Finding the Best Not the Most: Regularized Loss Minimization Subgraph Selection for Graph Classification

Shirui Pan^{a,*}, Jia Wu^a, Xingquan Zhu^b, Guodong Long^a, Chengqi Zhang^a

^aCentre for Quantum Computation & Intelligent Systems, FEIT, University of Technology Sydney, Australia ^bDepartment of Computer and Electrical Engineering & Computer Science, Florida Atlantic University, USA

Abstract

Classification on structure data, such as graphs, has drawn wide interest in recent years. Due to the lack of explicit features to represent graphs for training classification models, extensive studies have been focused on extracting the most discriminative subgraphs features from the training graph dataset to transfer graphs into vector data. However, such filter-based methods suffer from two major disadvantages: (1) the sub-graph feature selection is separated from the model learning process, so the selected most discriminative subgraphs may not best fit the subsequent learning model, resulting in deteriorated classification results; (2) all these methods rely on users to specify the number of subgraph features K, and suboptimally specified K values often result in significantly reduced classification accuracy.

In this paper, we propose a new graph classification paradigm which overcomes the above disadvantages by formulating subgraph feature selection as learning a *K*dimensional feature space from an *implicit* and *large* subgraph space, with the optimal *K* value being automatically determined. To achieve the goal, we propose a regularized loss minimization-driven (RLMD) feature selection method for graph classification. RLMD integrates subgraph selection and model learning into a unified framework to find discriminative subgraphs with guaranteed minimum loss *w.r.t.* the objective func-

^{*}Corresponding author. Address: Centre for Quantum Computation & Intelligent Systems, FEIT, University of Technology, Sydney, Ultimo, NSW 2007, Australia. Tel: +61 450768511. Fax: +61 2 9514 4535.

Email addresses: shirui.pan@uts.edu.au (Shirui Pan), jia.wu@student.uts.edu.au (Jia Wu), xzhu3@fau.edu (Xingquan Zhu), guodong.long@uts.edu.au (Guodong Long), chengqi.zhang@uts.edu.au (Chengqi Zhang)

tion. To automatically determine the optimal number of subgraphs K from the exponentially large subgraph space, an effective *elastic net* and a subgradient method are proposed to derive the stopping criterion, so that K can be automatically obtained once RLMD converges. The proposed RLMD method enjoys gratifying property including proved convergence and applicability to various loss functions. Experimental results on real-life graph datasets demonstrate significant performance gain.

Keywords: Feature Selection, Classification, Graph Classification, Sparse Learning

1 1. Introduction

Recent years have witnessed an increasing number of applications involving objects with structural relationships, including chemical compounds in Bioinformatics [1], brain networks [2], image structures [3], and academic citation networks [4]. For these applications, graph is a natural and powerful tool for modeling and capturing dependency relationships between objects.

Unlike conventional data, where each instance is represented in a feature-value vector format, graphs exhibit node-edge structural relationships and have no natural 8 vector representation¹. As a result, a common practice is to transfer graphs into vec-9 tors [5, 6, 7, 8, 9] in structure space or in Euclidean space, so that traditional machine 10 learning algorithms such as Support Vector Machines (SVM) and Decision Tree can 11 be applied. In the structure space (also referred to as quotient space) [7, 8], the dis-12 tance relations and nature of the original data are preserved, and some geometrical 13 and analytical concepts such as derivatives of functions on structures can be deter-14 mined, so that it can be applied to solve problems in structural pattern recognition. 15 In the Euclidean space, the structural relations may be lost, but it provides simpler 16 and more powerful analytical techniques for data analysis. Therefore, numerous ap-17 proaches [10, 9, 11, 12, 13, 14, 15, 16, 17, 18] have been proposed to represent graphs 18 in Euclidean space. The key idea of transferring graphs into vectors in Euclidean space 19 is to extract a set of subgraphs as features and use the presence/absence of the features 20

¹In this paper, we only consider graphs with labels but no other feature values on nodes and edges.

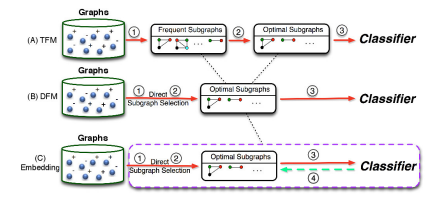


Figure 1: Subgraph-based methods for graph classification from the feature selection perspective. TFM methods (A) sequentially perform frequent subgraph mining ①, optimal feature selection ②, and classifier learning process ③. DFM methods (B) integrate the feature selection ② into the frequent subgraph mining ① process. Our embedding method RLMD (C) unifies all steps (①②③) into a whole framework, and iterates until convergence ④.

to represent each graph. From a feature selection perspective [19], these subgraphbased algorithms follow a filter approach for graph classification, *i.e.*, the subgraph feature selection and the subsequent model training are separated into two steps. In summary, existing filter-based graph classification methods roughly fall into the following two categories:

Two-step Filter Methods (TFMs): This type of method first mines a set of frequent 26 subgraphs as features and then applies a feature selection procedure to the discovered 27 subgraphs, and uses the selected subgraph features to learn a classifier (e.g., an SVM or 28 Naive Bayes), as shown in Fig. 1 (A). An early study [9] has shown that using frequent 29 subgraphs as features can achieve reasonable good classification results. However, be-30 cause TFMs separate subgraph feature discovery and feature evaluation into two steps, 31 they may suffer from severe disadvantage in that the number of discovered subgraphs 32 will grow exponentially when the minimum support value for subgraph mining is low. 33 As a result, it will make the feature selection step heavily time-consuming. On the other 34 hand, for relatively high minimum support values, many good subgraphs are pruned out 35 because they do not meet the frequency requirement, so cannot be found to represent 36 graphs. 37

Direct Filter Methods (DFMs): To improve the subgraph feature selection efficiency, 38 numerous approaches [11, 12, 15, 16, 17] have been proposed to combine subgraph 39 mining and feature selection into one step, representing a direct discriminative feature selection [18] scheme. So the feature selection is integrated into a subgraph mining 41 process (Fig. 1 (B)), with pruning rules derived from the anti-monotone property of 42 the significance (*p*-value) of each graph being used to reduce the search space. While 43 DFMs substantially overcome the subgraph feature selection bottleneck, they also have 44 a number of major disadvantages: (1) The subgraph selection is separated from the 45 model learning process, so the selected subgraphs features may not best fit the under-46 lying learning model, and (2) All these methods require users to specify the number of 47 subgraph features K, whereas the optimal number of subgraphs K required for training 48 a good classifier for graph classification is unknown and difficult to determine. Al-49 though subgraphs are selected using optimized measures, due to the redundancy inside 50 the feature set, the accuracy of the classifiers, when varying the number of selected 51 subgraph features K, is highly variable, as shown in Fig. 2. This is a common problem 52 for all existing filter-based graph classification methods. 53

The above observations motivate the proposed research which **aims** to *integrate* subgraph mining, feature selection, and model training into one single framework (Fig. 1 (C)) with the optimal number of subgraphs K being automatically determined for graph classification. To achieve this goal, we formulate subgraph feature selection as the problem of learning a K-dimensional feature space from a **huge subgraph space** in order to result in minimum regularized loss on the training data as follows:

$$\min_{\boldsymbol{w}} \frac{1}{n} \sum_{i=1}^{n} \mathcal{L}(\boldsymbol{y}_i, f(\boldsymbol{x}_i)) + \gamma R(\boldsymbol{w})$$
(1)

where $\{x_i, \dots, x_n\}$ are the vector representations of the training graphs, \mathcal{L} is a loss function measuring the difference between the prediction $f(x_i)$ and the true label y_i , and R(w) is a regularization term on parameters w to avoid over-fitting.

Indeed, the optimization in Eq.(1) has been widely studied [20, 21, 22] in machine
learning community, but mainly for data with vector format. Several significant challenges remain for graph data:

1. Implicit Subgraph Features: For graph classification, no subgraph features are

readily available (*i.e.*, x_i is unknown) for training the model in Eq.(1). Instead, the feature space used to represent graphs is implicit and needs to be discovered by subgraph mining procedure as needed.

K-dimensional Features from Huge Subgraph Space: The number of sub graph candidates representing graphs is exponentially large. Finding an optimal
 number of *K* subgraphs for different graph datasets (in order to result in best
 classifiers), is crucial but has not been addressed by existing research.

