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Sparsity preserving discriminant analysis for single training image face recognition

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ABSTRACT

Single training image face recognition is one of the main challenges to appearance-based pattern recognition techniques. Many classical dimensionality reduction methods such as LDA have achieved success in face recognition field, but cannot be directly used to the single training image scenario. Recent graphbased semi-supervised dimensionality reduction (SSDR) provides a feasible strategy to deal with such problem. However, most of the existing SSDR algorithms such as semi-supervised discriminant analysis (SDA) are locality-oriented and generally suffer from the following issues: (1) they need a large number of unlabeled training samples to estimate the manifold structure in data, but such extra samples may not be easily obtained in a given face recognition task; (2) they model the local geometry of data by the nearest neighbor criterion which generally fails to obtain sufficient discriminative information due to the highdimensionality of face image space; (3) they construct the underlying adjacency graph (or data-dependent regularizer) using a fixed neighborhood size for all the sample points without considering the actual data distribution. In this paper, we develop a new graph-based SSDR algorithm called sparsity preserving discriminant analysis (SPDA) to address these problems. More specifically, (1) the graph in SPDA is constructed by sparse representation, and thus the local structure in data is automatically modeled instead of being manually predefined. (2) With the natural discriminative power of sparse representation, SPDA can remarkably improve recognition performance only resorting to very few extra unlabeled samples. (3) A simple ensemble strategy is developed to accelerate graph construction, which results in an efficient ensemble SPDA algorithm. Extensive experiments on both toy and real face data sets are provided to validate the feasibility and effectiveness of the proposed algorithm.

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1. Introduction

One of the major challenges to appearance-based face recognition is the small sample size (SSS) problem (Duda et al., 2001). In particular, in many practical applications such as law enforcement, driver license or passport card identification, usually only one labeled sample per person is available. Under such scenario, most of the traditional methods including Eigenface (Turk and Pentland, 1991) and Fisherface (Belhumeur et al., 1997) will suffer serious performance drop or even fail to work. Therefore, special tricks, such as synthesizing virtual sample (Beymer and Poggio, 1995) and localizing the training image (Chen et al., 2004), are generally required to deal with the single training sample problem. One can refer to a recent survey (Tan et al., 2006) for more details on this topic.

Although much success has been achieved by synthetic sample techniques, such artificial process has trouble in capturing the real face data distribution due to the variations of pose, illumination and facial expression (Tan et al., 2006). An alternative and more natural way to deal with such problem is semi-supervised dimensionality reduction (SSDR) if considerable unlabeled samples are available. For example, the recent semi-supervised discriminant analysis (SDA) (Cai et al., 2007), a semi-supervised extension of typical linear discriminant analysis (LDA), has been successfully applied to the single training image face recognition problem.¹ Besides SDA, researchers have developed some special SSDR algorithms, such as SSLDA (Song et al., 2008), SSMMC (Song et al., 2008), and lapLDA (Chen et al., 2007) and reported that the semisupervised extensions can generally improve the performance over their supervised counterparts like LDA and MMC (Li et al., 2006, 2007). Despite being independently proposed, these SSDR algorithms share similar starting point and can be unified under a graph-based dimensionality reduction framework (Yan et al., 2007; Song et al., 2008). We will give a brief review on these methods in the next section.





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¹ Strictly speaking, it should be called "single *labeled* image face recognition problem". We abuse the terminology, i.e., single *training* image face recognition problem, just for keeping consistent with the expression in (Cai et al., 2007).

Despite the success of many graph-based SSDR algorithms in dealing with partially labeled face recognition problem (Cai et al., 2007; Song et al., 2008), there are still some problems that are not properly addressed, especially under the single labeled training image scenario. In particular,

- (1) Many existing graph-based SSDR algorithms are based on manifold assumption, implying that sufficiently many samples are required to characterize the data distribution (Belkin et al., 2006). For example, with a large number of auxiliary unlabeled training samples, SDA can remarkably improve the performance of LDA. However, it is generally uneasy to obtain a sufficient sampling for intrinsic highdimensional data such as face images.² Therefore, a natural question is: can we improve the performance of LDA just with very few extra unlabeled samples?
- (2) As pointed out in (Zhu, 2008), although graph is at the heart of the graph-based semi-supervised methods, its construction has not been studied extensively. Most of the current algorithms such as SDA and lapLDA construct their adjacency graphs by the nearest neighbor criterion on raw data set. However, the nearest neighbor criterion generally fails to obtain sufficiently discriminative information due to its poor performance in the original high-dimensional face space.
- (3) The underlying adjacency graphs (or data-dependent regularizers) involved in many SSDR algorithms are artificially defined beforehand and use a fixed neighborhood size for all the sample points. Not only does this ignore the actual data distribution, but also bring the difficulty of parameter selection, especially when only few labeled samples are available as in single training image face recognition.

