume_design_by] [Robert Ridgely]]
[[Felix Bastians] [film/film/costume_design_by] [English Language]]
[[Denzel Washington] [film/film/costume_design_by] [German Democratic Republic]]
[[Gina Gershon] [film/film/costume_design_by] [Shanghai Greenland F.C.]]
[[Peter Cook] [film/film/costume_design_by] [Carrie Fisher]]
[[Jim Taylor] [film/film/costume_design_by] [Private university]]
[[Guitar] [film/film/costume_design_by] [Screenwriter]]
[[Broadcast Film Critics Association Award] [film/film/costume_design_by] [Running back]]
[[Southern Methodist University] [film/film/costume_design_by] [Mellotron]]

i_d = 7

[[Grammy Award for Best R&B Performance] [film/film/costume_design_by] [Tunisia]]
[[Comedy-GB] [film/film/costume_design_by] [Louisiana State University]]
[[Yes Man] [film/film/costume_design_by] [Brooklyn]]
[[Justin Chambers] [film/film/costume_design_by] [David Petraeus]]
[[Akiva Schaffer] [film/film/costume_design_by] [United States Dollar]]
[[Manovian Voivodeship] [film/film/costume_design_by] [Germany]]
[[Kevin Betsy] [film/film/costume_design_by] [“One Flew Over the Cuckoos Nest”]]
[[Tombstone] [film/film/costume_design_by] [Drum]]
[[Piranha DD] [film/film/costume_design_by] [United States of America]]
[[Sire Records] [film/film/costume_design_by] [Primetime Emmy Award]]

i_d = 8

[[Coconut milk] [film/film/costume_design_by] [Goa]]
[[Academy Award for Best Picture] [base/eating/practicer_of_diet/diet] [A Beautiful Mind]]
[[Indiana Jones and the Temple of Doom] [film/film/costume_design_by] [Romania]]
[[Charterhouse School] [film/film/costume_design_by] [Marilyn Manson]]
[[Verizon Communications] [film/film/costume_design_by] [Japan]]
Table 10: Interpolation of each latent dimension for the RGVAE with graph matching.

### RGVAE STANDARD

**i_d = 1**

- [1976 Winter Olympics] [base/marchmadness/ncaa_basketball_tournament/seeds] [Iain Glen]
  - [Judy Greer] [film/actor/film] [Arkansas]
  - [Vincent Pastore] [film/film/genre] [Vice President-GB]
  - ["Ulysses Gaze"] [people/person/gender] [ESPN]
  - [Manchester University] [award/ranked_item/appears_in_ranked_lists] [Silver medal]
  - [Wesleyan University] [business/job_title/people_with_this_title] [Lisa Gay Hamilton]
  - [Iraq] [music/instrument/family] [Greg Daniels]
  - [Empire of the Sun] [people/person/profession] [Catherine Keener]
  - [Leslie Bricusse] [award/award_nominee] [Jane Fondal]
  - [Forward] [film/distributor/films_distributed] [Sandra Bullock]

**i_d = 2**

- [Saxophone] [award/award_winning_work/awards_won] [The Prince of Egypt]
  - [Rang De Basanti] [location/location/contains] [1992 Summer Olympics]
  - [Small Soldiers] [award/award_nominee] [New York City]
  - [Bill Hicks] [award/award_nominee] [Paul Williams]
  - [Hairspray] [user/jg/default_domain/olympic_games/sports] [Michael K. Williams]
  - [The Aviator] [film/actor/film] [Goalkeeper]
  - [Joel Grey] [award/award_nominee] [Vladimir Vladimirovich Nabokov]
  - [Vibraphone] [film/film/films_distributed] ["Jews harp"]
  - [Ice Age: Dawn of the Dinosaurs] [music/group_member/membership] [Club Atlas]
  - [Doctor Zhivago] [film/film/release_date_s] [Terror in the Aisles]