In this paper, we propose a *unified* regularized loss minimization-driven (RLMD) graph 74 classification framework. Our theme is to progressively select the most discriminative 75 subgraph features from the training data in order to achieve minimum regularized loss 76 for a well defined objective function. To integrate subgraph selection into the model 77 learning process (Challenge 1), we formulate an objective function and design a sub-78 gradient method to induce a measurement to assess the utility of each subgraph, so that 79 the best subgraph features can be identified and incrementally included to optimize 80 the objective function for maximum performance gain. To determine the optimal K81 value for each dataset (Challenge 2), we use an *elastic net* [21] and derive a stopping 82 condition, so that the K value can be automatically obtained when the algorithm con-83 verges. By using the automatically determined optimal K value, as shown in Fig. 2, 84 RLMD finds 180 best subgraphs and achieves the best performance, which is 6% more 85 accurate than the second best method. 86

⁸⁷ The main contributions of this paper are summarized as follows:

We propose an *embedded* and *theoretically convergent* graph classification algorithm, which can automatically determine the optimal number of subgraphs *K* for graph classification. This is a *unified* approach in the sense that (1) it can employ any differentiable loss function (including least squares, exponential, and logistic loss functions) for graph classification; and (2) it integrates subgraph mining, feature selection, and model learning into one single framework.

• We generalize the *column generation technique* of gBoost [23] for graph classification, and demonstrate that gBoost [23] is a special case of our loss minimization algorithm.

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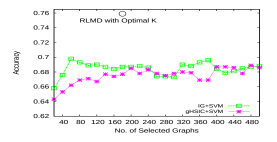


Figure 2: Classification accuracy for filter subgraph-based methods *w.r.t.* different numbers of subgraphs on the NCI-1 chemical compound dataset. IG is a TFM method which uses information gain to select subgraphs, whereas gHSIC [12] is a DFM method. All methods use SVM as a base classifier. The optimal number of subgraph features K is crucial, but difficult to decide for filter methods. In comparison, the proposed method (RLMD) automatically finds 180 best subgraphs and achieves the highest accuracy, which is 6% more accurate than the second best method.

- We propose the use of *elastic net* (which integrates two sparsity-inducing regularization norms, l₁-norm and l₂-norm) to produce a sparse and robust solution for discriminative subgraph selection.
- We derive a branch-and-bound rule according to the subgradient of our objective function to prune search space for optimal subgraph mining.
- Experimental results show that our algorithm RLMD outperforms two-step filter methods (TFMs), direct filter methods (DFMs), and gBoost algorithm with significant performance gain.

105 2. Related Work

¹⁰⁶ Our work is closely related to graph-based learning and graph classification.

107 2.1. Graph-based learning

Learning from data with dependency structures has been commonly addressed by existing research. Instead of considering samples as *I.I.D* observations, graph-based learning takes the relationships/correlations between samples to ensure effective learning. For example, graph-based approaches have been popularly used to propagate labels in semi-supervised learning [24, 25, 26], where training samples are connected through one or multiple graphs. A recent method [27] considers preserving global and

local structures inside the training data for feature selection. For large scale networks, 114 predicting linkage relationships between nodes (i.e. link prediction) can be used for 115 friendship recommendation in social networks [28], or suggesting potential interac-116 tions between proteins in bioinformatics research. A recent work [29] proposed to use 117 latent feature kernels to support link prediction on sparse graphs. All above methods 118 consider a large scale network with thousands (or millions) of integer-connected nodes 119 in the network. In contrast, we consider small graph classification problem, in which 120 each graph has a label indicating the property of the graph, and the graph normally 121 contains tens or several hundreds of nodes. The purpose is to predict the label of the 122 graph by using node and structure information inside the graphs, for purposes such 123 as chemical compound activity prediction [1] and gender classification using magnetic 124 resonance connectome (*i.e.* brain-graph) [2]. 125

126 2.2. Graph Classification

Existing methods for graph classification [18, 10, 9, 11, 12, 13, 14, 15, 16, 17, 23, 30, 31] can be roughly categorized into two groups: similarity-based methods and vector representation-based methods.

130 2.2.1. Similarity-based methods

These approaches aim to directly learn global similarities between graphs by using graph kernels [9, 32, 33, 34] or graph embedding [35]. Global similarities are then fed to similarity-based classifiers, such as KNN or SVM, for learning. One clear drawback of global similarity-based approaches is that the similarity is calculated based on global graph structures, such as random walks or embedding space. Therefore, it is not clear which substructures are more important for classifying graphs into different classes.

137 2.2.2. Vector representation-based methods

Another branch of methods transfer graphs into vector representations in structure space or in Euclidean space. In structure space [7, 8], geometrical and analytical concepts such as the angle between structures and the derivatives of functions on structures can be obtained, so that the structural pattern recognition problems can be formulated as optimization problems with certain cost functions. In Euclidean space, the goal is to transfer graphs into vector representations in Euclidean space so existing analytical
techniques can be applied for data analysis. Methods in this category are mainly filterbased approaches, including two-step filter methods (TFMs) or direct filter methods
(DFMs).

TFMs are straightforward approaches for graph classification which simply de-147 compose frequent subgraph generation and selection as two separated steps. An early 148 work [9] has shown that learning an SVM classifier based on the discovered frequent 149 subgraphs can achieve reasonably good accuracy for graph classification. On the other 150 hand, research [16, 15] also indicates that TFM methods may result in a bottleneck 151 for the subsequent feature selection module. Specifically, the number of frequent sub-152 graphs will grow exponentially if the minimum support threshold is low, which imposes 153 a great challenge for the subsequent feature selection task. This challenge has moti-154 vated many direct filter methods (DFMs), which seek to integrate subgraph discovery 155 and feature selection into one step. 156

For DFMs (a review on this category can be found in [18]), a key issue is to 157 define a proper measurement to assess the utility of each subgraph. Yan et al [17] pro-158 posed a LEAP algorithm to exploit the correlation between structural similarity and 159 significance similarity, so that a branch-and-bound rule can be derived to prune out 160 unpromising searching space efficiently. Ranu and Singh [16] proposed a scalable 161 GraphSig algorithm, which is able to mine significant subgraphs with low frequencies. 162 Thoma et al. [15] propose a CORK algorithm to find subgraph features. Recently, re-163 searchers have extended DFM to other graph applications, and have proposed effective 164 algorithms such as gSemi [11] for the semi-supervised setting, gCGVFL [36] for multi-165 view learning, gHSIC [12] for multi-label classification, and our recent multi-graph 166 classification for classifying graph bags, each containing multiple graphs [37, 38]. 167

Although filter methods for graph classification have been extensively studied, they all suffer from two major disadvantages: (1) the feature selection is not linked to the model learning process. As a result, the selected subgraph features may not best fit the underlying learning algorithms; and (2) the optimal number of subgraphs K for graph classification is difficult to decide and often varies from dataset to dataset, and inappropriately specified K value often results in significantly reduced classification accuracy. This is the common drawback for filter-based methods [19].

Embedded Methods. Our algorithm belongs to the embedded approach which inte-175 grates the subgraph selection into the model training process. In this subcategory, Saigo 176 et al [23] proposed a gBoost (its variants for imbalanced graph classification [39, 40] 177 and cost-sensitive learning [41] are proposed recently) algorithm which formulates the 178 graph classification as a linear program. As will be elaborated in Sec 4.6, our algorithm 179 is more general in the sense that it can adopt any differentiable loss function and use 180 more robust regularization to produce better performance. In fact, gBoost [23] can be 181 considered as a special case of our loss minimization problem. 182

3. Problem Definition

Definition 1. Connected Graph: A graph is denoted by $G = (\mathcal{V}, E, L = \{L_1, L_2\}, \mathcal{A} = \{\mathcal{A}_1, \mathcal{A}_2\})$, where \mathcal{V} is the vertex set, $E \subseteq \mathcal{V} \times \mathcal{V}$ is the edge set, $\mathcal{A} = \{\mathcal{A}_1, \mathcal{A}_2\}$ with \mathcal{A}_1 and \mathcal{A}_2 being the set of labels for vertices and edges, respectively; and $L = \{L_1, L_2\}$, $L_1 : \mathcal{V} \to \mathcal{A}_1, L_2 : E \to \mathcal{A}_2$ are labeling functions that assigns labels to a node or an edge, respectively. A connected graph is a graph such that there is a path between any pair of vertices.

In this paper, we focus on connected graphs and assume that each graph *G* has a class label $y, y \in \mathcal{Y} = \{-1, +1\}$. We only focus on binary-class classification tasks, but our methods can be easily extended to multi-class tasks.