To address the above issues, in this paper, we present a new graph-based SSDR algorithm called *sparsity preserving discriminant analysis* (SPDA) which is motivated by the recent progress in sparse representation (Qiao et al., 2010; Wright et al., 2009). Concretely, we highlight the favorable properties of SPDA and main contributions of this paper:

(1) SPDA can remarkably improve the performance of typical LDA only resorting to very few extra unlabeled samples, because it does not based on manifold assumption, but mainly focuses on the discriminative power which can be naturally achieved by minimizing a ℓ_1 -regularization objective function. We will give a detailed discussion on this point in Section 3.

- (2) Graph construction involved in SPDA relies on sparse representation classification criterion (Wright et al., 2009) which is generally superior to the nearest neighbor criterion, especially for high-dimensional data.
- (3) The "neighborhood" size and edge weight for each sample are automatically obtained in one single step by a ℓ_1 optimization problem. As a result, different sample will get different neighborhood sizes, which is more adaptive to complex data distribution.
- (4) Alternatively, we develop a simple ensemble SPDA algorithm to reduce the computational complexity involved in obtaining sparse representation for graph construction when a large number of unlabeled samples are provided. Also, as a byproduct, we formulate the kernelized version of SPDA.

(5) The idea behind SPDA is quite general, and can potentially be extended to other graph-based semi-supervised learning algorithms by integrating with different discriminant criteria or loss functions.

The rest of the paper is organized as follows. Section 2 briefly reviews several existing graph-based SSDR algorithms. In Section 3, we develop a new data-dependent regularizer and SPDA algorithm. In Section 4, we extend SPDA to kernel and ensemble versions. Section 5 shows the experimental results, followed by the conclusion and future work in Section 6.

2. Brief review of semi-supervised dimensionality reduction (SSDR)

Firstly, we want to make clear that why we employ SSDR instead of other Semi-supervised learning (SSL) algorithms for the single training image face recognition problem. Indeed, various SSL algorithms have been developed in the past few years. One can refer to (Zhu, 2008) for a detailed literature survey. However, as pointed out in (Cai et al., 2007), many of the existing SSL algorithm can only work on *transductive* setting, which requires both the training and test set are available during the learning process. Therefore, they are not always suitable for face recognition applications where the test set is generally not available during the training phrase. In contrast, SSDR first learns a subspace from the available training set (containing labeled and unlabeled samples), and then the forthcoming test sample is projected onto the subspace for further decision.

2.1. Semi-supervised discriminant analysis (SDA) (Cai et al., 2007)

SDA extends LDA to incorporate the manifold structure illustrated by both labeled and unlabeled data. Therefore, SDA aims to best preserve the discriminative information as well as the geometric structure in data. Given a set of data points $X = [x_1, x_2, ..., x_n]$ including both labeled and unlabeled samples, the SDA objective function is defined as follows:

$$\max_{w} \frac{w^{T} S_{b} w}{w^{T} S_{t} w + \lambda_{1} w^{T} w + \lambda_{2} J_{MR}(w)},$$
(1)

where S_b and S_t are respectively the inter-class and total scatter matrix calculated using the labeled training samples. $w^T w$ is the Tikhonov regularizer, and $J_{MR}(w)$ is a data-dependent manifold regularizer (Belkin et al., 2006). λ_1 and λ_2 are two parameters, controlling the balance among the three terms in denominator. Obviously, if $\lambda_1 = \lambda_2 = 0$, SDA becomes the standard LDA; if $\lambda_1 \neq 0$, $\lambda_2 = 0$, it becomes the regularized discriminant analysis (RDA) (Hastie, 2009).

The data-dependent regularizer in SDA plays a role in preserving the manifold structure in data. It is constructed using both labeled and unlabeled training samples as follows:

$$J_{\rm MR}(w) = w^T X L X^T w = \sum_{ij} (w^T x_i - w^T x_j)^2 p_{ij},$$
(2)

where *L* is the graph Laplacian (Belkin and Niyogi, 2003), p_{ij} is the edge weight between data point x_i and x_j . In particular,

$$p_{ij} = \begin{cases} \exp\{\|x_i - x_j\|/2\sigma^2\}, & x_i \in N_k(x_j) \lor x_j \in N_k(x_i), \\ 0, & \text{otherwise.} \end{cases}$$
(3)

Since SDA shares the similar objective function to LDA, one can solve SDA by the following generalized eigenvalue problem:

$$S_b w = \eta (S_t + \lambda_1 I + \lambda_2 X L X^T) w.$$
⁽⁴⁾

 $^{^{2}}$ A recent research (Meytlis and Sirovich, 2007) has shown that the face space is estimated to have at least 100 dimensions.

