# Improving Graph Classification by Means of Linear Combinations of Reduced Graphs 

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Keywords: Structural Pattern Recognition, Graph Matching, Genetic Algorithm, Multiple Classifier Systems.


#### Abstract

The development and research of graph-based matching techniques that are both computationally efficient and accurate is a pivotal task due to the rapid growth of data acquisition and the omnipresence of structural data. In the present paper, we propose a novel framework using information gained from diversely reduced graph spaces to improve the classification accuracy of a structural classifier. The basic idea consists of three subsequent steps. First, the original graphs are reduced to different size levels with the aid of node centrality measures. Second, we compute the distances between the reduced graphs in the corresponding graph subspaces. Finally, the distances are linearly combined and fed into a distance-based classifier to produce the final classification. On six graph datasets we empirically demonstrate that classifiers clearly benefit from the combined distances obtained in the graph subspaces.


## 1 INTRODUCTION

Graph classification is a prominent task in structural pattern recognition (e.g., community classification of social groups (Liu et al., 2015) or the determination whether or not molecular compounds are toxic (Jin et al., 2010), to name only two examples). From a very broad perspective, structural pattern classification is relatively similar to supervised statistical machine learning. Both approaches aim to extract useful formal representations out of given data and map those extracted representations to one of the available categories. Unfortunately, due to the inherent relational structure of graphs, one cannot directly apply methods, originally developed for statistical data, to graph data.

A prominent way to deal with this problem is to embed graphs into a vector space and eventually apply a standard machine learning framework on the embedded graphs. Yet, it is not a trivial task to find an embedding function that respects the inherent relations that may exist between the elements of a graph. Graph embedding approaches range from spectral methods (Caelli and Kosinov, 2004; Qiu and Hancock, 2006), to Graph Kernels (Kriege et al.,

[^0]2020), Dissimilarity Embeddings (Riesen and Bunke, 2010), and Graph Neural Networks (GNN) (Wu et al., 2021).

Research in the field of graph embedding has made great progress over the past few years. However, there is still a risk of loosing discriminating power during the embedding process. Moreover, some of the mentioned approaches still rely on graph matching, while others (e.g. GNN) suffer from their non-interpretability.

The present paper is focused on graph matching methods for graph classification (Conte et al., 2004), and in particular, on Graph Edit Distance (GED) (Bunke and Allermann, 1983; Sanfeliu and Fu, 1983). Actually, distance-based classifiers coupled with GED have shown reasonable classification accuracy on numerous classification tasks (Maergner et al., 2018; Fuchs and Riesen, 2021). The present paper aims to improve the classification accuracy of a distance-based classifier using GED. To achieve this goal, we propose a novel framework that gains extra information out of various subgraphs extracted from the original graphs.

More formally, the basic idea of our approach consists of mapping the graphs into various reduced graph spaces, where each graph space contains graphs that consist of the most important nodes only. Node centrality measures are employed for this graph re-
duction. Next, we compute the GED between the graphs in each reduction level and combine the distances or predictions obtained. The meta-parameters, viz. the weight coefficients for the linear combination, are either optimized via grid-search or by means of a genetic algorithm. Finally, the combined distances or predictions are used as basis for the final classification.

The remainder of this paper is organized as follows. In Section 2, we formally introduce the notion of graphs and give a brief review of graph matching methods. In Section 3, we describe in detail our novel method to produce differently sized graphs and combine the distances or predictions obtained in the reduced graph spaces. In Section 4, we present and discuss the results of our experimental evaluation, and in Section 5, we conclude the paper and discuss some future research ideas.

## 2 RELATED WORK

In this section, we provide a formal definition of graph structures and briefly review common graphbased matching methods (including the one actually used).

