A Unified Feature Selection Framework for Graph Embedding on High Dimensional Data

Marcus Chen, Ivor W. Tsang, Mingkui Tan, and Tat Jen Cham

Abstract—Although graph embedding has been a powerful tool for modeling data intrinsic structures, simply employing all features for data structure discovery may result in noise amplification. This is particularly severe for high dimensional data with small samples. To meet this challenge, this paper proposes a novel efficient framework to perform feature selection for graph embedding, in which a category of graph embedding methods is cast as a least squares regression problem. In this framework, a binary feature selector is introduced to naturally handle the feature cardinality in the least squares formulation. The resultant integral programming problem is then relaxed into a convex Quadratically Constrained Quadratic Program (QCQP) learning problem, which can be efficiently solved via a sequence of accelerated proximal gradient (APG) methods. Since each APG optimization is w.r.t. only a subset of features, the proposed method is fast and memory efficient. The proposed framework is applied to several graph embedding learning problems, including supervised, unsupervised, and semi-supervised graph embedding. Experimental results on several high dimensional data demonstrated that the proposed method outperformed the considered state-of-the-art methods.

Index Terms—Sparse graph embedding, sparse principal component analysis, efficient feature selection, high dimensional data

1 INTRODUCTION

HIGH dimensional data is ubiquitous in many real world applications, especially in this era of data [1], [2]. In microarray technology, a large sensor array can capture thousands of genes simultaneously. The latest camera phones such as the Nokia Lumia 1020 are able to capture photos of up to 41 megapixels. However, directly learning a classifier on high dimensional data may significantly degrade the performance of many applications, especially when data features are highly correlated and the sample size is relatively small. This is commonly referred to as the curse of dimensionality [3]. To alleviate this, one possible approach is to transform high dimensional data into a lower dimensional representation while preserving the intrinsic data structure. This is known as dimensionality reduction.

Graph embedding has been shown to be a powerful tool for dimensionality reduction [4], [5]. In particular, some popular dimensionality reduction methods such as Principal Component Analysis (PCA), Linear Discriminant Analysis (LDA) [6], Isomap [7], Locally Linear Embedding (LLE) [8], and Locality Preserving Projection (LPP) [4] can be formulated into graph embedding methods [5]. By employing full dimensional features for learning tasks, the graph embedding methods aim to learn a low dimensional projection, preserving some intrinsic data structures. Intrinsic data structures can have both local and global properties, depending on the applications. Local properties often refer to the local neighborhood relationship such as in LPP, while examples of global properties include class separation in LDA, the global variance in PCA, and the global shortest path between any pairs of data samples in the Isomap method. Graph embedding of high dimensional data suffers from two weaknesses. First, it is hard to interpret the resultant features when using all dimensions for embedding. Second, the original data inevitably contains noisy feature measurements. Simply incorporating these noisy features could make graph embedding unreliable and noisy [9], [10]. Therefore, it is important to select only the significant features for graph embedding.

Many feature selection methods have been proposed in different learning contexts [11], [12], [13], [14], [15], [16] with different feature importance measures. These methods can be categorized into two classes, namely, the supervised and unsupervised methods [17]. For the supervised methods, there are two main feature importance measures, distance based measures, and the correlation based measures. Specifically, the distance based measures define the important features as those that separate classes better and cluster the within-class samples, such as LDA-based feature selection methods [18]. In correlation based feature selection methods [19], the important features are those that correlate well with class labels and give better prediction results. In the unsupervised methods, due to the absence of class labels, several criteria have been proposed to evaluate the feature importance based on different learning contexts, such as information measure, variance measure, and locality measure, summarized as follows:
- Information measure: good features gain more information when included.
- Variance measure: good features capture more variance in the data.
- Locality measure: good features can preserve data locality better.

The existing feature selection methods are also often classified as filter, wrapper, and embedded approaches [20]. Filter methods aim to predefined some desired intrinsic data properties and rank feature subsets accordingly. These methods employ either forward selection or backward elimination search strategies, often resulting in local optimality. They do not consider the subsequent applications for feature evaluation, and are thus suboptimal in performance. He et al. [21] proposed to encode data locality via a Laplacian Score (LS) for feature ranking. Similarly, Zhao and Liu [22] proposed to evaluate features based on SPECTrum decomposition (SPEC) on the Laplacian matrix. These two methods employ feature level ranking, and do not consider the redundancy among features. To address this, Nie et al. [23] proposed a feature subset trace ratio method to rank feature subsets that best discriminate between-class samples and preserve within-class relationships.

In contrast to filter methods, wrapper methods are application dependent. The feature subset giving the best performance of a predetermined learning task will be chosen. The evaluation of each feature subset requires a complete run of the application, and is thus very computationally expensive. Generally, given a learning task, wrapper methods can perform better than filter methods [24]. Finally, embedded methods encapsulate the feature selection problem into their optimization objectives such as the solution sparsity [25], [26], [27], [28].