Table 1 Several special SSDR algorithms proposed recently.

Algorithm	Discriminant criterion Fisher	MMC ^a	Regularization term Tiknonov	Manifold
SDA (Cai et al., 2007) LapLDA (Chen et al., 2007) SSLDA (Song et al., 2008) SSMMC (Song et al., 2008)	$\sqrt[]{}$	\checkmark	\checkmark \checkmark	\checkmark \checkmark \checkmark

^a Maximum margin criterion.

2.2. Other SSDR algorithms

Although, in the recent years, many graph-based SSDR algorithms have been proposed independently, most of them share the same idea: the labeled sample points are used to maximize the discriminative power, while the unlabeled sample points are used to best preserve the geometric structure in data. As a result, they are similar to each other with different choices of discriminant criterion and regularization term. Table 1 gives several popular examples of those methods.

Since the discriminant criteria (e.g., Fisher criterion and MMC) are usually off-the-shelf, the data-dependent regularizer naturally plays an important role in the graph-based SSDR algorithms. Also, we notice that the data-dependent regularizer is generally determined by a graph constructed based on both labeled and unlabeled samples. For example, in SDA, the manifold regularizer roots in the above mentioned *k*-neighborhood graph (3). Therefore, in the next section, we will start to introduce our SPDA algorithm from constructing a novel graph.

3. Sparsity preserving discriminant analysis

3.1. Graph construction based on sparse representation

3.1.1. Motivation from sparsity

We first give the reasons why the sparse representation is suitable to graph construction.

- (1) Sparsity plays an important role in typical k-neighborhood graph. On one hand, sparsity implicitly characterizes the locality of data distribution; on the other hand, it can effectively save computational cost and storage space. However, for the typical k-neighborhood graph constructed by Eq. (3), its sparsity depends on artificially fixed neighborhood size. It seems to be unreasonable that all data points share an identical k, which may not characterize the manifold structure well, especially in undersampling case. This motivates us to consider whether we can automatically learn the sparsity from the data instead of artificial predefinition.
- (2) The sparsest representation is naturally discriminative. Since our ultimate goal is classification, we expect that the graph can contain as much discriminative information as possible. That is, two data points are linked by an edge if they are likely from the same class. For the typical kneighborhood graph, this desirability depends heavily on how well the nearest neighbor criterion works in original space (Chen et al., 2005). Unfortunately, the nearest neighbor criterion does not generally achieve good performance for raw high-dimensional data, e.g., face images (Meytlis and Sirovich, 2007). In contrast, the recent researches (Wright et al., 2009) showed that sparse representation has natural discriminative power and can work well under high-dimensional scenario. Moreover, the discriminative power is closely related to the class numbers rather than

the sample numbers (Wright et al., 2009). As a result, we might construct a graph which contains considerable discriminative information without requiring abundant unlabeled samples.

3.1.2. The objective function for graph construction

Instead of considering *k*-neighborhood and the pairwise similarity as in typical graph construction, we attempt to automatically construct a graph *G* and make it well preserve discriminative information based on sparse representation.

Given a set of sample points $X = [x_1, x_2, ..., x_n]$, where $x_i \in \mathbb{R}^m$, i = 1, 2, ..., n, we expect to reconstruct each sample point x_i using as few data points in X as possible. This can be expressed by the following ℓ_0 -minimization problem:

$$\min_{s_i} \|s_i\|_0 \tag{5}$$

s.t.
$$x_i = X s_i$$
,

where $s_i = [s_{i_1}, \dots, s_{i,i-1}, 0, s_{i,i+1}, \dots, s_{i_n}]^T$ is a *n*-dimensional column vector in which the *i*th element is equal to zero, implying the x_i is removed from X, and the element s_{ij} , $j \neq i$ denotes the contribution of x_j for reconstructing x_i . It is well known, (5) is a NP-hard problem. Here, we bypass this difficulty by solving the following ℓ_1 optimization problem³:

$$\begin{array}{l} \min_{s_i} & \|S_i\|_1 \\ \text{s.t.} & x_i = Xs_i, \end{array}$$
(6)

where ℓ_1 is used instead of ℓ_0 . It can be effectively solved by linear programming. Recent researches showed that if the optimal solution sought is sparse enough, the solution of ℓ_0 minimization problem is equal to the solution of ℓ_1 minimization problem (Baraniuk, 2007). After obtaining all of the optimal reconstruction coefficient \hat{s}_i for each x_i , we construct a sparse weight matrix *S* by

$$\mathbf{S} = [\hat{\mathbf{s}}_1, \hat{\mathbf{s}}_2, \dots, \hat{\mathbf{s}}_n]. \tag{7}$$

Then, the new constructed graph $G = \{X, S\}$, where X is the training sample set, S is the edge weight matrix.