Definition 2. Subgraph: Given two graphs $G = (\mathcal{V}, E, L = \{L_1, L_2\}, \mathcal{A} = \{\mathcal{A}_1, \mathcal{A}_2\})$ and $g_k = (\mathcal{V}', E', L' = \{L'_1, L'_2\}, \mathcal{A}' = \{\mathcal{A}'_1, \mathcal{A}'_2\})$, g_k is a subgraph of G (i.e., $g_k \subseteq G$) if there is an injective function $\hat{f}: \mathcal{V}' \to \mathcal{V}$, such that $\forall (a, b) \in E'$, we have $(\hat{f}(a), \hat{f}(b)) \in$ $E, L'_1(a) = L_1(\hat{f}(a)), L'_1(b) = L_1(\hat{f}(b)), L'_2(a, b) = L_2(\hat{f}(a), \hat{f}(b))$. If g_k is a subgraph of G ($g_k \subseteq G$), G is a supergraph of g_k ($G \supseteq g_k$).

Subgraph-based Graph Classification: Given a set of labeled graphs $\mathcal{T} = \{(G_1, y_1), \cdots, (G_n, y_n)\}$, subgraph-based graph classification **aims** to select an optimal set of discriminative subgraphs \mathcal{F}_1 from \mathcal{T} , and learn a classification model from the reduced subgraph space \mathcal{F}_1 to predict previously unseen test graphs with a maximum accuracy. Set \mathcal{F}_1 is optimal if the classifier learned from \mathcal{F}_1 has the highest classification accuracy, ²⁰³ compared to classifiers trained from any subset of \mathcal{T} . A major feature of our method is ²⁰⁴ that it can automatically determine the best set of subgraph features to represent each ²⁰⁵ graph datasets without requiring users to specify the number of subgraph features. This ²⁰⁶ essentially advances the existing subgraph feature-based graph classification methods ²⁰⁷ from finding the most discriminative subgraph features to finding the best subgraph set ²⁰⁸ for maximum accuracy gain.

4. Regularized Loss Minimization for Graph Classification

To support graph classification, state-of-the-art algorithms [23, 10] use a set of subgraphs discovered from the training graphs as features, where each subgraph g_k can be used to represent a graph G_i as follows:

$$\hbar_{g_k}(G_i) = 2I(g_k \subseteq G_i) - 1; \tag{2}$$

Here I(a) = 1 if *a* holds, and 0 otherwise. This rule simply maps a graph G_i into +1 if $g_k \subseteq G_i$, or -1 otherwise.

Let $\mathcal{F} = \{g_1, \dots, g_m\}$ be the full set of subgraphs for the training graphs. We can use \mathcal{F} as features to represent each graph G_i into a vector space as $\mathbf{x}_i = \{\hbar_{g_1}(G_i), \dots, \hbar_{g_m}(G_i)\}$, with $\mathbf{x}_i^k = \hbar_{g_k}(G_i)$. In the following subsection, G_i and \mathbf{x}_i are used interchangeably as they both refer to the same graph. Given the full subgraph features \mathcal{F} , the prediction function for the graph \mathbf{x}_i is a linear classifier:

$$f(\mathbf{x}_i) = \mathbf{x}_i \cdot \mathbf{w} + b = \sum_{g_k \in \mathcal{F}} w_k \hbar_{g_k}(G_i) + b$$
(3)

where $w = [w_1, \dots, w_m]'$ is the weight vector for all features \mathcal{F} , and *b* is the bias of the model. The predicted class of x_i is +1 if $f(x_i) > 0$ or -1 otherwise. Note that in practice, subgraph space \mathcal{F} is **implicit** and exponentially **large**, *i.e.*, the number of subgraphs grows exponentially with respect to the number of nodes.

224 4.1. Regularized Loss Minimization Formulation

In this paper, we propose to learn a *K*-dimensional feature space from the *implicit* and *large* subgraph space \mathcal{F} to achieve the lowest regularized empirical risks for the graph dataset, with *K* being automatically determined. Eq.(1) can be reformulated as the following objective function:

$$\min \mathcal{J}(\boldsymbol{w}, b) = \min_{\boldsymbol{w}, b} \underbrace{\frac{1}{n} \sum_{i=1}^{n} \mathcal{L}(y_i, \boldsymbol{x}_i \cdot \boldsymbol{w} + b)}_{C} + \underbrace{\gamma_1 \|\boldsymbol{w}\|_1 + \gamma_2 \|\boldsymbol{w}\|_2^2}_{R}$$
(4)

The first term C measures the loss on the training graphs, where $\mathcal{L}(y_i, f(x_i))$ can be any 229 loss function measuring the misclassification penalty of a graph G_i . The second part R 230 consists of regularization terms to enforce sparse and robust solutions. Parameters γ_1 231 and γ_2 are used to trade-off these parts ($\gamma_1 \ge 0, \gamma_2 \ge 0$). For the regularization, our 232 objective is to obtain a sparse and stable solution on w, i.e., low dimensional subgraph 233 features for final graph classification. Here, we combine both ℓ_1 and ℓ_2 norm, which is 234 known as *elastic net* in machine learning [21]. The motivation of our regularization is 235 as follows: 236

The ℓ_1 -norm regularizer $(\sum_k |w_k|)$ can produce solutions with many coefficients be-237 ing 0, which is known as lasso [20] and has been widely applied for variable selections. 238 Although ℓ_1 regularization can produce a sparse solution, it suffers from two major 239 disadvantages: (1) the number of selected variables is limited by the number of obser-240 vations; and (2) the lasso penalized model can only select one variable from a group 241 of correlated variables and does not care which one is selected [21]. In contrast, ℓ_2 242 regularization, which is widely used in SVM formulation (($||\boldsymbol{w}||_2^2 = \sum_{k=1}^m |w_k|^2$)), can 243 produce more stable and robust classification results. However, ℓ_2 formulation cannot 244 produce a sparse solution. By combining ℓ_1 and ℓ_2 norm, known as elastic net [21], we 245 can overcome these issues and enjoy the sparse and stable properties. 246

247 4.2. Sparse Subgraph Learning: Challenges and Solution Overview

Challenges: For explicit vector data with moderate feature size, the problem defined in Eq.(4) can be effectively solved in traditional supervised learning. However, for graph data the challenges are evident: (1) the feature set \mathcal{F} is unavailable (**implicit**) unless we enumerate all subgraphs from the training graphs, which is NP-complete; and (2) the whole subgraph set is exponentially **large**, and only a small subset of subgraphs are useful for classifiers to achieve maximum graph classification accuracy.

Solution Overview: To solve the aforementioned challenges, we propose a regularized 254 loss minimization-driven (RLMD) subgraph selection method for graph classification. 255 Driven by our formulation in Eq.(4), our principle is to iteratively mine the best sub-256 graph feature to reduce the empirical loss on the training graphs. To this end, we resort 257 to the subgradient method in the functional space to define the utility of each sub-258 graph, and embed the feature selection/ranking into the subgraph mining/enumeration 259 process. To handle the exponentially large subgraph space, we derive an effective 260 branch-and-bound pruning scheme to reduce the search space. After a new subgraph 261 is selected, we include and re-solve the new restricted objective function of Eq.(4) by 262 using currently selected subgraphs. To find optimal K value, we derive a stopping cri-263 terion for our feature selection procedure based on the subgradient in the functional 264 space, so that K can be automatically obtained once the algorithm converges. 265

Logistic Loss Function: A Running Example. Our method is based on the gradient/subgradient on the functional space of the objective function Eq.(4). In this paper, we use the logistic loss function as an example to illustrate how subgraph selection is performed by subgradient methods, and the logistic loss function is given as follows:

$$\mathcal{L}(y_i, f(x_i)) = \log\left(1 + \exp\left\{-y_i f(\boldsymbol{x}_i)\right\}\right)$$
(5)

Note that our algorithm is a general method in the sense that any other differentiable loss function, such as least square loss $\mathcal{L}(y, f_t) = \frac{1}{2}(y-f_t)^2$ or exponential loss $\mathcal{L}(y, f_t) = \exp\{-yf_t\}$, can be directly used in our algorithm. As discussed latter in Section 4.6, our method is also applicable to convex locally Lipschitz but non-differentiable loss functions such as the hinge loss used by the (margin) perceptron and linear SVM.