Among the sparsity induced methods, the sparse regression method, termed as LASSO [29], is popular in the literature due to its simplicity and efficiency in computation. Its formulation of the feature selection problem is as follows:

$$\min_w \|X^Tw - y\|_2^2 + \gamma\|w\|_1,$$

where $X \in \mathbb{R}^{d \times n}$ is the $d$-dimensional data matrix of $n$ samples, $y$ is the response variable vector, and $w$ is the weight vector. The regularization constant $\gamma$ controls the sparsity of solution; a larger $\gamma$ generally results in a sparser solution, and vice versa. However, LASSO can only choose at most $n$ variables even when $d \geq n$, and the solution is not well defined unless $\|w\|_1$ is bounded by a certain value [30]. The LASSO shrinkage may produce biased estimates for large coefficients of $w$ [31]. Zhou and Hastie [30] then proposed the Elastic net method, which adds an $L_2$ norm term, i.e., $\gamma\|w\|_1 + \beta\|w\|_2^2$. However, both LASSO and the Elastic net method do not guarantee the choice of the same set of features when regressing over different principal components ($y$). To address this issue, Cai et al. [32] proposed a simple heuristic method to rank the features based on the weight of different subspaces. This method is termed as MultiClusters Feature Selection (MCFS) with preserving data locality as its main learning objective.

Recently, the Convex Relaxations for Subset Selection method (CRSS) [33] was shown to achieve better results than the Least Angle Regression (LAR) method [34]. Bach et al. [33] proposes two search methods. One is a branch and bound search method and is combinatorial in computational complexity. The other one is based on the random generation of Gaussian vectors. Both LASSO and CRSS select features based on sequential iterations of principal components. In this manner, tuning parameters for different components is complicated [10]. Therefore, they do not optimize for the collective objectives. There has been little work of estimating multiple sparse components simultaneously. The most similar method by Vu et al. [10], Fantope Projection and Selection (FPS), optimizes the collective objectives for sparse PCA via a Fantope projection approach.

Both the graph embedding and feature selection methods define their own paradigms to preserve data intrinsic structures. In existing work in the literature, these two tasks have been done independently or mutually exclusively. This paper instead proposes a novel paradigm to unify these two schemes by performing feature selection and graph embedding simultaneously. As an application illustrated in Fig. 1b, the proposed paradigm is applied to the LPP graph embedding of some face images. The proposed method can automatically choose some important pixels that capture the variations in illumination conditions, poses, and facial expressions. The main contributions of this paper are summarized as follows.

- By exploiting the least squares formulation of graph embedding, we introduce a binary feature selector to directly constrain the desired number of features. We further reformulate the resultant problem as a convex semi-infinite programming problem (SIP). This novel feature selection scheme can be applied to unsupervised, supervised, and semi-supervised learning tasks in preserving the corresponding intrinsic data structures via low dimensional embeddings.
- By exploiting the observation that only a few constraints are active in the resultant SIP problem, we...
The proposed method addresses the learning in a low rank of two random matrices, $R^k$ and $R^k$. Moreover, the proposed method is guaranteed to converge globally.

- The proposed method addresses the learning in a holistic way, resulting in both generalized graph embedding and the desired cardinality of the features. A wide range of datasets have been tested in the experiments to verify the effectiveness of the proposed framework for unsupervised, supervised, and semi-supervised learning tasks.

The organization of the rest of the paper is as follows. In Section 2, we briefly review some graph embedding methods for dimensionality reduction and introduce the least squares formulation of graph embedding. Section 3 details the proposed approach. In Section 4, we conduct some experiments to compare our results with the current state-of-the-art algorithms. Section 5 concludes this paper.

## 2 Related Work and Preliminaries

In this section, we first briefly review the recent literature on graph embedding and feature selection. An overview of the generalized subspace problem is also provided since this is a foundation of our method.

### 2.1 Graph Embedding for Dimensionality Reduction

Coherent structures in high dimensional data, such as neighboring pixels in images, naturally induce a high correlation among dimensions. To alleviate the curse of dimensionality, scientists have proposed to transform data into a low dimensional manifold via graph embedding [5], [21]. The number of data samples, data dimensionality, and the number of classes are denoted by $n$, $d$, and $k$, respectively. $X \in \mathbb{R}^{d \times n}$ is a zero-mean data matrix, and $Y = \{y_1, y_2, \ldots, y_n\} \in \mathbb{R}^{k \times n}$ if available represents class label information. Furthermore, the symmetric positive semi-definite matrix, $S \in \mathbb{R}^{n \times n}$, encodes the desired data properties. Graph embedding for a class of dimensionality reduction methods aims to find the projection vector $\omega$ for the following generalization subspace problem [35]:

$$
XSX^T \omega = \lambda XX^T \omega. \tag{2}
$$

Many dimensionality reduction methods such as PCA, LDA, CCA, LPP, and Hypergraph Spectral Learning (HSL) can be formulated into the above graph embedding framework [5]. The definitions of $S$ for the above mentioned methods are tabulated in Table I. More details can be found in [35] and references therein.