In practice, the constraint $x_i = Xs_i$ in (6) does not always hold due to noise or insufficient training samples. We extend it by incorporating a reconstructive compensation term t_i as follows:

$$\min_{s_i} \quad \|s_i\|_1 + \lambda \|t_i\|_p \tag{8}$$

s.t.
$$x_i = Xs_i + t_i$$
,

which is equivalent to

.....

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$$\min_{s_i} \|s_i\|_1 + \lambda \|x_i - Xs_i\|_p, \tag{9}$$

³ In fact, suboptimal solutions can be found by a variety of strategies such as greedy-based (Mallat and Zhang, 1993) and Bayesian-based (Ji et al., 2008) methods. Here, we consider the ℓ_1 strategy simply due to that the equivalence of the ℓ_0 and ℓ_1 problem has been studied deeply from a mathematical perspective.

where $t_i = x_i - Xs_i \in \mathbb{R}^m$ can be seen as a compensation (or error tolerance) for reconstructing x_i . $||t_i||_p$ denotes the ℓ_p -norm, a special measure of the compensation t_i . From the Bayesian viewpoint, $||t_i||_p$ essentially corresponds to different prior distribution (or assumption) about t_i . For example, $||t_i||_2$ is related to Gaussian prior,⁴ while $||t_i||_1$ is related to Laplacian prior.⁵ λ is a regularized parameter to control the trade-off between the sparsity of reconstructive coefficient and the reconstructive compensation. Although the parameter selection problem has been studied in-depth (Mallat and Zhang, 1993; Ji et al., 2008; Hastie, 2009), there is currently no reliable method in theory to assign optimal value for λ . Therefore, we simply set $\lambda = 1$ in all our experiments.

It is worthwhile to point out that the above graph construction manner differs from the one in our previous SPP algorithm (Oiao et al., 2010), though both are motivated by sparse representation (Wright et al., 2009). More specifically, (1) in SPP we used two independent sparse representation models, which are directly developed from (Wright et al., 2009). In contrast, here we reveal the inherent relationship between these models and unify them in one single objective function (8). As a result, the graph construction models behind SPP are just two special instances of (8), and we can develop new graph construction model from the unified objective according to different priors. (2) In SPP, we require sum-to-one constraint as in LLE (Roweis and Saul, 2000). However, we ignore such constraint in SPDA, since we mainly concern discrimination. Not only does this save computational cost, but also, more fortunately, we achieve higher recognition rate than SPP (see Table 4 in Section 5 for details).

3.2. Sparsity preserving regularization

Now we propose the new data-dependent regularizer based on the previously-constructed graph $G = \{X, S\}$. Revisiting the manifold regularizer (2) in SDA, it implies that if x_i and x_j are "close" to each other, then their low-dimensional representation $y_i = w^T x_i$ and $y_j = w^T x_j$ should be close to each other as well. However, for the newly constructed graph G, its edge weight \hat{s}_{ij} is not a rigorous similarity measure, and thus we can not construct the data-dependent regularizer as in SDA.

Note that the relationship between x_i and x_j is characterized by $x_i \approx \sum_{j=1}^{n} \hat{s}_{ij} x_j$ instead of simple "closeness", and hence we expect that their low-dimensional representations y_i and y_j preserve such relationship as well, i.e., $y_i \approx \sum_{j=1}^{n} \hat{s}_{ij} y_j$, which is motivated by LLE (Roweis and Saul, 2000). Therefore, we propose the data-dependent regularizer by minimizing the following objective function:

$$J_{Sparsity}(w) = \sum_{i=1}^{n} \|y_i - Y\hat{s}_i\|^2 = \sum_{i=1}^{n} \|w^T x_i - w^T X\hat{s}_i\|^2,$$
(10)

where $Y = [y_1, y_2, ..., y_n]$ is the low-dimensional representation of the original data. Since the regularizer aims to preserve the sparse reconstructive relationship, we call it *sparsity preserving regularizer*. Then, with simple algebraic formulation (see Appendix), it can be rewritten as

$$w^T X L_s X^T w, \tag{11}$$

where $L_s = I - S - S^T + SS^T$. Although, the data-dependent regularizer can potentially be incorporated into many semi-supervised learning algorithms, we only focus on SSDR in this paper.