The partial derivative of the loss term *C* in Eq.(4) on the subgraph feature g_k is defined as $\frac{\partial C}{\partial w_k}$. For logistic loss function,

$$\frac{\partial C}{\partial w_k} = \frac{1}{n} \sum_{i=1}^n \frac{\partial \mathcal{L}(y_i, f(\mathbf{x}_i))}{\partial f(\mathbf{x}_i)} \frac{\partial f(\mathbf{x}_i)}{\partial w_k}$$

$$= -\frac{1}{n} \sum_{i=1}^n \frac{y_i \mathbf{x}_i^k}{1 + e^{y_i f(\mathbf{x}_i)}} = \sum_{i=1}^n y_i \alpha_i \mathbf{x}_i^k$$
(6)

Here, $\alpha_i = -\frac{1}{n(1+e^{y_i/(x_i)})}$ can be regarded as a weight associated with graph G_i for the subgraph mining process.

279 4.3. RLMD Subgraph Selection for Graph Classification

Because we aim to learn a sparse solution of subgraph features (K-dimensional 280 feature space) from graph data, some subgraphs/features g_k with zero weights, *i.e.*, 281 $w_k = 0$ will not be used for learning the classification model. Thus it makes sense 282 to partition the subgraph features \mathcal{F} into two disjoint subsets \mathcal{F}_1 and \mathcal{F}_2 . \mathcal{F}_1 stores 283 active features which are used to learn the classification model and this set is frequently 284 updated as desired, and \mathcal{F}_2 includes unselected graphs with 0 weights (*i.e.*, for $g_k \in$ 285 $\mathcal{F}_2, w_k = 0$). Then we can iteratively select the best feature from \mathcal{F}_2 to \mathcal{F}_1 , and solve 286 the following restricted subproblem: 287

$$\min \mathcal{J}_{t}(\boldsymbol{w}, b) = \min_{\boldsymbol{w}, b} \underbrace{\frac{1}{n} \sum_{i=1}^{n} \mathcal{L}(y_{i}, \boldsymbol{x}_{i}(t) \cdot \boldsymbol{w} + b)}_{C'} + \underbrace{\gamma_{1} ||\boldsymbol{w}||_{1} + \gamma_{2} ||\boldsymbol{w}||_{2}^{2}}_{R'} \qquad (7)$$
$$= \mathcal{J}_{t}(\boldsymbol{w}(t), b(t))$$

where $x_i(t)$ is the feature representations for graph G_i based on the active set \mathcal{F}_1 in the t-th iteration, and (w(t), b(t)) is the solution of Eq.(7). Note that $\mathcal{J}_t(w, b)$ is used here to denote the restricted subproblem Eq.(7) while $\mathcal{J}(w, b)$ is referred to original problem in Eq.(4).

The optimal number of subgraphs *K* can be automatically determined by setting $K = |\mathcal{F}_1|$ once the algorithm converges. Note that given a solution (w(t), b(t)), the loss term *C'* in Eq.(7) equals to *C* in Eq.(4) because the prediction of of graph mainly depends on the active set \mathcal{F}_1 , *i.e.*, C' = C. In the following, we will derive the stopping condition of our algorithm, and prove its convergence.

Stopping Condition for Optimal *K* value: Our objective function Eq.(4) is convex and non-smooth, *i.e.*, it may be non-differentiable at a point *w*. When it is non-differentiable at *w*, we can compute its generalized gradient (*i.e.*, subgradient) instead. According to the optimization conditions, when reaching the optimum, we will have

$$0 \in \frac{\partial C}{\partial w_k} + \gamma_1 o_k + 2\gamma_2 w_k; \tag{8}$$

³⁰¹ where o_k is the subgradient with respect to w_k

$$o_k \in \begin{cases} \operatorname{sign}(w_k) : w_k \neq 0 \\ [-1,1] : w_k = 0 \end{cases}$$
(9)

where sign(a) = 1 if a > 0 otherwise -1.

According to Eq.(8) and Eq.(9), we can now state the optimal condition for our sparse subgraph learning problem.

Proposition 1. Optimal Solution: Let $\hat{w} = [\hat{w}_1, \dots, \hat{w}_m]$. Suppose that (\hat{w}, \hat{b}) is the optimal solution of our objective function Eq.(4), then Eq.(10) and Eq.(11) hold.

$$\frac{\partial C}{\partial \hat{w}_k} + \gamma_1 sign(\hat{w}_k) + 2\gamma_2 \hat{w}_k = 0 \quad \text{if} \quad \hat{w}_k \neq 0 \tag{10}$$

$$\left|\frac{\partial C}{\partial \hat{w}_k}\right| \le \gamma_1 \quad \text{if} \quad \hat{w}_k = 0 \tag{11}$$

Eq.(11) holds because for $\hat{w}_k = 0$, the third term of Eq.(8) disappears. Combining Eq.(8) and Eq.(9) will result in Eq.(11).

To reduce the objective value \mathcal{J}_t in Eq.(7), we propose to select a subgraph in \mathcal{F}_2 whose weight violates Eq.(11), and update the selected active set \mathcal{F}_1 with the newly selected feature and re-optimize the restricted subproblem Eq.(7) with current features. This process will repeat until no candidate violates Eq.(11). In other words, Eq.(11) is a stopping condition and determines the number of subgraphs being selected for RLMD's subgraph selection process. Utility of Subgraphs: Eq.(11) can be used naturally to induce a criterion for quantify-

ing the utility value of a subgraph. The larger $|\frac{\partial C}{\partial w_k}|$ is, the more informative it will be for reducing the objective function. Accordingly, we formally define the informative score as follows:

Definition 3. *Informative Score:* For a subgraph pattern g_k , its informative score for graph classification is defined as follows:

$$\Theta(g_k) = \left|\frac{\partial C}{\partial w_k}\right| = \left|\frac{\partial C'}{\partial w_k}\right| = \left|\sum_{i=1}^n y_i \alpha_i \boldsymbol{x}_i^k\right|$$
(12)

321 where $\alpha_i = -\frac{1}{n(1+e^{y_i f(x_i)})}$.

Note that the informative score directly depends on the weight of each graph α_i , which is calculated based on the active set \mathcal{F}_1 . Intuitively, the best subgraph of \mathcal{F}_2 is the one with the maximum informative score, because it is more likely to violate the stopping condition Eq.(11).

Algorithm 1 Regularized Loss Minimization-Driven Subgraph Selection (RLMD) for

	ph Classification
Req	uire:
ł	$\{(G_1, y_1), \cdots, (G_n, y_n)\}$: Training Graphs;
	S _{max} : Maximum number of iterations;
Ensi	ire:
1	w, b: Parameters for classifier model
1: 6	$\alpha_i = 1/n; \ \mathcal{F}_1 \leftarrow \emptyset; \ t \leftarrow 0;$
2:	while $t < S_{max}$ do
3:	Mine an optimal subgraph features g^* with maximum informative score defined by Eq.12; //Algo
	rithm 2;
4:	if $\Theta(g^{\star}) \leq \gamma_1 + \varepsilon$ then
5:	break;
6:	end if
7:	$\mathcal{F}_1 \leftarrow \mathcal{F}_1 \bigcup g^{\star};$
8:	Solve Eq.(7) based on \mathcal{F}_1 to get the new solution ($w(t), b(t)$);
9:	Update the graph weights on each training graph
	$\alpha_i = -\frac{1}{n(1+e^{y_i f(x_i(t))})}$
10:	$t \leftarrow t + 1;$
11:	end while
12:	$K = \mathcal{F}_1 ;$
13:	return w, b;

RLMD Algorithm: Algorithm 1 illustrates the detailed steps of RLMD for graph classification. Initially, the weights for all training graphs are equally set as 1/n, and the active set \mathcal{F}_1 is initialized to be empty.

In the next step, the algorithm mines an optimal subgraph g^* from \mathcal{F}_2 which has the highest informative scores defined by Eq.(12). This step involves a subgraph mining procedure, which will be addressed in the next subsection. On steps 4-5, if current optimal subgraph no longer violates the optimal condition Eq.(11), the algorithm terminates. Here, we have relaxed the convergence condition to ε tolerance; this is because in the last few iterations, the maximum score will only change subtly (we set $\epsilon = 0.005$ in our experiments).

On step 7, we add the newly selected subgraph g^* to the existing subgraph set \mathcal{F}_1 , and re-solve the following restricted subproblem Eq.(7). To solve this restricted

objective function, we use the MALSAR toolbox ² in our experiments. It is worth noting that because this step only involves a very small number of features, it is very efficient in practice.

Subsequently, the algorithm updates the weight α_i for each graph G_i . This will help compute the derivative of $\frac{\partial C}{\partial w_k}$ for subgraph mining in next round. After the algorithm terminates, the optimal number of subgraphs *K* can be easily obtained as $K = |\mathcal{F}_1| = t$ on step 12.