Both PCA and LPP are unsupervised as they do not consider class labels. They are often used for general pre-processing, clustering, or visualization. For classification, supervised graph embedding, such as LDA, generally can achieve better performance since class label information is considered. The graph embedding can be readily extended to the semi-supervised setting, which utilizes large unlabeled datasets and small labeled datasets to model intrinsic data structures [36], [37], [38], [39]. To achieve this, a possible model is a weighted graph whose vertices are both labeled and unlabeled samples and edges reflect samples similarity. For example, the semi-supervised discriminative analysis (SDA) method [40] builds upon the LDA and LPP graph embeddings.

### 2.2 The Least Squares Formulation for Graph Embedding

Solving the generalized eigenvalue problem in (2) is very expensive for large-scale and high dimensional problems. To reduce the computational burden for large-scale problems, Sun et al. [35] formulates it into a least squares problem, as in (4). Specifically, since $S \in \mathbb{R}^{d \times n}$ is a symmetric positive semi-definite matrix, it can be decomposed as $S = HH^T$, where $H \in \mathbb{R}^{n \times r}$ and $r \leq n$ is the number of significant singular values of $S$. Furthermore, let $HP = QR$ be the QR-decomposition of $H$ with the permutation matrix $P$, where $Q \in \mathbb{R}^{n \times r}$ is a matrix with $r$ orthonormal columns. Moreover, let $R = U_2 \Sigma V_R^T$ be the compact singular value decomposition (SVD) of $R$, and the regression response $T \in \mathbb{R}^{n \times r}$ is computed as follows:

$$
T = QU_R. \tag{3}
$$

Then the generalized eigenvalue problem can be cast as the following least squares regression problem,

$$
\min_W \|X^T W - T\|^2_F, \tag{4}
$$

where $X \in \mathbb{R}^{d \times n}$ and $W \in \mathbb{R}^{d \times r}$ are data matrix and weight matrix, respectively. In practice, to improve the robustness to noise and avoid overfitting, a regularization term could be added as follows:

$$
\min_W \|X^T W - T\|^2_F + \gamma \|W\|^2_F. \tag{5}
$$

## 3 General Framework for Feature Selection

After transforming the generalized eigenvalue problem into a regression problem, the formulation (5) can benefit from many existing efficient least squares solvers. However, the regularizer $\gamma \|W\|^2_F$ may not produce sparse solutions. In other words, it cannot achieve the feature selection task. To
induce sparsity, we introduce a binary vector \( p \in \{0, 1\}^d \), whose entries are 1 for the selected features and 0 otherwise. To select \( m \) desired features, exactly \( m \) entries in \( p \) will be set to 1, where \( m < d \). Let \( \mathcal{P} = \{ p : p \in \{0, 1\}^d, p^\top 1 = m \} \) be the domain of \( p \), and 1 \( \in \mathbb{R}^d \) denotes a vector with all entries equal to 1. The proposed least squares formulation for graph embedding based feature selection is cast as the following optimization problem,

\[
\min_{p \in \mathcal{P}} \min_{W} \frac{1}{2} \left\| \Xi \right\|_F^2 + \frac{\gamma}{2} \left\| W \right\|_F^2, \quad \text{s.t.} \quad \Xi = X^\top \text{diag}(p)W - T,
\]

where \( T \in \mathbb{R}^{n \times r} \) is the response matrix, \( \text{diag}(p) \) is the matrix whose diagonal is the feature selector vector \( p \), and \( \Xi \in \mathbb{R}^{n \times r} \) denotes the residual matrix.

The proposed formulation has several advantages over the conventional approaches, which impose sparsity directly on \( W \) like the sparsity objective \( \sum_i \left\| W_i \right\|_1 \) in the sparse PCA (SPCA) method [41]. First, it selects features naturally with the desired cardinality. This is much more efficient than the sparsity induced methods, in which a regularizer constant controls cardinality. Second, the proposed model can be transformed to a convex programming problem [42], based on which an efficient solver can be developed. The similar schemes used in [42] and [18] are designed for the Fisher score method and classification model can be transformed to a convex programming problem even for small-sized \( p \) and \( m \).

### 3.1 Sparse Graph Embedding for Feature Selection

The optimization problem in (8) has a combinatorial number of constraints. However, only a few of them are active. Exploiting this observation, we adopt the cutting plane algorithm to solve the QCQP problem (8). The cutting plane algorithm iteratively finds the most active constraint and adds it to the active constraint set \( \mathcal{I} \), which is initialized to an empty set \( \emptyset \). The active constraint set \( \mathcal{I} \) is always a subset of \( \mathcal{P} \), i.e., \( \mathcal{I} \subseteq \mathcal{P} \). Given the updated set \( \mathcal{I} \), we solve the following subproblem,

\[
\max_{V \in \mathbb{R}^{d \times k}, p \in \mathbb{R}} \theta \quad \text{s.t.} \quad \theta \leq f(V, p), \quad \forall p' \in \mathcal{I}.
\]

We term our proposed procedure Sparse Graph Embedding (or SparGE for short), described in Algorithm 1.