3.3. Sparsity preserving discriminant analysis (SPDA)

Similar to SDA, we extend LDA⁶ to semi-supervised version based on the newly proposed data-dependent regularizer. Naturally, the objective function can be defined as follows:

$$\max_{w} \frac{w^{t} S_{b} w}{w^{T} (S_{t} + \lambda_{1} I + \lambda_{2} X L_{s} X^{T}) w},$$
(12)

where S_b and S_t are respectively the inter-class and total scatter matrices, which are calculated just using the labeled training samples. *I* is an identity matrix related to Tikhonov regularizer, and $w^T X L_s X^T w$ is the sparsity preserving regularizer. The solution of (12) can be easily achieved by the following generalized eigenvalue problem:

$$S_b w = \eta (S_t + \lambda_1 I + \lambda_2 X L_s X^T) w.$$
⁽¹³⁾

The algorithmic procedure is shown as follows. Concretely, we assume the training sample set $X = [x_1, \ldots, x_l, x_{l+1}, \ldots, x_{l+u}] = [X_L, X_U]$, where the first *l* training samples $\{x_i\}_{i=1}^l$ are labeled and from *c* classes (there are l_k samples in the *k*th class), the last *u* training samples $\{x_i\}_{i=1}^{l+u}$ are unlabeled. Without loss of generality, the sample points in X_L are ordered according to their labels.

Algorithm 1. Sparsity Preserving Discriminant Analysis

- Step 1. Calculate $S_b = X_L H_L X_L^T$ and $S_t = X_L X_L^T$ based on the labeled training samples in X_L , where, $H_L = diag(H^1, H^2, ..., H^c)$ is a block-diagonal matrix, and H^k is a $l_k \times l_k$ matrix with all elements equal to $1/l_k$.
- Step 2. Construct graph $G = \{X, S\}$. The weight matrix S is calculated based on all training samples in X using (6) or (8).
- Step 3. Calculate the data-dependent (sparsity preserving) regularizer $w^T X L_s X^T w$, where $L_s = I S S^T + S^T S$.
- Step 4. Calculate the projections by the generalized eigenvalue problem (13), and the projection matrix $W = [w_1, w_2, ..., w_d]$, where w_i are the eigenvectors corresponding to the largest d eigenvalues.

4. Extensions of SPDA

In this section, we extend the proposed algorithm to its kernelized version (for improving the flexibility of SPDA) and ensemble version (for reducing the computational complexity), respectively.

4.1. Kernel SPDA

As described above, SPDA only focuses on linear dimensionality reduction, and thus it may fail to deal with the highly nonlinear structure in data. Fortunately, we can easily extend SPDA to perform in reproducing kernel hilbert space (RKHS) like other graphbased dimensionality reduction algorithms.

Let $\phi: x \to F$ be a function mapping the data points in the input space to the feature space. According to the kernel trick, we expect to replace the explicit mapping with the inner product $K(x_i, x_j) = (\phi(x_i) \cdot \phi(x_j))$. Furthermore, we assume $\phi_L = [\phi(x_1), \phi(x_2), \dots, \phi(x_l)]$, $\phi_U = [\phi(x_{l+1}), \phi(x_{l+2}), \dots, \phi(x_n)]$ and $\phi = [\phi_L, \phi_U]$, then the inter-class scatter matrix and the total scatter matrix in the feature space can respectively be denoted as

$$S_{b}^{F} = \phi_{L}H_{L}\phi_{L}^{T} = \phi \begin{pmatrix} H_{L} & 0 \\ 0 & 0 \end{pmatrix} \phi^{T} = \phi H\phi^{T}, \ S_{t}^{F} = \phi_{L}\phi_{L}^{T} = \phi \begin{pmatrix} I & 0 \\ 0 & 0 \end{pmatrix} \phi^{T} = \phi \widetilde{I}\phi^{T},$$
(14)

where H_L is defined as in the SPDA algorithm.

⁴ In this sense, the (9) has the same mathematical expression as the popular LASSO (Tibshirani, 1996) in statistics.

⁵ It has been validated applicable to face images with partial occlusion (Wright et al., 2009). In this paper, we use this prior by empirically modeling the variations of expression and illumination as partial corruption on clear face images.

⁶ Of course, we can consider other discriminant criteria such as MMC if necessary.