### 2.1 Graph Structure

A graph $G=(V, E)$ in a graph space $\mathcal{G}$ is defined as a set of nodes $V$ and a set of edges $E \subset(V \times V)$ between these nodes. In case of directed graphs, an edge starts at node $u \in V$ and ends at node $v \in V$ and is denoted by $(u, v) \in E$. If the graph is undirected, an edge is defined as $(u, v) \in E \leftrightarrow(v, u) \in E$. A node (and/or edge) labeling function $\mu: V \mapsto L_{V}$ (and/or $v: E \mapsto L_{E}$ ) is defined in the case of labeled nodes (and/or labeled edges). In the remainder of this paper, we limit our work to simple, undirected graphs, i.e., graphs that have at most one edge between each pair of nodes and no edge between a node and itself. The size of a graph is commonly defined as the cardinality of its node set $|V|$.

### 2.2 Graph Matching Methods

We focus our research on structural methods in pattern recognition using inexact graph matching algorithms. In this scenario, the matching constraints are generally relaxed such that matchings between arbitrary graphs are possible. Inexact matching is particularly useful when (small) errors appear in the input graphs or when one wants to derive a numerical degree of similarity/dissimilarity between two graphs
(rather than checking for (sub-)graph isomorphism only).

Actually, various inexact graph matching methods have been proposed over the years (Conte et al., 2004). The authors of (Escolano et al., 2011), for instance, proposed an inexact graph matching that depends on spectral features that represent a graph as a bag of partial node coverages. (Kashima et al., 2003) introduced a probabilistic method for defining graph matching based on sequences of node indices and a random walk. (Escolano et al., 2017) proposed a new similarity measure that computes the mutual information between graphs with a combination of copula functions to perform graph comparison.

In the present paper, we employ Graph Edit Distance (GED) (Bunke and Allermann, 1983; Sanfeliu and Fu, 1983) as basic matching paradigm. GED is a well known graph matching technique with great adaptability power. It has gained interest as a dissimilarity measure on a broad range of problems and applications (Riesen and Bunke, 2010; Cortés and Serratosa, 2015). GED defines the dissimilarity between two graphs $G$ and $G^{\prime}$ by computing the least amount of transformation (i.e., edit operations) required to convert $G$ into $G^{\prime}$. Three edit operations on both nodes and edges are typically defined, viz. insertion, deletion, and substitution. Each of those edit operations can have a custom cost that emphasizes the severity of the given graph transformation. This cost is typically formalized by means of a cost function $c(\cdot)$. The goal of GED is to find an edit path between $G$ and $G^{\prime}$, i.e., a set $\left\{e_{1}, \ldots, e_{k}\right\}$ of $k$ edit operations $e_{i}$, that transforms $G$ into $G^{\prime}$, while minimizing the overall transformation cost.

In order to compute the optimal edit path, GED computations are typically based on combinatorial search methods with exponential complexity, which makes GED actually unfeasible on large graphs However, different GED approximations (Riesen and Bunke, 2009; Fischer et al., 2014) have been proposed allowing us to find a reasonable approximation of GED in polynomial time. In the present paper we actually employ the approximation proposed in (Riesen and Bunke, 2009).

As discussed in the next section, we propose to compute GED on differently sized graphs and eventually combine the outcome. Hence, our novel method is somehow related to the recently proposed hierarchical framework for inexact graph matching (Riba et al., 2020). They use a community detection method to construct a hierarchy of compressed graphs. The matching is then performed in a coarse-to-fine fashion. Starting the matching at the lowest level (i.e., the level where the graphs are the most compressed) and


Figure 1: Example of our graph reduction method with different levels of $\lambda$ on two molecular graphs from the AIDS dataset.
going up level by level if necessary.

## 3 OUR NOVEL METHOD

In the present paper we want to research the benefits and limitations of the multi-level information gained from reduced graphs in a multiple classifier scenario. That is, we investigate whether or not the simultaneous use of this extra information improves the classification performance of a distance-based classifier. The basic idea is to combine the information gained at different reduction levels and perform the classification on this combination.