**Algorithm 1. Sparse Graph Embedding for Feature Selection**

Input: data \( X \in \mathbb{R}^{d \times n} \), a positive semi-definite matrix \( S \), the desired feature cardinality \( m \).

1. Initialize \( \mathcal{I} = \emptyset \), and compute \( T \) according to (3). Assign \( t := 1 \).

2. Iterate the following two steps until convergence.
   a) Update \( V \) by solving the subproblem in (9).
   b) Find the most active constraint, which is indicated by \( p^* \), by solving \( p^* = \arg \max_p f(V, p) \), based on \( V \).

Output: \( \mathcal{I} = \{ p^1, p^2, \ldots, p^t \} \), with each \( p^j \) indexing the selected features.

Given \( V \), Step 2b) of Algorithm 1 requires us solving

\[
p^j = \arg \max_p f(V, p) = \arg \max_p \left\| \text{diag}(p) XV \right\|_F^2
\]

in order to find the most active constraint of problem (8). Let \( A = XV \in \mathbb{R}^{d \times r} \), and define \( s_i = \sum_{j=1}^d (A_{i,j})^2 \). The optimization problem becomes:

\[
\arg \max_p \left\| \text{diag}(p) XV \right\|_F^2 = \arg \max_p \sum_{i=1}^d s_i p_i.
\]

Apparently, problem (10) can be solved readily by sorting \( s \) and then setting its top \( m \) values in \( s \) to 1 and the rest to 0. In other words, the most active constraint can be identified by choosing the features with the \( m \) highest values in \( s \). The algorithm for the most active constraint analysis is summarized in Algorithm 2. The most active
constraint \( p' \) obtained is then added to the active constraint set \( \Pi := \Pi \cup \{ p' \} \).

**Algorithm 2. The Most Active Constraint Selection**

Input: data \( X \in \mathbb{R}^{d \times n} \), dual variable \( V \), the desired number of features \( m \), and the selection vector \( p \).

1. Set all the entries of \( p \) to 0.
2. Compute \( s_i = \sum_{k=1}^{d} (A_i)^2 \), \( \forall i = 1, \ldots, d \).
3. Sort \( s \) in descending order.
4. Set \( m \) entries of \( p \) w.r.t. the top \( m \) values of \( s \).

Output: \( p \) which defines the most active constraint.

### 3.2 The Subproblem Optimization

After updating the active constraint set \( \Pi \), we then solve the subproblem in (9) with reduced constraints as defined by \( \Pi \). Since the number of constraints in \( \Pi \) is no longer large, this problem is readily solved by a sub-gradient method, such as simpleMKL [18], [42]. However, solving this problem w.r.t. the dual variables \( V \) can be very expensive, in particular when \( n \) is very large.

Assume there are \( k \) active constraints in \( \Pi \), i.e., \( k = |\Pi| \). Even though there are a large number of features in \( X \), at most \( m k \) features are chosen by \( \Pi \subseteq \mathbb{P} \), where \( m k \ll d \). Based on this observation, the subproblem in (9) might be solved more efficiently w.r.t. the primal variables \( W \). To be more specific, following [43], we have the following proposition.

**Proposition 1.** The subproblem in (9) can be equivalently addressed in the following primal form:

\[
\min_{\Omega} \frac{\tau}{2} \left( \sum_{k=1}^{C} \| \Omega_k \|_F^2 \right)^2 + \frac{1}{2} \| \Xi \|_F^2, \tag{11}
\]

where \( \Xi = T - \sum_{k=1}^{C} X^T \text{diag}(p') \Omega_k^T \) denotes the regression residual matrix and \( \Omega_k \) denotes the weight matrix defined on the features indicated by \( p' \). Moreover, the dual variable \( V \) in (9) can be recovered by \( V = \Xi \), which is required for the most active constraint selection.

The proof of this proposition is included in Appendix A. Problem (11) is  a non-smooth problem due to the regularization term \( \frac{\tau}{2} \left( \sum_{k=1}^{C} \| \Omega_k \|_F^2 \right)^2 \). However, there are at most \( m k \) (where \( m k \ll d \)) features involved in this problem, making it easier to be solved. For convenience, we define \( \Omega = [\Omega_1, \Omega^2, \ldots, \Omega^C] \in \mathbb{R}^{d \times \ell} \) by stacking \( \Omega_k \in \mathbb{R}^{d \times \ell} \). Let \( P(\Omega) = \frac{\tau}{2} \left( \sum_{k=1}^{C} \| \Omega_k \|_F^2 \right)^2 \) and \( f(\Omega) = \frac{1}{2} \| \Xi \|_F^2 \). Following [44], we propose to solve the primal problem using the accelerated proximal gradient method (APG), which iteratively minimizes the following quadratic approximation of (11):

\[
Q(\Omega, \Omega_\ell) = f(\Omega_\ell) + \langle \nabla f, \Omega - \Omega_\ell \rangle + \frac{\tau}{2} \| \Omega - \Omega_\ell \|_F^2 + P(\Omega) \]

\[
= \frac{\tau}{2} \| \Omega - \Omega_\ell \|_F^2 + P(\Omega) + f(\Omega_\ell) - \frac{1}{2\tau} \| \nabla f \|_F^2, \tag{12}
\]

where \( \nabla f \) denotes the gradient of \( f \) at point \( \Omega_\ell \), \( \tau > 0 \) denotes the Lipschitz constant of \( f(\Omega) \), and \( G = \Omega - \frac{1}{\tau} \nabla f \).