According to the Representer Theorem (Scholkopf et al., 2001), the projection w^F sought in feature space can be expressed as $w^F = \phi \alpha$, where $\alpha = [\alpha_1, \alpha_2, ..., \alpha_n]^T$ is a coefficient vector that represents w^F in the feature space. Let $K = \phi^T \phi$ be the kernel matrix, the objective function of kernel SPDA can be expressed as follows:

$$\max_{w} \frac{\alpha^{T} K H K^{T} \alpha}{\alpha^{T} (K \widetilde{I} K^{T} + \lambda_{1} K + \lambda_{2} K L_{s} K^{T}) \alpha}.$$
(15)

The optimal solution $\hat{\alpha}$ can be obtained by solving the following generalized eigenvalue problem:

$$KHK^{T}\alpha = \eta(KIK^{T} + \lambda_{1}K + \lambda_{2}KL_{s}K^{T})\alpha.$$
(16)

Thus, given a new data point *x*, its low-dimensional representation is $(w^F)^T \phi(x) = \hat{\alpha}^T K(\cdot, x)$, where $K(\cdot, \cdot)$ is a kernel function.

4.2. Ensemble SPDA

According to (Wright et al., 2009), the sparsity of the ideal solution sought is mainly related to the class numbers rather than the sample numbers. Therefore, intuitively, a large number of unlabeled samples do not necessarily help improve the performance of SPDA significantly, and conversely incur high computational burden since SPDA constructs graph based on all the training samples. Here, we introduce a very simple ensemble strategy to speed up the proposed SPDA algorithm based on the above observation.

In particular, given a set of training samples $X = [x_1, ..., x_l, x_{l+1}, ..., x_{l+u}] = [X_L, X_U]$ as mentioned before, we randomly partition the unlabeled sample set X_U into q small sample sets X_{U1} , $X_{U2}, ..., X_{Uq}$, and thus we can generate a series of new training sets $X_1 = [X_L, X_{U1}], ..., X_q = [X_L, X_{Uq}]$. Then, we perform SPDA on each new training sets and the test sample is classified by voting strategy. Concretely, the ensemble SPDA algorithmic procedure is shown as follows.

Algorithm 2. Ensemble SPDA

Step 1. Partition the training set *X* into *q* sub-sets X_1, X_2, \ldots, X_q .

Step 2. Implement SPDA on each sub-set, and get *q* subspaces.

Step 3. Project the test sample *x* onto each subspace, and then

implement classification (e.g., 1NN) on each subspace.

Step 4. Vote to decide the class label of the test sample.

5. Experiments

5.1. Illustrative examples

In this subsection, we intuitively illustrate why the proposed algorithm might work well through two illustrative experiments on toy (5.1.1) and face (5.1.2) data sets, respectively.

5.1.1. Illustrative experiment on toy data

For simplicity of our illustration here, we only consider binary classification problem and assume that each class lies in a onedimensional subspace embedding in three-dimensional ambient space. We randomly sample three (one labeled and two unlabeled) data points from per class for training. Fig. 1a gives an instance of so-generated training sample points which are respectively signed with pentacle and square. In order to approximate practical problem, the data points are corrupted by Gaussian additive white noise with standard deviation 0.1.

Based on the training data, we construct the typical neighborhood graph and the sparse reconstruction graph, respectively. In particular, Fig. 1b gives the neighborhood graph, where the neighborhood size k = 2. It is easy to see that the edges on the graph link the data points which are close to each other, yet from different class. Obviously, other locality-oriented graph construction manners such as the one involved in LLE may also suffer from the fact that the samples from different class are close to each other. In contrast, Fig. 1c gives the sparse reconstruction graph behind SPDA. With the sparsity constraint, the non-zero reconstructive coefficients for a given data point more possibly match the data points in the same class. As a result, so-constructed graph tends to contain more discriminative information than typical neighborhood graph. More specifically, we classify 100 randomly generated test samples using 1-nearest neighbor (1NN) classifier on the obtained one-dimensional subspace by SDA and SPDA, respectively. The average classification accuracies corresponding to SDA and SPDA are 70.34% and 87.22%, respectively.

5.1.2. Illustrative experiment on face data

The previous toy problem showed that locality-oriented graph construction manners may affect the performance significantly. How about on real-world data sets? Here, we take AR database⁷ as an example to compare the proposed algorithm with LLE, since the graph behind SPDA is constructed by ℓ_1 -minimization optimization problem which is closely related to the least square graph construction hidden in LLE.

More specifically, we assume the face data set $X = [x_1, x_2, ..., x_n]$, where the samples are ordered according to their labels for the convenience of illustration. Then, given a face image $x_i \in X$, by solving (8) or (9), we obtain a sparse reconstruction coefficient s_i in which the non-zero values model the contribution of each support point⁸ or support face to represent x_i . In other words, simply by solving (8) or (9), we get both the graph and its corresponding edge weights simultaneously, which is contrary to the scheme of LLE, where the graph and its edge weights are estimated separately. In particular, for a specific face image, Fig. 2 gives an illustration of support faces and corresponding coefficient found by SPDA and those by LLE.