### 3.1 Graph Reduction

Our approach depends on graph subspaces. Hence, we conduct a fast yet deterministic mapping of our graphs to different graph subspaces. Our reduction method is based on centrality measures for nodes in a network.

In the present paper, we use the PageRank centrality score (Brin and Page, 1998). Note, however, that any other node centrality measure could be used as well (Newman, 2018). The basic concept of PageRank relies on the fact that a node's influence increases if it is connected to other influential nodes. To avoid that the most prestigious nodes in a graph widespread its high-centrality score too all its neighbors, the influence of a node is diluted proportionally to the number of neighbors.

Given the centralities for each node, we sort them according to their respective centrality score from the least to the most influential one. Given a reduction factor $\lambda \in[0,1]$ we then keep the $\lambda|V|$ most influential nodes in the graph while the other nodes (including
their edges) are discarded.
The reduction factor $\lambda$ roughly corresponds to the percentage of remaining nodes of the original graphs (i.e., if we set $\lambda=0.6$, approximately $60 \%$ of the nodes remain in the reduced version of the graph). We vary the reduction factor such that it takes the following values $\lambda \in\{1.0,0.8,0.6,0.4,0.2\}$, where $\lambda=1.0$ represents the full graphs (i.e., graphs where no nodes are removed).

In Fig. 1 we show two examples of graph reductions with different levels of $\lambda$ on two graphs from the AIDS dataset (the datasets are described in the next Section).

From now on, we describe a reduced graph (with reduction factor $\lambda$ ) as $G_{\lambda}=\left(V_{\lambda}, E_{\lambda}\right)$. Reducing the $N$ graphs of a dataset for a given $\lambda$ gives us a reduced graph space $\mathcal{G}_{\lambda}=\left\{G_{\lambda}^{(1)}, \ldots, G_{\lambda}^{(N)}\right\}$. Repeating this procedure for all possible values of $\lambda$ we produce multiple reduced graph spaces.

In each graph subspace $\mathcal{G} \lambda$ we are now able to compute graph dissimilarities between pairs of graphs. Formally, for each graph $G_{\lambda}^{(i)}$ we produce a distance vector $d_{i}=\left[d_{i, 1}, d_{i, 2}, \ldots, d_{i, N}\right]$ between itself and all other graphs and combine them to create a distance matrix $D_{\lambda}$.

Given these distance matrices for different levels, we pursue two different strategies of combining them to come up with a final classification. Both strategies are briefly described in the next two subsections. In both scenarios a distance based classifier, viz. a $K$ -nearest-neighbor classifier (KNN), is employed. The KNN has the clear advantage that it directly operates on the resulting distances without any additional training and can thus be used as indicator for the quality of the underlying distance.

Table 1: Statistics of the graph datasets. We show the number of graphs $(|G|)$ with the size of the training, validation and test set ( $\operatorname{tr}$, va, te), the number of classes $(|\Omega|)$ and the average number of nodes and edges $(\emptyset|V|, \emptyset|E|)$ per dataset.

| Dataset | $\|G\|$ (tr, va, te) | $\|\Omega\|$ | $\emptyset\|V\|$ | $\emptyset\|E\|$ |
| :--- | :---: | :---: | :---: | :---: |
| AIDS | $2,000(250,250,1,500)$ | 2 | 9.5 | 10.0 |
| Mutagenicity | $4,337(1,500,500,2,337)$ | 2 | 30.3 | 30.8 |
| NCI1 | $4,110(1,500,500,2,110)$ | 2 | 29.9 | 32.3 |
| Proteins | $1,113(660,220,223)$ | 2 | 39.1 | 72.8 |
| Enzymes | $600(360,120,120)$ | 6 | 32.6 | 62.1 |
| IMDB Binary | $1,000(600,200,200)$ | 2 | 19.8 | 96.5 |

### 3.2 Combination of the Distances

The first idea is to linearly combine the given distance matrices $D_{\lambda}$ with $\lambda \in\{1.0,0.8,0.6,0.4,0.2\}$ into a meta-distance matrix

$$
\begin{equation*}
\mathcal{D}=\sum_{\lambda} \omega_{\lambda} D_{\lambda} \tag{1}
\end{equation*}
$$

Eventually, we use this matrix $\mathcal{D}$ to perform the classification with a KNN. Note the parameter $\omega_{\lambda} \in$ $[0,1]$ which is used to weight the relative importance of the reduced graph space at level $\lambda$.