**i_d = 3**

- [Real Sociedad] [common/topic/webpage/common/webpage/category] [Palmitic acid]
  - [Soul music] [people/person/profession] [Alameda]
  - [Viggo Mortensen] [medicine/disease/risk_factors] [Love Story]
  - [Finland] [sports/sports_position/players] [DVD]
  - [Crazy Heart] [tv/tv_producer/programs_produced] [Los Angeles]
  - [Drew Barrymore] [tv/tv_writer/tv_programs] [Road cycling]
| id | Movie/TV Show | Notes |
|----|----------------|-------|
| 4  | BAFTA Award for Best Screenplay | [award/award_nominee] [John Ortiz] |
|    | [Humford] [film/film/release_date] [Howard Gordon] |
|    | [Grammy Award for Best Musical Album] [people/person/places_lived] [Batman Begins] |
|    | [Cats & Dogs] [sports/sports_team/colors] [Musician-GB] |
| 5  | Saint Marys College | [government/political_party/politicians_in_this_party] [Joel Coen] |
|    | [Eva Longoria] [people/person/profession] [The Lives of Others] |
|    | [United States Congress] [award/award_nominee] [Tony Award for Best Actor] |
|    | [Midfielder] [music/instrument/instrumentalists] [Elizabeth] |
|    | [Brad Pitt] [film/actor/film] [South Sudan] |
|    | [Django Unchained] [user/ktrueman/default_domain] [France] |
|    | [McMaster University] [music/artist(track_contributions)] [Philippines] |
|    | [The Stand] [people/person/profession] [Curly Howard] |
|    | [Jerome Flynn] [award/award_nominee] [Piano] |
|    | [Brighton Rock] [location/location/contains] [Forward] |
| 6  | Jennifer Aniston | [people/person/place_of_birth] [Peru] |
|    | [Timothy Busfield] [film/film/release_date] [Defender] |
|    | [Nirupa Roy] [food/food/nutrients] [Wake Forest University] |
|    | [Kidderminster F.C.] [award/award_winning_work(awards_won_winner)] [United States Dollar] |
|    | [BAFTA Award for Best Actress] [location/statistical_region] [Forest Lawn Memorial Park] |
|    | [Babylon 5] [award/award_nominee] [Hong Kong] |
|    | [Johann Wolfgang von Goethe] [film/actor/film] [Sunday Bloody Sunday–GB] |
|    | [St. Augustine] [people/marriage_union_type/unions_of_this_type] [Columbia Pictures] |
|    | [Brokeback Mountain] [award/award_nominee] [Oscar] |
|    | [Shanghai Knights] [film/film/cinematography] [United States Dollar] |
| 7  | Cape Fear | [award/award_nominee] [DVD] |
|    | [Busta Rhymes] [film/film/language] [United States Dollar] |
|    | [The Bourne Identity] [people/person/profession] [Ally Sheedy] |
|    | [Taxi Driver] [people/person/profession] [Ipswich Town F.C.] |
|    | [France] [film/actor/film] [Greensboro] |
|    | [The Constant Gardener] [award/award_nominee] [Academy Award for Best Screenplay] |
|    | [North Atlantic Treaty Organization (NATO)] [people/person/profession] [Makeup Artist-GB] |
|    | [Trey Parker] [time/event/locations] [University of the West Indies] |
|    | [Mockumentary] [people/deceased_person/place_of_death] [Newport Beach] |
|    | [Phillips Academy] [music/performance_role/regular_performances] [Police procedural] |
| 8  | Gladiator | [people/person/profession] [Clarinet] |
|    | [Jenna Fischer] [sports/sports_team_location/teams] [Spain] |
|    | [Israel] [base/local_food сезон/month/produce_available] [Yale University] |
|    | [Tompkins County] [organization/organization/place_founded] [William Shakespeare] |
|    | [Bas-Rhin] [film/film/release_date] [Actor–GB] |
|    | [Virgin Records] [location/hud_foreclosure_area] ["Peoples Action Party"] |
|    | [Farida Jalal] [education/educational_institution/school_type] [Primetime Emmy Award] |
|    | [Piano] [award/award_winning_work(awards_won)] [Marty Stuart] |
|    | [John C. Reilly] [location/location/contains] [Jim Broadbent] |
|    | [Tim Roth] [film/film/language] [Egypt] |
| 9  | Ennio Morricone | [award/award_winner(awards_won_winner)] [98th United States Congress] |
|    | [Mali Finn] [base/marshmadness/ncaa_basketball_tournament/seeds] [Angie Dickinson] |
|    | [David Walliams] [music/performance_role/regular_performances] [Toy Story 3] |
|    | [Marriage] [award/award_ceremony/awards_presented] [New York City] |
|    | [Philip Yordan] [soccer/football_team/current_roster] [English Language] |
|    | [63rd Primetime Emmy Awards] [film/film/other_crew] [Stephen Frears] |
|    | [Hud] [film/film/other_crew] [Potassium] |
|    | [Arkansas] [film/film/featured_film_locations] [Special Effects Supervisor] |
|    | [Fran Drescher] [film/film_distributor/films_distributed] [Giant] |
|    | [Charlotte Gainsbourg] [travel/travel_destination/climate] [United Kingdom] |
Table 11: Interpolation of each latent dimension for the RGVAE with standard loss function.