From the experiment result, we note that the support faces found by SPDA with a ℓ_1 -minimization criterion are more discriminative than those by LLE with the least square criterion – two of three faces with the same identity as the prototype are correctly found by SPDA scheme.

5.2. One (labeled) training image face recognition

In this subsection, we perform one training image face recognition experiments on three publicly available face databases: CMU PIE, Extended Yale B and AR databases.

5.2.1. Database description

CMU PIE face database contains 68 subjects with 41,368 face images as a whole. The face images were captured under varying pose, illumination and expression. Similar to (Cai et al., 2007), in the experiment, we choose the frontal pose (C27) with varying lighting which leaves us 43 images per person. The size of each face image is cropped to have 32×32 pixels as shown in Fig. 3(top).

Extended Yale B database (Lee et al., 2005) contains 2414 frontview face images of 38 individuals. For each individual, about 64 pictures were taken under various laboratory-controlled lighting

⁷ See the next subsection for the description about this database. Here we just used the face images taken in the first session.

 $^{^{\}mbox{8}}$ Here, support point denotes the face image which contributes to represent the given face image.



Fig. 2. Illustration of three support faces with corresponding coefficients (bottom row) for a given face image (the first image on the left side of the approximately equal mark in bottom row) and the reconstruction coefficient distribution (upper row) using SPDA (left) and LLE (right), respectively.

conditions. In our experiments, we simply use the cropped images⁹ with the resolution of 32×32 as shown in Fig. 3(middle). The database may be substantially more challenging than the above PIE database due to much larger illumination variations.

AR database consists of over 4000 face images of 126 individuals. For each individual, 26 pictures were taken in two sessions (separated by two weeks) and each section contains 13 images. These images include front view of faces with different expressions, illuminations and occlusions. In our experiments, we only use the images without occlusion in the AR face database provided and preprocessed by Martinez and Kak (2001). This sub-dataset contains 1400 face images corresponding to 100 person (50 men and 50 women), where each person has 14 different images taken in two sessions. The original resolution of these image faces is 165×120 . Here, for computational convenience, we resize them to 66×48 as shown in Fig. 3(bottom).

5.2.2. Experimental setting

On each face database, we perform two groups of experiments with different unlabeled training sample numbers. Table 2 gives the specific experimental setting. For example, for PIE database, experiment 1 denotes that three images are randomly selected from each class as the training set, and the rest images as the testing set. Among the three training images, only one image is randomly selected and labeled, which leaves the rest two images unlabeled; while, in experiment 2, the labeled training samples keep the same, but the number of the total training samples per subject increases to 30. In fact, the experiment 2 is also considered in (Cai et al., 2007) where the authors justified that their proposed SDA algorithm achieved better performance than some popular algorithms, e.g., LPP (He and Niyogi, 2003) and LapSVM (Belkin et al., 2006). For all the experiments here, we report the averaged results over 30 random training/test splits.

Based on each data partition, we compare SPDA with Baseline, unsupervised SPP, supervised LDA¹⁰ and semi-supervised SDA. The baseline approach denotes the 1-NN classifier on the original face space without dimensionality reduction. For LDA, the face subspace is learnt only using the labeled samples; for SPP, the face subspace is learnt using all the training samples without label information; for SDA and SPDA, the face subspace is learnt using both labeled and unlabeled samples. Then, based on the learnt subspace, 1-NN classifier is employed to evaluate the recognition rate on the test data. As descript previously, SPDA suffers from high computational cost when a large number of unlabeled samples are considered. Therefore, for the experiment 2 on each database, we adopt the ensemble version of SPDA where the unlabeled samples are simply and randomly partitioned into q = 10 small sub-sets.

5.2.3. Parameter selection

LDA and SPP are both parameter-free. SDA contains four parameters: two regularized parameters and two free parameters for graph construction. Here, we use the same parameter values for SDA as in (Cai et al., 2007). For convenience of comparison, the two regularized parameters in SPDA are assigned the same values as in SDA. In addition, for all the above algorithms, the subspace dimension is set to c - 1, where c is the class number. Table 3 gives specific parameter values for SDA and SPDA.

⁹ We directly download the cropped image data from http://www.cs.uiuc.edu/ homes/dengcai2.

¹⁰ Strictly speaking, under the single training sample case, typical LDA fails to work since the intra-class variation cannot be obtained. Here, we simply replace the intraclass scatter matrix using a constant matrix as in (Zhao et al., 1999).



Fig. 3. Some face images from PIE (top), Yale B (middle) and AR (bottom) databases.

Table	2	

Data set description and partition.

Database	Samples sizes per class	Class numbers	Experiment	Experiment 1		Experiment		
			Train	Labeled	Train	Labeled		
PIE	64	68	3	1	30	1		
Yale B	43	38	3	1	30	1		
AR	14	100	3	1	10	1		

Table 3

Parameter setting for SDA and SPDA.