### 3.3 Combination of the Predictions

The second combination idea is defined as follows. Instead of combining the distances, we combine the predictions obtained at each reduced graph subspace.

Formally, we retrieve the class prediction of a KNN for each graph sample at level $\lambda$. Thus, we have a prediction vector $p_{\lambda}=\left[p_{1}, p_{2}, \ldots, p_{n}\right]^{T}$ for each graph subspace where $p_{i}$ corresponds to the prediction of the $i$-th graph. We now linearly combine the weighted prediction vectors with

$$
\begin{equation*}
\mathcal{P}=\sum_{\lambda} \omega_{\lambda} p_{\lambda} \tag{2}
\end{equation*}
$$

That is, we conduct a weighted majority voting (Kittler, 2002).

### 3.4 Optimization of the Parameters $\omega$

The parameter vector $\boldsymbol{\omega}=\left(\omega_{\lambda_{1}}, \ldots, \omega_{\lambda_{t}}\right)$ contains the $t$ parameters $\omega_{\lambda_{i}}$ for all reduction factors $\lambda_{i}$ where $t$ corresponds to the graph subspaces actually available. That is, this parameter weights the importance of each reduced graph space. We aim for a linear combination of the reduced graph subspaces, and we add a constraint on $\boldsymbol{\omega}$ such that the sum of the parameters equals 1. Formally, we have

$$
\begin{equation*}
\sum_{\lambda_{i}} \omega_{\lambda_{i}}=1 \text { and } \omega_{\lambda_{i}} \in[0,1] \tag{3}
\end{equation*}
$$

Our goal is now to optimize the linear coefficients to obtain $\boldsymbol{\omega}^{*}$ such that the meta-distance matrix $\mathcal{D}$ or the combined predictions $\mathcal{P}$ achieve the best classification accuracy. The optimization is performed with two different methods.

The first method is to use a grid search over the parameter space. Note, however, that the search space has a size of $O\left(\mathrm{D}^{t}\right)$, where D is the number of possible values for $\omega_{\lambda_{i}}$ and $t$ is the total number of subspaces to be combined. In order to have a reasonable search time, we use five reduction levels only ( $\lambda \in\{1.0,0.8,0.6,0.4,0.2\}$ ) and 11 possible weighting factors $\left(\omega_{\lambda_{i}} \in\{1.0,0.9, \ldots, 0.1,0.0\}\right.$ leading to $11^{5}=161,051$ different possibilities. To reduce the risk of overfitting during the grid search optimization method, we apply a 5 -fold cross-validation.

The second optimization method is to use a Genetic Algorithm (GA) (Eiben and Smith, 2015). Each $\omega_{\lambda_{i}}$ represents a gene in the chromosome $\omega$. The fitness function of each chromosome is the classification accuracy of the KNN coupled with corresponding weighted combination. This score defines how well a given weighting vector solves the optimization problem. The first step of a genetic algorithm is to randomly generate a set of chromosomes. Then, GA evaluates those chromosomes with the fitness function, selects the best performing chromosomes and combines them with a crossover operation to create the next generation of chromosomes. The crossover is done uniformly among the genes of the parents. Each newly created chromosome has a chance to mutate with probability $p_{m}=0.1$. These three steps are repeated until a specified criterion is reached. In the present paper we use a fixed number of optimization iterations. GAs allow a more efficient search procedure over the parameter space compared to grid search. On the other hand, it is more prone to overfitting without offering well-defined regularization techniques to prevent it.