Note that our algorithm 1 generalizes the column generation technique in gBoost [23] by iteratively selecting the most violated subgraph in each iteration until convergence. Our algorithm 1 relies on ε and γ_1 , which serve as a stopping condition to determine *K*. In practice, ε is a subtle value insensitive to the algorithm performance. Meanwhile, γ_1 is much easier to set than asking users to specify *K* values because γ_1 is chosen in a

much smaller range, as we will demonstrate in Section 5.2.4.

351 4.4. Theoretical Study

350

Theorem 1. (*Convergence Property:*) Algorithm 1 guarantees that the restricted objective function Eq.(7) will monotonically decrease.

PROOF. Suppose in the t-th iteration, the optimal objective value based on current t features (i.e., $|\mathcal{F}_1| = t$) is obtained with (w(t), b(t)), i.e.,

$$\mathcal{J}_{t}(\boldsymbol{w}(t), b(t)) = \underbrace{\frac{1}{n} \sum_{i=1}^{n} \mathcal{L}(y_{i}, \boldsymbol{x}_{i}(t) \cdot \boldsymbol{w}(t) + b(t))}_{C = C'} + \underbrace{\gamma_{1} ||\boldsymbol{w}(t)||_{1} + \gamma_{2} ||\boldsymbol{w}(t)||_{2}^{2}}_{\mathcal{R} = \mathcal{R}'} = (C + \mathcal{R})_{|(\boldsymbol{w}(t), b(t))|}$$

then in the t + 1-th iterations, the optimal objective value for Eq.(7) is

$$\min \mathcal{J}_{t+1}(\boldsymbol{w}, b) = \min(C + \mathcal{R}) \le (C + \mathcal{R})_{|(\boldsymbol{w}(t), 0|, b(t))} = \mathcal{J}_t(\boldsymbol{w}(t), b(t))$$

Here [w(t), 0] means that the weights for subgraphs selected in the t-th iteration remain

unchanged while the weight for newly selected subgraph in the t+1-th iteration is 0.

²http://www.MALSAR.org

Thus the objective value of the restricted problem Eq.(7) based on the currently selected features \mathcal{F}_1 always monotonously decreases in two successive iterations. Because the objective function value is non-negative (bounded), we can ensure that it will finally converge as iteration continues. The proof is complete.

Suppose the algorithm converges in the *K*-th iteration with a solution (w(K), b(K)), and the objective value for Eq.(7) is $\mathcal{J}_K(w(K), b(K))$. By adding m - K zeros for subgraphs in \mathcal{F}_2 to w(K), *i.e.*, $\hat{w}(K) = [w(K), 0 \cdots]$, we obtain a solution $(\hat{w}(K), b(K))$ for Eq.(4).

Corollary 1. *Optimal Solution Guarantees:* If Algorithm 1 converges with solution (w(K), b(K)) for Eq.(7), then ($\hat{w}(K), b(K)$) is an optimal solution for Eq.(4).

PROOF. According to our Proposition 1, $(\hat{w}(K), b(K))$ is an optimal solution of Eq.(4), because $\forall w_k = 0 \ (g_k \in \mathcal{F}_2)$, we have $\Theta(g_k) < \gamma_1$ based on our stopping condition. Thus we will have

 $\mathcal{J}_K(\boldsymbol{w}(K), \boldsymbol{b}(K)) = \mathcal{J}(\hat{\boldsymbol{w}}(K), \boldsymbol{b}(K)) = \min \mathcal{J}(\boldsymbol{w}, \boldsymbol{b})$

where $J_K(w(K), b(K))$ and $\mathcal{J}(\hat{w}(K), b(K))$ refer to the objective values of the restricted subproblem Eq.(7) and Eq.(4), respectively.

We have proved that objective value for Eq.(7) is monotonously decreasing (Theorem 1) and its recovered solution ($\hat{w}(K), b(K)$) is an optimal solution to Eq.(4) (Corollary 1).

371 4.5. Optimal Subgraph Mining

In order to mine optimal subgraph g^{\star} on step 3 of Algorithm 1, we need to perform the subgraph enumeration procedure. In RLMD, we employ the frequent subgraph mining-based algorithm gSpan [42]. The key idea of gSpan is that each subgraph has a unique DFS Code, which is defined by a lexicographic order of the discovery time during the search process. By employing a depth first search strategy on the DFS Code tree (where each node is a subgraph), gSpan can enumerate all frequent subgraphs efficiently. During the subgraph mining process, the search space is exponentially large, which requires an effective pruning scheme to reduce the search space. In this subsection, we will derive the upper-bound of the informative score for each subgraph, which helps prune the search space and speed up the subgraph mining.

Theorem 2. (*Upper-bound Score:*) Let g and g' be two subgraph patterns, and $g \subseteq g'$, for the subgraph g, we define

$$A_1(g) = 2 \sum_{\{i \mid y_i = +1, g \in G_i\}} \alpha_i$$

$$A_2(g) = 2 \sum_{\{i \mid y_i = -1, g \in G_i\}} \alpha_i$$

$$A_3 = \sum_{i=1}^n \alpha_i y_i$$

385

$$\hat{\Theta}(g) = \begin{cases} \max\{|A_1(g) - A_3|, |A_2(g)|\} & : \quad A_3 \ge 0\\ \max\{|A_2(g) + A_3|, |A_1(g)|\} & : \quad A_3 < 0 \end{cases}$$

then $\Theta(g') \leq \hat{\Theta}(g)$, where $\Theta(g')$ is defined in Eq.(12).

³⁸⁷ **PROOF**. We start with the definition of $\Theta(g')$:

$$\begin{split} \Theta(g') &= |\sum_{i=1}^{n} y_{i} \alpha_{i} \mathbf{x}_{i}'| \\ &= |\sum_{i=1}^{n} y_{i} \alpha_{i} \cdot [2I(g' \subseteq G_{i}) - 1]| \\ &= |2 \sum_{g' \subseteq G_{i}} y_{i} \alpha_{i} - \sum_{i=1}^{n} \alpha_{i} y_{i}| \\ &= |A_{1}(g') - A_{2}(g') - A_{3}| \\ &\leq \begin{cases} \max\{|A_{1}(g') - A_{3}|, |A_{2}(g')|\} : A_{3} \ge 0 \\ \max\{|A_{2}(g') + A_{3}|, |A_{1}(g')|\} : A_{3} < 0 \\ \\ &\leq \begin{cases} \max\{|A_{1}(g) - A_{3}|, |A_{2}(g)|\} : A_{3} \ge 0 \\ \max\{|A_{2}(g) + A_{3}|, |A_{1}(g)|\} : A_{3} \ge 0 \\ \\ &\max\{|A_{2}(g) + A_{3}|, |A_{1}(g)|\} : A_{3} < 0 \\ \\ &= \hat{\Theta}(g) \end{split}$$

The first inequality holds because for $\alpha_i < 0$, $A_1(g') \le 0$ and $A_2(g') \le 0$, so the upperbound depends on A_3 . If $A_3 \ge 0$, $A_1(g')$ and A_3 will have different signs, then the upper-bound is the maximum between $\{|A_1(g') - A_3|, |A_2(g')|\}$. The case is similar for $A_3 < 0$. The second inequality holds because $|A_1(g')| \le |A_1(g)|$ and $|A_2(g')| \le |A_2(g)|$ for $g \subseteq g'$.

Algorithm 2 Optimal Subgraph Mining

Require:

 $\{(G_1, y_1), \cdots, (G_n, y_n)\}$: Training graphs;

- α_i : Weight for each graph example;
- \mathcal{F}_1 : Already selected subgraph set;

Ensure:

 g^* : The optimal subgraph;

- 1: $\eta = 0;$
- 2: while Recursively visit the DFS Code Tree in gSpan do
- 3: $g_p \leftarrow$ current visited subgraph in DFS Code Tree;
- 4: **if** g_p has been examined **then**
- 5: continue;
- 6: end if
- 7: Compute scores $\Theta(g_p)$ for subgraph g_p according Eq.(12);
- 8: **if** $g_p \notin \mathcal{F}_1$ & $\Theta(g_p) > \eta$ then
- 9: $\eta = \Theta(g_p);$
- 10: $g^{\star} \leftarrow g_p;$
- 11: **end if**
- 12: **if** $\hat{\Theta}(g_p) > \eta$ **then**
- 13: Depth-first search the subtree rooted from node g_p ;
- 14: end if
- 15: end while
- 16: **return** g^* ;
- Theorem 2 states that for any super graph of a subgraph g, its informative score is upper-bounded by $\hat{\Theta}(g)$. This rule can prune unpromising candidates effectively.