Algorithms	Reg. para. λ_1	Reg. para. λ_2	Neighbor k	Edge weights
SDA	0.01	0.1	2	Cosine
SPDA	0.01	0.1	Auto	Auto

5.2.4. Experimental results and overall observations

Based on the above experimental setting, Table 4 reports the classification accuracies corresponding to different algorithms and databases, where E1 and E2 denote the first and second groups of experiments, respectively.

From the experimental results on the three popular face databases, we can achieve several observations as follows:

- (1) Among the discussed dimensionality reduction methods, LDA generally achieve relatively low accuracies due to the fact that only one labeled sample per class is used to learning the face subspace.
- (2) Despite its unsupervised nature, SPP can outperform LDA with the help of extra training samples. However, the performance of SPP does not always be improved with the increase of training samples. Interestingly, SPP can even achieve better performance than SDA in some of the experiments, which benefits from the natural discriminative power of sparse representation.
- (3) Semi-supervised SDA and the proposed SPDA always outperform LDA if considerable unlabeled training samples are available. That is, the extra unlabeled training samples can generally help improve the performance.

Table 4

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	Baseline (%)	LDA (%)	SPP (%)	SDA (%)	SPDA (%)
PIE					
E1	25.88 ± 1.2	25.88 ± 1.2	62.55 ± 2.0	30.91 ± 2.1	67.47 ± 1.8
E2	26.21 ± 1.6	26.21 ± 1.6	51.29 ± 3.1	59.57 ± 3.2	70.44 ± 3.0
Yale	В				
E1	12.60 ± 1.2	12.60 ± 1.2	17.95 ± 3.1	16.1 ± 1.6	31.27 ± 3.6
E2	13.01 ± 1.4	13.01 ± 1.4	14.28 ± 3.2	26.77 ± 2.5	35.44 ± 3.3
AR					
E1	24.55 ± 1.4	24.55 ± 1.4	44.57 ± 2.6	22.48 ± 1.6	58.46 ± 2.0
E2	24.69 ± 2.4	24.69 ± 2.4	55.06 ± 3.1	26.22 ± 2.1	61.23 ± 2.5

(4) SPDA consistently outperforms SPP and SDA on all the used face databases. This illustrates both label information and well-constructed graph (or equivalently, data-dependent regularizer) play important roles in the ultimate recognition rates. More importantly, the proposed SPDA algorithm can remarkably improve the performance of LDA even when only few unlabeled training samples are available.

6. Conclusion and future works

In this paper, we developed a new semi-supervised dimensionality reduction method called sparsity preserving discriminant analysis (SPDA). The newly proposed algorithm does not only model the "locality" automatically, but also remarkably improves the performance of typical LDA only resorting to very few additional unlabeled samples. As a result, SPDA algorithm is more applicable to face recognition problem with only a few training samples.

From the experimental results, we can find that SPDA is more effective than the popular SDA algorithm, but has still a big gap from practical face recognition applications. Therefore, in the future work, we will attempt to integrate the typical strategies (e.g., synthesizing virtual samples, localizing the training images) with the proposed algorithm and expect to further improve the performance.

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Appendix. The formulation for sparsity preserving regularizer

$$J_{Sparsity}(w) = \sum_{i=1}^{n} \|y_i - Y\hat{s}_i\|^2 = \sum_{i=1}^{n} \|w^T x_i - w^T X\hat{s}_i\|^2$$
$$= w^T \left(\sum_{i=1}^{n} (x_i - X\hat{s}_i)(x_i - X\hat{s}_i)^T\right) w.$$

Let e_i be a *n*-dimensional unit vector with the *i*th element 1, 0 otherwise, then the above equation is equal to

$$w^{T}\left(\sum_{i=1}^{n} (Xe_{i} - X\hat{s}_{i})(Xe_{i} - X\hat{s}_{i})^{T}\right)w$$

= $w^{T}X\left(\sum_{i=1}^{n} (e_{i} - \hat{s}_{i})(e_{i} - \hat{s}_{i})^{T}\right)X^{T}w$
= $w^{T}X\left(\sum_{i=1}^{n} e_{i}e_{i}^{T} - \hat{s}_{i}e_{i}^{T} - e_{i}\hat{s}_{i}^{T} + \hat{s}_{i}\hat{s}_{i}^{T}\right)X^{T}w = w^{T}XL_{s}X^{T}w$

where $L_{s} = I - S - S^{T} + SS^{T}$ and $S = [\hat{s}_{1}, \hat{s}_{2}, ..., \hat{s}_{n}]$.

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