1. In practice, the optimization is conducted on those selected features only.

### 3.3 Handling High Dimensional Sparse Problems

Given an ultrahigh dimensional sparse data matrix, removing the data mean (zero-centering) could make the matrix very dense. The data matrix \( (X_{1:n}) \) can be used instead for regression to remove the data offset. As for the proposed framework, zero-centering can be performed in each subproblem. Zero-centering could also affect the computation of some regression responses \( T \), such as PCA as in Table I, which assumes zero mean. In this case, \( X X^T \) can be first computed and centering can then be applied to both rows and columns as follows:

\[
X \left( I - \frac{1}{n} 1_n 1_n^T \right), \tag{16}
\]

\[
S = \left( I - \frac{1}{n} 1_n 1_n^T \right) X X^T \left( I - \frac{1}{n} 1_n 1_n^T \right). \tag{17}
\]

The overall APG algorithm for solving problem (11) is summarized in Algorithm 4, where \( F(\Omega) = \frac{\tau}{2} \left( \sum_{k=1}^{C} \| \Omega_k \|_F^2 \right)^2 + \frac{1}{2} \| \Xi \|_F^2 \). Interested readers can find more details and the convergence derivation of Algorithm 1 and Algorithm 4 in [44].
4.1 Experiments on Unsupervised Sparse Embedding

The proposed SparGE-PCA method chooses feature subsets that maximize the variance in the data in an unsupervised manner. It optimizes over all principal components simultaneously. On the other hand, most of the sparse PCA methods [41], [47], [48], [49] select features sequentially over the principal components (PC). It is assumed that the feature subset derived from the first PC should be more important, but this may not be true. A simple counterexample is to find one feature explaining the most variance of the covariance matrix $\{3.2, 0.0; 0.3, 3; 0.3, 3\}$. In this case, SPCA [41] will select either the second or the third feature based on the first PC, but the correct selection should actually be the first feature. Only the recent concurrent work on Fantope Projection and Selection (FPS) by Vu et al. [10] shares a similar optimization objective as our proposed method.

To compare the proposed SparGE-PCA method with FPS and SPCA, we conducted experiments on both simulated data and a real gene data dataset shown in Table 3. The $\%$ of explained variance $r_\Sigma$ was used to measure the quality of the selected feature set as follows:

$$r_\Sigma = \frac{\text{trace}(\Sigma_f)}{\text{trace}(\Sigma)} \times 100\%,$$

where $\Sigma_f$ and $\Sigma$ are the covariances of the selected features and of all features, respectively. A larger $r_\Sigma$ indicates a better feature subset. The results of 50 independent runs are reported. For SPCA, we chose the features with the highest absolute magnitudes of the weight matrix, similar to [32].

4.1.1 The Results on Simulated Data

Optimizing for global variance, a good sparse embedding method should be able to identify a feature subset explaining the most variance and also removing noisy features. To test the optimality of the selected features of the proposed SparGE-PCA method, a simulation experiment is carried out. In this experiment, a toy data is generated with 50 significant features and 500 noisy features. Its covariance is shown in Fig. 2a. The explained variance should converge at 50 features. Non-zero values in the off-diagonal entries indicate a correlation among the first 50 significant features.

As shown in Fig. 2c, FPS, SPCA, and the proposed SparGE-PCA method all converged to the optimal variance at 50 features in accordance to the groundtruth. Computationally, SPCA was an order of magnitude slower than the proposed method, as shown in Fig. 2d. FPS shared a similar computational time with SPCA.

4.3 Computational Complexity

There are a few components in the proposed framework. Given $S \in \mathbb{R}^{n \times r}$ with $r$ significant singular values, the proposed framework requires a Cholesky decomposition $S = HH^T$ and a QR-decomposition of $HR$, SVD of $R$, finding $p^t$, and solving the subproblem. The decomposed matrices are compact, i.e., $Q \in \mathbb{R}^{n \times r}$, $R \in \mathbb{R}^{r \times r}$. Table 2 shows the computational complexity of different components. For $r \leq n$ and the desired feature cardinality $m_k \ll d$, the overall computational complexity is $O(ndr)$. This is much more efficient than FPS [10], whose complexity is $O(d^3 + nd^2)$ especially for high dimensional data applications where $d \gg m_k$, $d \gg n$, and $d \gg r$.