Optimal Subgraph Exploration Algorithm: Our optimal subgraph mining algorithm is listed in Algorithm 2. The minimum value η in the optimal set is initialized on step 1. Duplicated subgraph features are pruned on steps 4-5, and the informative score $\Theta(g_p)$ for g_p is calculated on step 7. If g_p is not selected before $(g_p \notin \mathcal{F}_1)$ and $\Theta(g_p)$ is larger than η , we replace the optimal subgraph g^* with the current g_p and update the optimal score η (steps 8-11).

⁴⁰¹ A branch-and-bound pruning rule, according to Theorem 2, is subsequently used to ⁴⁰² prune the search space on step 12. Lastly, the optimal subgraph g^* is returned on step ⁴⁰³ 16. The above pruning process is a key feature of our algorithm, because we do not require a support threshold value for subgraph mining (whereas all filter subgraph mining methods will require users to predefine a threshold value in order to discover subgraphs).

408 4.6. Relation to gBoost

Our RLMD subgraph selection algorithm advances the existing *column generation* style techniques employed in gBoost [23] for graph classification. The learning objective function for gBoost is

$$\max_{\rho, w, \xi} \quad \rho - \frac{1}{v_n} \sum_{i=1}^n \xi_i$$
s.t.
$$y_i \sum_{k=1}^m h_{g_k}(G_i) w_k + \xi_i \ge \rho;$$

$$\sum_{k=1}^m w_k = 1;$$

$$w_k \ge 0, \xi_i \ge 0;$$
(13)

From [43], we know that this formula is equivalent to the following linear programming:

$$\min_{\substack{w,\xi \\ w,\xi \\ s.t. \\ w_k \ge 0, \xi_i \ge 0;}} \sum_{k=1}^m w_k + C \sum_{i=1}^n \xi_i (14)$$

Eq.(14) is actually a ℓ_1 svm formulation, and can also be formulated as the regularized loss minimization formulation problem:

$$\min \|\boldsymbol{w}\|_1 + C \sum_{i=1}^n \mathcal{L}_h(\mathbf{y}_i, f(\boldsymbol{x}_i))$$
(15)

Here, $\mathcal{L}_h(y_i, f(\mathbf{x}_i)) = \max(1 - y_i f(\mathbf{x}_i), 0)$, which is known as hinge loss in machine learning.

Compared to our objective function in Eq.(4), we can find that gBoost (Eq.15) is a special case of Eq.(4), with the ℓ_2 regularization term being 0. Although the hinge loss function is non-differentiable, our subgradient method still applies, as long as $\frac{\partial C}{\partial w_k}$ in Eq.(6) is properly defined. This observation shows the following advantages of our algorithm: (1) gBoost employs a hinge loss function which is similar to SVM and

requires the problem to be formulated as a linear programming. Our algorithm gen-423 eralizes and advances gBoost by removing the linear programming constraint and can 424 employ any differentiable loss function, in addition to the logistic loss function con-425 sidered in our paper. This generalization has great attractiveness in many applications, 426 especially when the probability estimation for classification is required (the logistic 427 function can provide some probabilistic information compared to the hinge loss func-428 tion); (2) while gBoost employs ℓ_1 norm regularization to obtain a sparse solution, our 429 algorithm considers an additional norm ℓ_2 . This combined norm (known as *elastic net*) 430 enables a sparse and more stable solution. 431

432 5. Experiment

433 5.1. Experimental Settings

Benchmark Data: We validate the performance of the proposed algorithm on two
types of graph classification datasets.

Anti-cancer activity prediction (NCI): The NCI graph collection³ is a benchmark 436 for predicting the biological activity of small molecules for different types of cancers. 437 Each NCI dataset belongs to a bioassay task for anticancer activity prediction, such 438 as Breast cancer or Leukemia cancer. Each molecule is represented as a graph, with 439 atoms representing nodes and bonds denoting edges. A molecule is positive if it is 440 active against a certain type of cancer, or negative otherwise. Table 1 summarizes nine 441 NCI graph classification tasks used in our experiments, where columns 2-4 denote the 442 number of positive molecules, the total number of graphs, and the type of cancer of 443 each dataset. In our experiments, we randomly select 1000 graphs from each dataset 444 with balanced class distributions for graph classification. 445

Predictive Toxicology Challenge Dataset (PTC): The PTC challenge includes a number of carcinogenicity classifications for the toxicology prediction of chemical compounds⁴. The dataset we selected contains 417 compounds with four types of test animals: MM (male mouse), FM (female mouse), MR (male rat), and FR (female

³http://pubchem.ncbi.nlm.nih.gov

⁴http://www.predictive-toxicology.org/ptc/

ID	#Pos	#Total	Learning tasks
1	1793	37349	Non-Small Cell Lung
33	1467	37022	Melanoma
41	1350	25336	Prostate
47	1735	37298	Central Nerv Sys
81	2081	37549	Colon
83	1959	25550	Breast
109	1773	37518	Ovarian
123	2715	36903	Leukemia
145	1641	37043	Renal

rat). Each compound has labels selected from {CE, SE, P, E, EE, IS, NE, N}. Similar
to [44], we set {CE, SE, P} as positive labels, and {NE,N} as negative labels.

Baseline Methods: In our experiments, we consider three types of baseline methods, namely the two-step filter methods (TFMs), direct filter methods (DFMs), and embed-

- ded methods, as follows:
- IG+SVM is a TFM method that simply mines a set of frequent subgraphs, and then performs feature selection by using Information Gain. A SVM classifier is trained by using selected subgraph features for graph classification.
- **TOP+SVM** is similar to IG+SVM except that it selects the top *K* subgraphs based on their frequency rather than their information gain values.
- gSemi+SVM [11] is a DFM method, which integrates the feature selection into
 the subgraph mining process. The measurement for feature selection mainly con siders the *must-link* and *cannot link* constraints between graph samples within
 the same or between different classes.
- **gHSIC+SVM** [12] is another DFM method which exploits the correlations between features and labels.
- **gBoost** [23] is a state-of-the-art embedded method which formulates the feature selection as a linear problem and selects subgraph features which best fit the

468 objective function.

• **RLMD** is our proposed method which employ a logistic loss function together with an *elastic net* for regularization, and automatically determines optimal number of subgraphs *K*.

We conduct 10-fold cross-validation on all graph datasets and report the average 472 results and standard errors of 10 folds in the final result. The parameters for γ_1 are 473 selected from $\{0.005, 0.01, 0.03, 0.05\}$, and γ_2 is selected from $\{0.01, 0.03, 0.05\}$. We 474 will further analyze the impact of γ_1 and γ_2 in wider ranges in Section 5.2.4. For the 475 filter methods (IG+SVM, TOP+SVM, gSemi+SVM, and gHSIC+SVM), the minimum 476 support for frequent subgraph mining is set to 10% on NCI graph datasets and 1% 477 on PTC classification tasks, and an SVM classifier is trained with C parameter from 478 the range $\{0.1, 1, 10, 100, 1000, 10000\}$. For the gBoost algorithm, the parameter v is 479 selected from $\{0.1, 0.2, 0.3, 0.4\}$. Following [23], we select the best average results 480 of 10-fold cross-validation for each baseline algorithm by varying these parameters, 481 which represents the best performance each baseline can achieve. 482

For fairness of comparison, we increase the number of features to be selected for 483 the filter methods (IG+SVM, TOP+SVM, gSemi+SVM, and gHSIC+SVM), and in-484 crease the iterations for the embedded methods (gBoost and RLMD), then collect and 485 compare the performance of all algorithms under the same number of features. We set 486 $S_{max} = 200$, which defines the maximum number of features used to learn the classifier 487 models. Note that for RLMD, the algorithm may stop before reaching the maximum 488 iterations/subgraphs we set, *i.e.*, the optimal K is obtained. When RLMD stops, the 489 optimal number of subgraph features has been discovered and RMLD will not add 490 additional subgraphs to the feature set. We also compare RLMD under the optimal 491 subgraph value to other baselines with the same number of K features (the purpose is 492 to show that the optimal subgraph features discovered by RLMD are indeed optimal 493 for graph classification). 494

495 5.2. Experimental Results

496 5.2.1. Results on NCI Graph Dataset

For the NCI graph datasets, we vary the number of selected subgraph features from
20 to 200 for filter methods, and the number of iterations for gBoost and RLMD from
1 to 200. The accuracies and AUC values are shown in Fig. 3.