| $i_{d_1}$ = 1 | $i_{d_2}$ = 2 | $i_{d_3}$ = 3 | $i_{d_4}$ = 4 |
|---------------|---------------|---------------|---------------|
| [[Amman] /user/tsegaran/random/taxonomy_subject/entry [Bill Cosby]] | [[UTV] /base/localfood/seasonal_month/produce_available [Drama]] | [[Dick Powell] /film/film/release_date_s [Alex Borstein]] | [[Acid house] /user/tsegaran/random/taxonomy_subject/entry [Big Love]] |
| [[Kristen Stewart] /people/person/taxonomy_subject/entry [United States of America]] | [[Cave of Forgotten Dreams] /film/film/release_date_s [The Lord of the Rings]] | [[Jeff Richmond] /film/film/personal_appearances [Never Say Never Again]] | [[The Amazing Spider-Man] /user/tsegaran/random/taxonomy_subject/entry [Animaniacs]] |
| [[Dean Koontz] /user/tsegaran/random/taxonomy_subject/entry [Fukushima Prefecture]] | [[2008 Summer Olympics] /time/event/locations [Chulalongkorn]] | [[Richard Burton] /user/tsegaran/random/taxonomy_subject/entry [Pinellas County]] | [[(Heavy metal] /user/tsegaran/random/taxonomy_subject/entry [Mark Ronson]] |
| [[Western Front] /user/tsegaran/random/taxonomy_subject/entry [Member of Congress]] | [[(Omarion] /award/award_category/disciplines_or_subjects [Taito Corporation]] | [[Cavendish] /government/politician/government_positions_held [Drama]] | [[Southeast Asia] /location/hud/county_place/county [Franklin Medal]] |
| [[Cyran de Bergerac] /user/tsegaran/random/taxonomy_subject/entry [Star Wars Episode III]] | [[(Economist] /sports/sports_team/roster [Tommy Lee Jones]] | [[(Jon Stewart] /film/film/country [Warren Ellis]] | [[Jon Stewart] /film/film/administrative_division [95th United States Congress]] |
| [[(Country] /user/tsegaran/random/taxonomy_subject/entry [Lois Smith]] | [[(Independent Spirit Award for Best Director] /film/film/written_by [Defender]] | [[(Marv Wolfman] /location/administrative_division [95th United States Congress]] | [[(Real Betis] /film/film/release_date_s [Obesity]] |
Table 12: Interpolation of each latent dimension for the RGVAE with $\delta = 0.6$ and graph matching.
Table 13: Interpolation of each latent dimension for the RGVAE with $\delta = 0.6$ and standard loss.