<table>
<thead>
<tr>
<th>Modules</th>
<th>Cholesky &amp; QR decompositions</th>
<th>SVD</th>
<th>Finding $p^t$</th>
<th>Subproblem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Details</td>
<td>$S = HH^T$, $H = QR$</td>
<td>$U_R \Sigma_R V_R^T$</td>
<td>Compute regression $XY$, $s$ in (10), and sort $s$</td>
<td>$O(mnr)$</td>
</tr>
</tbody>
</table>

**Algorithm 4. Accelerated Proximal Gradient for Solving Problem (11)**

- **Initialization:** Initialize the Lipschitz constant $L_t = L_{t-1}$ and set $\Omega^{-1} = \Omega^0$ by warm start, $\tau_0 = L_t$, $\eta \in (0, 1)$, parameter $\varrho^{-1} = \varrho^0 = 1$, and $k = 0$.
- **Set** $V^0 = \Omega^k$ + $\varrho^{-1}$ $(\Omega^k - \Omega^{-1})$.
- **Set** $\tau = \eta \tau_k$.

- **Repeat**
  - **Set** $G = V^k + \frac{1}{2} \nabla f(V^k)$, compute $S_t(G)$.
  - **If** $F(S_t(G)) \leq Q(S_t(G), V^k)$
  - **Set** $t_{k+1} = \tau$, stop, break;
  - **Else**
  - **Set** $\tau = \min(\eta^{-1} \tau, L_t)$.

- **Until** convergence $F(S_t(G)) \leq Q(S_t(G), V^k)$
- **Set** $\Omega^{k+1} = S_t(G)$.
- **Let** $\varrho^{k+1} = \frac{1 - \sqrt{1 + (4 \varrho^k)^2}}{2}$.
- **Let** $k = k + 1$.
- **Quit** if the stopping condition is achieved. Otherwise, go to step 1.
- **Let** $L_t = \eta^2 \tau_k$ and return.

**TABLE 2**

<table>
<thead>
<tr>
<th>Approximation</th>
<th>Proposed Framework</th>
<th>SVD</th>
<th>Finding $p^t$</th>
<th>Subproblem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complexity</td>
<td>$O(n^2)$</td>
<td>$O(r^3)$</td>
<td>$O(ndr + d^2) + m \log d$</td>
<td>$O(mnr)$</td>
</tr>
</tbody>
</table>

**TABLE 3**

<table>
<thead>
<tr>
<th>S/N</th>
<th>Data</th>
<th># dimensions</th>
<th># instances</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Toy data</td>
<td>550</td>
<td>5,000</td>
</tr>
<tr>
<td>2</td>
<td>Ramaswamy data</td>
<td>16,063</td>
<td>144</td>
</tr>
</tbody>
</table>
4.1.2 The Results on Real Data Sets

PCA is often used for analyzing high dimensional data with small samples, especially biological data. In this experiment, the microarray data, Ramaswamy data [41], is used to find the meaningful genes from very high dimensional data. The data has a very high dimensionality of 16,063 (genes) and 144 samples only. Only the first 25 principal components were used to select features.

In this experiment, the FPS method was unable to handle such a high dimensionality since its computational cost is too high. The proposed SparGE-PCA method performed similarly to SPCA and FPS in the explained variance on a toy dataset with the first 80 important features, but it was at least an order of magnitude faster than SPCA and FPS. PCA with 25 subspaces using all of the dimensions explains about 99.8 percent of the total variance.

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complexity is $O(d^3 + nd^2)$ when performing a Fantope Projection. Therefore, only the SparGE-PCA and SPCA methods were chosen for comparison.

The proposed SparGE-PCA method outperformed SPCA significantly in the explained variance, by about 10 percent between 200 and 350 features as shown in Fig. 3a. Both SPCA and the proposed SparGE-PCA method converged to 70 percent in the explained variance with 500 features (only about 3.1 percent of the total features). Computationally, SPCA was at least two orders of magnitude slower than the proposed method as shown in Fig. 3b.

4.2 Experiments on Feature Selection for Clustering
Besides sparse graph embedding for PCA, our proposed framework can be used to identify important features for clustering tasks. As discussed in Section 1, unsupervised LPP can model the local data structure well, and thus its embedding could improve clustering performance. Since image datasets, such as digits and faces images, usually lie on a low dimensional manifold, four popular image datasets, namely MNIST, COIL20, ORL, and USPS as shown in Table 4 were chosen.

To evaluate the quality of selected features, we apply k-means clustering on the data with the chosen features, where $k$ is set to the number of classes. The normalized mutual information defined in [21] is used as the performance measure. The clustering baseline employed all of the features. Besides the baseline, the feature ranking methods such as Laplacian Scores (LS) and MultiClusters Feature Selection (MCFS) were also chosen for a comparative evaluation.

The result was shown in Fig. 4 with up to 250 features only. Interestingly, the baseline results could be achieved with as few as 50 to 70 features. Both MCFS and the proposed SparGE-LPP method performed much better than the LS feature selection method. The proposed SparGE-LPP method also outperformed MCFS on MNIST, COIL20, and USPS. Note that MCFS employed a simple ranking method to choose features from a sequential learning framework. Theoretically, it is unclear whether this approach could converge to optimality. Furthermore,
it does not learn the collective weights based on the common feature subsets. In contrast, the proposed framework could achieve both weight learning and feature subset selection simultaneously.