Comparison with Filter methods: The results in Fig. 3 show that with the in-500 crease in the number of features/iterations, the filter methods (TOP+SVM, IG+SVM, 501 gSemi+SVM, and gHSIC+SVM) are inferior to RLMD. This is because filter methods 502 separate the feature selection module from the model learning process. The subgraph 503 features selected from filter methods may not fit the underlying learning model very 504 well (we use SVM in our experiments). This is actually an observed common draw-505 back of filter methods [19]. The performance among these filter-based methods varies 506 from one graph dataset to another, and none of them significantly outperforms others. 507 For instance, gSemi+SVM outperforms TOP+SVM, IG+SVM, and gHSIC+SVM on 508 NCI-1 (Fig. 3.A) when the number of selected graphs is considerably large (≥ 160), 509 but is worse than gHSIC on NCI-109 (Fig. 3.G). This may be attributed to the inherent 510 differences underneath the graph datasets. 511

How many subgraphs to select: Another drawback of filter methods, shown in our experiments, is that the performance of filter methods varies significantly *w.r.t.* different numbers of selected features (K). Indeed, all these filter methods only select subgraph features with maximum discriminative score regardless of the redundancy among the features. Adding redundant features may decrease the performance of an algorithm. Further analysis of subgraph features is presented with a case study in the next subsection.

In contrast, for embedded methods, the above drawbacks can be handled effectively. Our algorithm RLMD unifies the feature selection and model learning into a whole framework, so that the feature selection process is driven by the well-defined objective function, and the selected features can further enhance the learning models. At the same time, RLMD is guaranteed to be convergent given an appropriate γ_1 value, which means that we do not need to specify the total number of selected graphs *K*. For

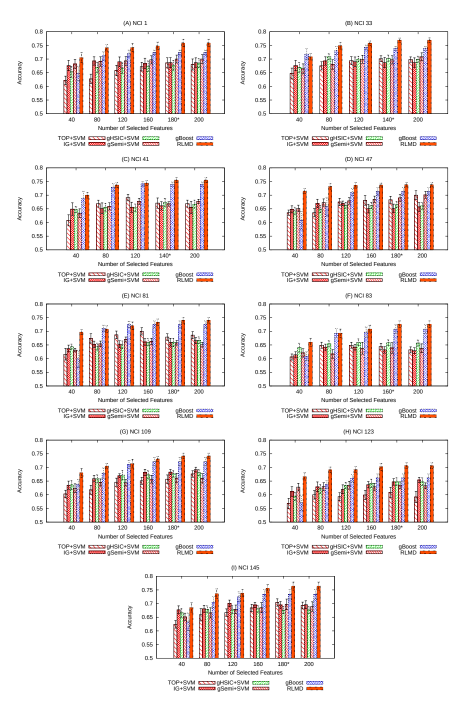


Figure 3: The classification accuracy and standard error on NCI dataset *w.r.t.* the number of selected graphs. The optimal number of subgraphs selected by RLMD (once it converges for all 10 folds experiments) is marked by a star * at *x*-axis.

instance, in Fig. (3).A, RLMD reaches convergence with 180 features.

RLMD vs. gBoost: It is evident that RLMD outperforms gBoost for most datasets. 526 This is mainly because gBoost only uses ℓ_1 -norm regularization to produce a sparse 527 solution. As pointed out in [21], the lasso (ℓ_1 -norm) has several drawbacks: (1) when 528 the number of features (m, which is exponentially huge) is much bigger than the num-529 ber of observations (n), the ℓ_1 norm selects at most n variables before it saturates; and 530 (2) when the pairwise correlations in a group of variables are very high, lasso tends to 531 select only one variable from the group and does not discriminate which one it selects. 532 By contrast, RLMD uses an elastic net (combination of ℓ_1 and ℓ_2 norm), which encour-533 ages a grouping effect, where strongly correlated features will be included/excluded. 534 As a result, RLMD results in a similar sparsity of representation to gBoost, but often 535 outperforms gBoost. 536

ID	RLMD	TOP +SVM	IG +SVM	gHSIC +SVM	gSemi +SVM	gBoost
1	0.759 ±0.014	0.686±0.019	0.687 ± 0.018	0.677 ± 0.018	0.700±0.015	0.724±0.009
33	0.769±0.008	0.701 ± 0.011	0.687 ± 0.019	0.703 ± 0.012	0.697 ± 0.015	$0.739{\scriptstyle \pm 0.008}$
41	0.755±0.009	0.677 ± 0.017	$0.649{\scriptstyle\pm0.015}$	0.672 ± 0.015	0.670 ± 0.009	0.740 ± 0.008
47	0.738±0.009	0.683±0.013	0.652 ± 0.015	0.665 ± 0.012	0.690 ± 0.012	0.714 ± 0.014
81	0.740 ±0.010	0.679 ± 0.012	0.660 ± 0.014	0.659 ± 0.015	0.658 ± 0.009	0.725 ± 0.013
83	0.726 ±0.012	0.644 ± 0.012	0.632 ± 0.012	0.658 ± 0.011	0.639 ± 0.020	0.708 ± 0.016
109	0.742±0.010	0.657 ± 0.016	0.683 ± 0.009	0.677 ± 0.012	0.661 ± 0.017	0.721±0.015
123	0.707 ±0.010	0.609 ± 0.019	0.647 ± 0.012	0.648 ± 0.014	0.635 ± 0.012	0.663 ± 0.013
145	0.764 ±0.015	0.704 ± 0.013	$0.695{\scriptstyle\pm0.012}$	0.676 ± 0.013	0.697 ± 0.020	$0.734{\scriptstyle\pm0.018}$

Table 2: Averaged accuracies and standard errors on NCI Graphs with Optimal K.

Overall Performance with Optimal *K*: In Table 2, we summarize the performance of our algorithm under optimal *K* value with other methods, where filter methods use the same number of subgraphs (*K*) for graph classification, and gBoost runs until convergence. The result in Table 2 clearly demonstrates that RLMD outperforms twostep filter methods (TOP+SVM and IG+SVM), direct filter methods (gHSIC+SVM and gSemi+SVM), and gBoost algorithm in NCI datasets.

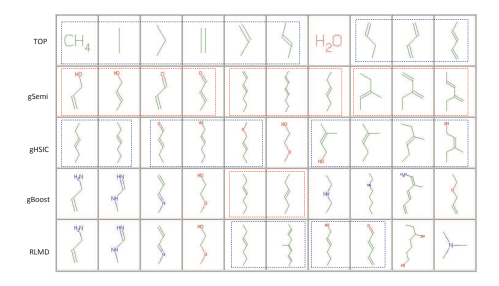


Figure 4: Case Study: comparison of discriminative subgraph features discovered by different algorithms. Subgraph features with high similarities are grouped and highlighted in the dashed rectangles. Subgraphs mined by filter methods are similar to each other and share high redundancy.

543 5.2.2. Case Study: Subgraph Feature Comparison

In this subsection, we use NCI-1 dataset as a case study to investigate subgraphs discovered by different algorithms. In our experiments, the top-10 subgraph features are discovered and illustrated in Fig. 4.

It is evident that the features for all filter methods (TOP, gSemi, and gHSIC) share 547 high correlations. For the gSemi algorithm, for instance, the top-10 subgraphs form 548 3 groups. In each group, the subgraph features are very similar to each other. This is 549 because the subgraph mining algorithm follows the depth-first-search (DFS) scheme, 550 and subgraphs from the same sub-tree are very close to each other in terms of their ge-551 ometrical structure. Because these methods consider each subgraph independently, the 552 selected subgraphs may have high redundancy, which imposes a great challenge in de-553 termining the optimal K subgraphs for graph classification and also causes fluctuating 554 results when the K values are varied. 555

In contrast, the subgraph correlations for gBoost and RLMD are much smaller.

The subgraphs discovered by gBoost and RLMD are highly overlapping (the first 5 subgraphs are identical). As pointed out by [21], ℓ_1 regularization tends to select only one subgraph from a group of features and is not selective about which one is included, thus the redundancy among the features in gBoost is minimal. By using elastic norm, RLMD retains several group effects (some discriminative features may be included and excluded simultaneously), and usually achieves better results. This result is consistent with observations reported in [21] for vector data.

564 5.2.3. Results on PTC Tasks

We also conducted extensive experiments on the PTC datasets. The accuracies and AUC values (*i.e.*, the area under ROC curves) are reported in Tables 3 and 4, where the results are obtained after RLMD converges, and K = 200 for all filter methods.

Table 3: Accuracies and standard errors on PTC graphs with Optimal K for 10-fold cross-validation.