Table 2: Classification accuracy [\%] obtained on the test set with linear combinations of reduced graphs. We present the results obtained by a KNN for the baseline and our two combination methods that are Combination of Distance Matrices (CoDM) and Combination of Predictions (CoP). The best result per dataset is shown in boldface. ( $/ / \bullet$ : statistically significantly better/worse than the baseline on a $5 \%$ level using a Z-test.).

| Method $\backslash$ Dataset | AIDS | Mutagenicity | NCI1 | Proteins | Enzymes | IMDB-Binary |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Baseline KNN | 98.53 | 71.33 | 70.33 | 73.82 | 41.67 | 66.00 |
| CoDM - Grid Search | $99.13 \circ$ | 71.84 | 72.09 | 73.39 | 45.83 | 64.50 |
| CoDM - Genetic Algorithm | $99.13 \circ$ | $\mathbf{7 2 . 6 6}$ | $\mathbf{7 3 . 2 2} \circ$ | 75.54 | $\mathbf{4 8 . 3 3} \circ$ | 66.00 |
| CoP - Grid Search | $\mathbf{9 9 . 3 3} \circ$ | 72.32 | 70.52 | 73.82 | 41.67 | $\mathbf{7 0 . 0 0}$ |
| CoP - Genetic Algorithm | $99.13 \circ$ | 71.84 | 70.52 | $\mathbf{7 6 . 3 9} \circ$ | $37.50 \bullet$ | $\mathbf{7 0 . 0 0}$ |

## 4 EXPERIMENTAL EVALUATION

### 4.1 Datasets

We evaluate our novel procedure on six datasets in total. Two datasets are retrieved from the IAM graph repository (Riesen and Bunke, 2008) ${ }^{1}$ (AIDS, Mutagenicity) and the four others from TUDataset (Morris et al., 2020) $)^{2}$ (NCI1, Proteins, Enzymes, IMDB Binary).

Table 1 shows some graph properties such as the number of graphs, the number of classes, and the average number of nodes and edges per dataset. The first three datasets (AIDS, Mutagenicity, and NCI1) represent small molecular compounds. The AIDS dataset contains molecules from two categories, viz. molecules that have an effect against the HI virus or not. Graphs in Mutagenicity dataset represent chemical compounds that can be classified as mutagen/nonmutagen. The graphs in the NCI1 dataset (Wale and Karypis, 2006) represent chemical compounds that are positive or negative lung cancerous cells. The Proteins graphs represent proteins that are categorized as enzymes/non-enzymes, while the graphs in Enzymes correspond to tertiary protein structures compiling 6 enzymes (Borgwardt et al., 2005). The graphs from IMDB-Binary (Yanardag and Vishwanathan, 2015) represent individual movies from two categories.

The split of the datasets into training, validation, and test sets is done as follows. The graphs from the IAM datasets are divided according to the provided splitting. The partitioning of the NCI1 dataset is done to match the size of the splitting of the Mutagenicity dataset. The remaining datasets are split according to the $60-20-20 \%$ split rule for the training, validation, and test set, respectively.

[^1]
### 4.2 Experimental Setup

In our experiment we want to verify whether or not using extra information gained from the reduced versions of the graphs can help improving the overall classification accuracy. To this end, we first run a basic KNN classifier on all datasets using the original graphs (i.e., $\lambda=1.0$ ). This actually builds the baseline of our evaluation. We individually optimize KNN's hyperparameters on the validation set. The optimized parameters consist of $k$ that designate the number of neighbors used by the KNN and $\alpha$ that ponders the relative importance of node and edge edit operation costs during GED computation. The obtained parameters are presented for each dataset in Table 3.