4.3 Experiments on Supervised Feature Selection

In this section, we compare the proposed SparGE-LDA method with some chosen feature selection methods on classification tasks. Six datasets as shown in Table 5 are used for comparison. The first three datasets ranging from text, images, and microarray data, have medium dimension and small sample sizes, and are collected from the Arizona State University (ASU) feature selection repository [50].

<table>
<thead>
<tr>
<th>S/N</th>
<th>Data</th>
<th># dimensions</th>
<th># instances</th>
<th># classes</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PCMAC</td>
<td>3,289</td>
<td>1,943</td>
<td>2</td>
<td>Text</td>
</tr>
<tr>
<td>2</td>
<td>ORL10P</td>
<td>10,304</td>
<td>100</td>
<td>10</td>
<td>Face image</td>
</tr>
<tr>
<td>3</td>
<td>GLI-85</td>
<td>22,283</td>
<td>85</td>
<td>2</td>
<td>Microarray data</td>
</tr>
<tr>
<td>4</td>
<td>Real-Sim</td>
<td>20,958</td>
<td>72,309</td>
<td>2</td>
<td>Text</td>
</tr>
<tr>
<td>5</td>
<td>RCV1</td>
<td>47,236</td>
<td>training:20,242, testing: 67,739</td>
<td>2 Text</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>News20</td>
<td>62,060</td>
<td>training:15,935, testing: 3,993</td>
<td>20 Text</td>
<td></td>
</tr>
</tbody>
</table>

Unless indicated otherwise, datasets were split for 10-fold cross validation.

Fig. 5. Comparison of different feature selection methods on classification tasks. Classification results of PCA as pre-processing is also included as a benchmark for the first three datasets.
other three higher dimensional and large-scale text datasets are from the LIBSVM [51] repository.

Several popular feature selection methods, such as Relief-F [52], max-dependency Max-Relevance and Min-Redundancy (MRMR) [53], Fisher score, and SVM Recursive Feature Elimination (SVM-RFE) are chosen for comparison. Beside these, we also compared the most recent method by Nie et al. [23], the Trace Ratio Criterion (TRC) method. TRC was shown to outperform many methods in the literature. PCA is also included as a benchmark for classification using SVM. However, PCA cannot handle datasets such as Real-Sim, RCV1, and News20, which have both ultrahigh dimensions and large data sample sizes. Therefore, no comparison with PCA for these three datasets is included. For RCV1 and News20 datasets, the accuracies on the test sets are reported. For the rest, the mean accuracies of 10-fold cross-validation are reported.

The results are shown in Fig. 5. We do not report the results of Relief-F, TRC, and MRMR on the three high dimensional and large-scale datasets because these methods are computationally inefficient on these datasets. Generally, both TRC and the SparGE-LDA method can handle global feature subset directly, they outperformed the feature-level selection processes such as MRMR, Relief-F, and Fisher score methods. Compared with TRC, our SparGE-LDA outperformed for the ORL10P and GLI-85 significantly, was marginally better than TRC for the PCMAC dataset. In the TRC method, the optimization enforces only one feature in each column of the weight matrix; it is thus more constrained and may thus have a poorer performance.

On the other hand, the SparGE-LDA method outperformed SVM-RFE for the ORL10P, PCMAC, Real-Sim, RCV1, and News20 datasets. On the first three small datasets, SVM-RFE performed well too as it directly optimized for the classification methods. However, SVM-RFE is a greedy method, and its performance dropped significantly in the higher dimensional and large-scale data shown in Fig. 5d, 5e, and 5f. Compared to SVM-RFE, SparGE-LDA achieved 20 percent better in performance on the News20 dataset, 5 percent on the Real-Sim dataset, and 10 percent on the RCV1 dataset. On the other hand, the Fisher score method, the counterpart of SparGE-LDA, significantly underperformed on the last three high dimensional datasets.

Computationally, the Fisher score method was the fastest, but SparGE-LDA was also efficient and completed the tasks within minutes as shown in Table VI. SVM-RFE was generally fast, but it was very slow on the News20 dataset, which had the highest dimension and number of classes among the three ultrahigh dimensional datasets.

### 4.4 Experiments on Semi-Supervised Feature Selection

So far, the experiments demonstrated the effectiveness of our proposed framework for both unsupervised and supervised learning settings. By incorporating small labeled samples, the semi-supervised discriminant analysis based on the proposed framework (SparGE-SDA) can achieve a good classification rate.

Two datasets (COIL and ORL) from Table 4 are used for evaluation. The proposed SparGE-SDA method was compared with the SVM-RFE and Fisher score feature selection methods. The classification accuracies are tabulated in Table 7. The proposed SparGE-SDA method was much better than the Fisher score method and consistently better than SVM-RFE.