ID	RLMD	TOP +SVM	IG +SVM	gHSIC +SVM	gSemi +SVM	gBoost
MR	0.655 ±0.013	0.606±0.024	0.607±0.025	0.596±0.019	0.601±0.020	0.608±0.027
MM	11	0.060 ± 0.025			0.603±0.019	0.622 ± 0.018
FR	0.704±0.018	0.607 ± 0.029	0.584±0.029	$0.635{\scriptstyle\pm0.026}$	0.619 ± 0.023	0.678 ± 0.008
FM	$0.615{\scriptstyle\pm0.018}$	$0.592{\scriptstyle\pm0.031}$	0.594 ± 0.026	0.581 ± 0.021	0.575 ± 0.018	0.603 ± 0.017

Table 4: AUC values and standard errors on PTC graphs with Optimal K for 10-fold cross-validation.

ID	RLMD	TOP +SVM	IG +SVM	gHSIC +SVM	gSemi +SVM	gBoost
MR	0.681±0.021	0.597 ± 0.025	0.596±0.025	0.560±0.021	0.580±0.021	0.649 ± 0.033
MM	0.680 ±0.020	0.593 ± 0.025	0.590 ± 0.025	0.583 ± 0.024	0.600 ± 0.022	0.600 ± 0.034
FR	0.673 ±0.022	0.600±0.029	0.575 ± 0.029	$0.618{\scriptstyle\pm0.021}$	0.623 ± 0.021	0.640 ± 0.026
FM	0.614 ±0.013	$0.585{\scriptstyle\pm0.032}$	0.587 ± 0.024	0.582 ± 0.021	0.580 ± 0.017	0.583±0.017

The results in Tables 3 and 4 show that RLMD achieves considerable performance gains over all filter methods (TFM and DFM) and gBoost algorithm for all PTC datasets.

570 Note that for PTC classifications, AUC values are more important because they are all

⁵⁷¹ imbalanced classification tasks.

572 5.2.4. Parameter Analysis

In this subsection, we study the impact of parameters γ_1 and γ_2 on algorithm per-

formance. Both γ_1 and γ_2 values are selected from {0, 0.01, 0.03, 0.05, 0.07, 0.09,

⁵⁷⁵ 0.11, 0.15, 0.2}, and the results under 10-fold cross-validation on NCI-1 and NCI-33

⁵⁷⁶ are shown in Fig. 5.

Impact of γ_1 **values:** The experimental results in Fig. 5 show that γ_1 plays a more

⁵⁷⁸ important role for the final classification model. With the increase of γ_1 from 0 to 0.2, the classification performance drops rapidly in terms of accuracy.

Table 5: Impact of different γ_1 values on NCI-1 dataset with $\gamma_2 = 0.03$, $S_{max} = 200$.

1	1 -		,			
γ_1	0	0.01	0.03	0.05	0.15	
#Selected Subgraphs	200	180	130	65	0	
Accuracy	0.775	0.759	0.711	0.679	0.5	
AUC	0.831	0.811	0.795	0.713	0.5	

579

To better understand the impact of γ_1 , we also summarize the number of subgraphs 580 selected with different γ_1 values in Table 5. The results show that increasing γ_1 values 581 will result in fewer subgraphs being selected for the final classifier model, because a 582 larger ℓ_1 norm regularizes more elements as 0. For $\gamma_1 = 0$, there is no sparse solu-583 tion. In other words, every subgraph should be used for graph classification. In this 584 case, RLMD will only terminate when all subgraphs are incorporated for learning the 585 model, or the maximum number of iterations S_{max} is reached. As the subgraph space 586 is exponentially large, it is impractical to use all subgraph features to learn the model. 587 The algorithm relies on S_{max} to terminate (200 is set in our experiment). The result 588 shows that $\gamma_1=0$ even achieves better classification result, which is attributed to the fact 589 that although ℓ_1 regularization introduces a sparse solution, it may be biased in some 590 applications [45], so the accuracy may drop. For other cases with γ_1 being considerably 591 large ($\gamma_1 = 0.5$), the regularization term dominates the objective function Eq.(4) with 592 no subgraph being used for classification, which results in poor classification accuracy. 593

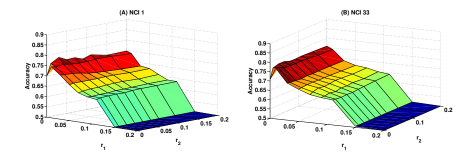


Figure 5: The accuracies with different γ_1 and γ_2 values.

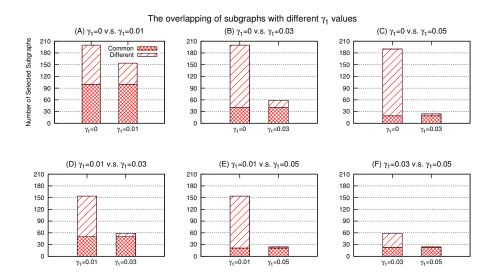


Figure 6: The overlapping of subgraphs (common subgraphs *vs.* different subgraphs) for different γ_1 values on NCI-1 dataset with a 70%-30% splitting on the NCI-1 dataset, *i.e.*, 70% graphs are randomly selected as training graphs, and 30% are used as test graphs. $\gamma_2 = 0.03$ and $S_{max} = 200$.

Table 6: Impact of different γ_2 values on NCI-1 dataset with $\gamma_1 = 0.01$.							
γ_2	0	0.03	0.07	0.15	0.2		
Accuracy	0.748	0.759	0.74	0.741	0.72		
AUC	0.810	0.811	0.806	0.811	0.78		

Table 7:	Impacts of	different ϵ	values	on NCI-1	dataset.

ε	0.01	0.005	0.001	0.0001	
Accuracy	0.748	0.759	0.761	0.760	
AUC	0.803	0.811	0.814	0.812	

Note that the convergence property of our algorithm is dependent on γ_1 . In our experiments, we notice that γ_1 is very easy to set (in a small range [0.01,0.03]) for obtaining satisfactory results. This is much easier than requiring users to specify the number of subgraph features *K* needed for each graph dataset, because users may not have any prior knowledge about the selection of *K* values for different datasets, and different *K* values often result in significant changes in the algorithm performance.

Interplay between γ_1 **and subgraph selection:** We further compare the common sub-600 graphs selected by different γ_1 values, and report the results in Fig. 6. The results show 601 that the subgraphs selected by using a smaller γ_1 values contain many subgraphs which 602 are selected by using a larger γ_1 value. This observation is further evident in Fig.6.(E) 603 and (F). The reason is that a smaller γ_1 value will result in more subgraph features to be 604 selected, which increases the possibility of covering a small subgraph set selected by 605 using a larger γ_1 value. In other words, a slightly smaller γ_1 value will result in more 606 subgraph feature candidates to be explored and be beneficial for the classification task. 607 **Impact of** γ_2 values: We also vary γ_2 from 0 to 0.2, and report the results in Table 608 6. The results show that a small regularization value $\gamma_2 = 0.03$ outperforms the case 609 of $\gamma_2 = 0$, where the ℓ_2 regularization effect disappears (only ℓ_1 is used). This result 610 is consistent with observations from a previous study [21]. This may be because ℓ_1 611 ignores the correlated subgraphs in a group of features. When γ_2 keeps increasing, 612 the classification performance drops because the larger ℓ_2 regularization dominates the 613 objective function and the loss minimization term has less effect. 614

Impact of ϵ values: We vary the ϵ values from 0.01 to 0.0001 to study the final classification performance of our algorithm, and report the final classification results in Table 7. The results show that as long as ϵ is subtle (from 0.001 to 0.0001), our classification can achieve similar classification results, *i.e.*, an ϵ -tolerance accuracy result to the

619 optimal solution.

620 6. Conclusion

In this paper, we proposed a regularized loss minimization-driven (RLMD) graph 621 classification method. We argued that existing filter-based subgraph selection methods 622 simply focus on finding most discriminative subgraph features, and suffer severe disad-623 vantages in determining the optimal number of subgraphs for graph classification and 624 separating feature selection from the model learning phase. As a result, they might be 625 able to find most discriminative subgraph features, but cannot form high accuracy clas-626 sifiers because they cannot determine how many discriminative features are needed to 627 train classifiers with the best performance gain. By integrating subgraph mining, dis-628 criminative subgraph selection, and model learning into one unified framework, RMLD 629 is able to automatically determine the optimal number of discriminative subgraphs for 630 best graph classification results. Our algorithm generalizes the state-of-the-art gBoost 631 algorithm in the sense that it can employ any differentiable loss function and achieve 632 better classification accuracy by using an elastic net regularization. Experimental re-633 sults on real-world graph datasets show a clear performance gain over existing two-step 634 filter methods (TFMs), direct filter methods (DFMs), and embedding methods. 635

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