Table 3: Optimal $k$ and $\alpha$ obtained during the hyperparameter optimization on the validation set.

| Dataset | $\boldsymbol{k}$ | $\boldsymbol{\alpha}$ |
| :--- | :---: | :---: |
| AIDS | 1 | 0.7 |
| Mutagenicity | 5 | 0.6 |
| NCI1 | 5 | 0.7 |
| Proteins | 3 | 0.9 |
| Enzymes | 1 | 0.9 |
| IMDB Binary | 5 | 0.9 |

For the experiments with the reduced graphs, we set the reduction factors to $\lambda \in\{1.0,0.8,0.6,0.4,0.2\}$. We reused the hyperparameters $k$ and $\alpha$ found during the individual optimization. Eventually, we compute all distance matrices $D_{\lambda}$ between training and validation graphs, and perform the optimization of the linear coefficicients with both grid search and genetic algorithm. The optimized parameters (using either grid search or a genetic algorithm) are finally used to classify the test graphs. In Fig. 2, we plot the optimized weights $\boldsymbol{\omega} *$ found during optimization. The plot shows the importance of the reduced graph spaces for all experiments conducted. We observe that, in general, the original graph space tends to have
a strong impact in the majority of the tested settings. However, it is also clearly observable that all other graph subspaces significantly contribute to the final distance and/or prediction.


Figure 2: Importance of each reduced graph space in the linear combination for all optimization methods, combination methods, and datasets.

### 4.3 Results

In Table 2, we present the classification accuracy obtained by our combination methods on the test set. Our two combination methods (Combination of Distance Matrices (CoDM) and Combination of Predictions ( CoP )) are optimized with both grid search and GA.

In general, we observe that combining the distances or the predictions of the reduced graphs improves the classification accuracy when compared to the baseline on all datasets. That is, on all datasets the overall best result (shown in boldface) is achieved by one of the combination methods.

On half of the datasets (Mutagenicity, NCI1, Enzymes) the classifier that relies on the combination of the distances (CoDM) optimized with GA achieves the overall best accuracies. We observe statistically significant improvements over the baseline on NCI1 and Enzymes by nearly $3 \%$ and $7 \%$, respectively. On the three other datasets, the classifier that uses the combination of the predictions ( CoP ) on the reduced graphs leads to the overall best classification accuracies. The improvements on the AIDS and Proteins dataset compared to our reference system are actually statistically significant.

In general, the optimization using the GA produces better results than the grid search for both combination methods (CoDM and CoP). In 9 out of 12 comparisons the GA achieves the same or better classification accuracies as the grid search. However, the drawback of this optimization technique is the risk of overfitting that clearly appears on the Enzymes dataset using CoP.

## 5 CONCLUSION

In present paper, we propose a novel framework that combines multi-level information computed on reduced versions of graphs. More specifically, we produce multiple reduced graph spaces and use GED to compute the distances between the graphs in those reduced graph spaces. In order to reduce the original graphs we evaluate the importance of each node by means of PageRank and discard the least important nodes in an iterative procedure. We linearly combine the weighted distances or the weighted predictions obtained in the various reduced graph subspaces. We optimize the linear combinations using two strategies, viz. a grid search and a genetic algorithm. By performing an empirical evaluation on a wide range of applications, we demonstrate the benefit of using extra information gained from reduced graphs. Particularly, we show that combining distances or predictions of multi-level graphs assists a distance-based classifier and improves its classification power. That is, on all data sets at least one of the combinations outperforms the reference system. On four out of six datasets we observe statistically significant improvements over the reference system.

In future work we plan to improve the performance of the distance-based classifier by using other graph reduction methods. In particular, we aim at investigating methods that keep more coherent topological information of the graphs and would summarize the inner information of the nodes during the reduction process. Non-linear combinations of the dissimilarities or predictions could also be a rewarding avenue to be pursued. Finally, further empirical analysis can be done with our novel method. Specifically, we can compare the computational complexity of our multi-level framework to the baseline KNN. Additionally, a comparison of our method to other state-of-the-art methods can be conducted.

## ACKNOWLEDGEMENTS

This material is based upon work supported by the Swiss National Science Foundation (SNSF) under Grant Nr. 200021_188496.

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