### 4.5 Experiments on Data Visualization

In this section, we intend to identify the important face features that can preserve the data locality, i.e., the neighborhood relationships of these image samples. The data locality is visualized using two dimensional linear embedding of face images by LPP. From the CMU-PIE database [54], 170 face images of one person shown in Fig. 1a were chosen for this experiment. The linear embedding of these images is shown in Fig. 6. In Fig. 6a, it can be observed that the illumination increases from left to right, and her face turns from left to right. The faces with expressions are far apart from the rest showing on the top, indicating a large difference. A similar observation can be drawn using the proposed SparGE-LPP with only 20 features. Variations in illumination, poses, and expressions are much more gradual on the

### Table 7

<table>
<thead>
<tr>
<th>Datasets</th>
<th># feats</th>
<th>1 labeled samples</th>
<th>2 labeled samples</th>
<th>3 labeled samples</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SparGE-SDA</td>
<td>SVM-RFE</td>
<td>Fisher</td>
<td>SparGE-SDA</td>
</tr>
<tr>
<td>COIL</td>
<td>100</td>
<td>59.7</td>
<td>56.8</td>
<td>31.9</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>61.2</td>
<td>57.6</td>
<td>37.0</td>
</tr>
<tr>
<td></td>
<td>300</td>
<td>64.2</td>
<td>57.9</td>
<td>50.0</td>
</tr>
<tr>
<td>ORL</td>
<td>100</td>
<td>59.0</td>
<td>48.5</td>
<td>25.3</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>64.7</td>
<td>52.2</td>
<td>40.0</td>
</tr>
<tr>
<td></td>
<td>300</td>
<td>65.0</td>
<td>53.9</td>
<td>48.5</td>
</tr>
</tbody>
</table>
embedding shown in Fig. 6b, indicating a better fit of the underlying manifold.

5 Conclusion

This paper proposes a novel unified framework to select features for generalized graph embedding. It utilizes a feature selector to directly optimize feature subsets for graph embedding in modeling the intrinsic data structures, enabling a more robust embedding, especially for high dimensional data with a small sample size. Its efficiency and effectiveness have been demonstrated with a series of experiments for clustering, classification, and visualization. In the experiments, the proposed methods outperformed the current state-of-the-art algorithms for unsupervised, supervised, and semi-supervised learning tasks. The proposed framework demonstrated its computational and memory efficiency in handling ultrahigh dimensional data for classification.

Appendix A: Proof of Proposition 1

In Proposition 1, the dual form of problem (11) has the same form as in problem (9).

Let $\Omega = [\Omega^1, \Omega^2, \ldots, \Omega^K]$ be the stack of $\Omega^i$. Define the cone $\mathcal{Q} = \{(\Omega, v) \in (\mathbb{R}^{d \times r}, \mathbb{R})|\|\Omega\|_F \leq v\}$. Let $z_i = \|\Omega^i\|_F, z = \sum_{i=1}^K z_i$. The optimization problem in (11) is equivalent to the following problem:

$$
\min_{z, \Sigma} \frac{\lambda}{2} z^2 + \frac{1}{2} \|\Sigma\|^2_F,
$$

s.t. $\Sigma = T - \sum_{i=1}^K X^T \text{diag}(p^i) \Omega^i$,

$$
\sum_{i=1}^K z_i \leq z, \quad (\Omega^i, z_i) \in \mathcal{Q},
$$

The Lagrangian function of (19) can be written as:

$$
\mathcal{L} = \frac{\nu}{2} z^2 + \frac{1}{2} \|\Sigma\|^2_F - tr \left( V^T \left( \sum_{i=1}^K X^T \text{diag}(p^i) \Omega^i - T + \Theta \right) \right) + \eta \left( \sum_{i=1}^K z_i - z \right) - \sum_{i=1}^K \left( tr(\xi^i) + \mu_i z_i \right),
$$

where $V \in \mathbb{R}^{n \times r}$, $\eta \in \mathbb{R}$, $\xi^i \in \mathbb{R}^{n \times r}$, and $\mu_i \in \mathbb{R}$ are the Lagrangian dual variables for the corresponding constraints. By setting the derivatives of $\mathcal{L}$ w.r.t. $z, z_i, \Omega^i$, and $\Sigma$ to zero, we obtain the KKT conditions as follows:

$$
g z = \eta, \xi^i = -X \text{diag}(p^i) V, \Sigma = -V \|\xi^i\|_F \leq \eta.
$$

Substitute these results into the Lagrangian function, and we obtain the dual problem as follows:

$$
\max_{V, p^i} tr(V^T T) - \frac{1}{2} tr(V^T V) - \frac{1}{2} \eta^2
$$

s.t. $\|X \text{diag}(p^i) V\|_F \leq \eta, t = 1, \ldots, k.$

(20)

Setting $\theta = tr(V^T T) - \frac{1}{2} tr(V^T V) - \frac{1}{2} \eta^2$ and $f(V, p^i) = tr(V^T T) - \frac{1}{2} \|X \text{diag}(p^i) V\|^2_F - \frac{1}{2} tr(V^T V)$, then the problem becomes as follows:

$$
\max_{V \in \mathbb{R}^{n \times r}, p^i \in \mathbb{R}} \theta
$$

s.t. $\theta \leq f(V, p^i), \quad \forall p^i \in \Pi.$

(22)

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