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# Research Article **Evolutionary Discriminant Feature Extraction with Application to Face Recognition**

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Evolutionary computation algorithms have recently been explored to extract features and applied to face recognition. However these methods have high space complexity and thus are not efficient or even impossible to be directly applied to real world applications such as face recognition where the data have very high dimensionality or very large scale. In this paper, we propose a new evolutionary approach to extracting discriminant features with low space complexity and high search efficiency. The proposed approach is further improved by using the bagging technique. Compared with the conventional subspace analysis methods such as PCA and LDA, the proposed methods can automatically select the dimensionality of feature space from the classification viewpoint. We have evaluated the proposed methods in comparison with some state-of-the-art methods using the ORL and AR face databases. The experimental results demonstrated that the proposed approach can successfully reduce the space complexity and enhance the recognition performance. In addition, the proposed approach provides an effective way to investigate the discriminative power of different feature subspaces.

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## 1. Introduction

Biometrics has become a promising technique for personal authentication. It recognizes persons based on various traits, such as face, fingerprint, palmprint, voice, and gait. Most biometric systems use the images of those traits as inputs [1]. For example, 2D face recognition systems capture facial images from persons and then recognize them. However, there are many challenges in implementing a real-world face recognition system [2-4]. A well-known challenge is the "curse of dimensionality," which is also a general problem in pattern recognition [5]. It refers to the fact that as the dimension of data increases, the number of samples required for estimating the accurate representation of the data grows exponentially. Usually, the spatial resolution of a face image is at least hundreds of pixels and usually will be tens of thousands. From the statistical viewpoint, it will require tens of thousands of face samples to deal with the face recognition problem. However, it is often very difficult, even impossible, to collect so many samples. The dimensionality reduction techniques, including feature

selection and extraction, are therefore widely used in face recognition systems to solve or alleviate this problem. In this paper, we will present a novel evolutionary computationbased approach to dimensionality reduction.

The necessity of applying feature extraction and selection before classification has been well demonstrated by researchers in the realm of pattern recognition [5, 6]. The original data are often contaminated by noise or contain much redundant information. Direct analysis on them could then be biased and result in unsatisfied classification accuracy. On the other hand, the raw data are usually of very high dimensionality. Not only does this lead to expensive computational cost but also causes the "curse of dimensionality." This may lead to poor performance in applications such as face recognition.

Feature extraction and selection are slightly different. Feature selection seeks for a subset of the original features. It does not transform the features but prunes some of them. Feature extraction, on the other hand, tries to acquire a new feature subset to represent the data by transforming the original data. Mathematically, given an  $n \times N$  sample

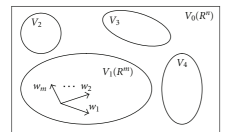


FIGURE 1: Linear feature extraction: from the subspace viewpoint.

matrix  $X = [x_1x_2\cdots x_N]$  (*n* is the original dimension of samples, and *N* is the number of samples), a linear feature extraction algorithm could use an  $n \times m$  transform matrix *W* to transform the data as  $Y = W^T X = [y_1y_2\cdots y_N]$ , where "*T*" is the transpose operator. Here,  $0 < m \ll n$  is the dimension of the transformed feature subspace. Figure 1 illustrates this process. Suppose that the original data lie in the *n*-dimensional space  $V_0$ . Feature extraction is then to find out one of its subspaces which has the best discriminability and is called feature subspace, say the *m*-dimensional space  $V_1$ . In linear cases, an optimal projection basis of the feature subspace,  $\{w_1, w_2, \ldots, w_m \in \mathbb{R}^n\}$ , can be calculated such that certain criterion is optimized. These basis vectors compose the column vectors in the transform matrix *W*.

Feature extraction is essentially a kind of optimization problem, and several criteria have been proposed to steer the optimization, for example, minimizing reconstruction error, maximizing reserved variance while reducing redundancy, and minimizing the within-class scatterance while maximizing the between-class scatterance, and so forth. Using such criteria, many feature extraction algorithms have been developed. Two well-known examples are Principal Component Analysis (PCA) [7] and Linear Discriminant Analysis (LDA) [5]. They represent two categories of subspace feature extraction methods [8] that are widely used in face recognition [3, 9–17]. In the context of face recognition, various feature subspaces have been studied [16, 17], for example, the range space of  $S_b$  and the null space of  $S_w$ . Here,  $S_b$  and  $S_w$  are the between-class and within-class scatter matrixes, defined as  $S_b = (1/N) \sum_{j=1}^{L} N_j (M_j - M) (M_j - M)$  $(M)^T$  and  $S_w = (1/N) \sum_{j=1}^L \sum_{i \in I_j} (x_i - M_j) (x_i - M_j)^T$ , where  $M = \sum_{i=1}^{N} x_i / N$  is the mean of all the N training samples, and  $M_j = \sum_{i \in I_j} x_i / N_j$  is the mean of samples in the *j*th class (j = 1, 2, ..., L). A significant issue involved in these methods is how to determine *m*, that is, the dimension of the feature subspace. Unfortunately, neither PCA nor LDA gives systematic ways to determine the optimal dimension in the sense of classification accuracy. Currently, people usually choose the dimension by experience [9, 10, 18]. For example, the dimensionality of PCA-transformed space is set to 20 or 30 or (N - 1), where "N" is the number of samples, and the dimensionality of LDA-transformed space is set to (L-1), where "L" is the number of classes. However, such method does not necessarily guarantee the best classification performance as we will show in our experiments. In addition, it is impractical or too expensive to search the whole solution

space blindly in real applications such as face recognition because of the very high dimensionality of the original data.

Recently, some researchers [18-32] have explored the use of evolutionary computation (EC) methods [28] for feature selection and extraction. In these methods, the solution space is searched in guided random way, and the dimensionality of the feature subspace is automatically determined. Although these methods successfully avoid the manual selection of feature subspace dimensionality and good results have been reported on both synthetic and realworld datasets, most of them have very high space complexity and are often not applicable for high dimensional or large scale datasets [29]. In this paper, by using genetic algorithms (GA) [30], we will propose an evolutionary approach to extracting discriminant features for classification, namely, evolutionary discriminant feature extraction (EDFE). The EDFE algorithm has low space complexity and high search efficiency. We will further improve it by using the bagging technique. Comprehensive face recognition experiments have been performed on the ORL and AR face databases. The experimental results demonstrated the success of the proposed algorithms in reducing the space complexity and enhancing the recognition performance. In addition, the proposed method provides a way to experimentally compare the discriminability of different subspaces. This will benefit both researchers and engineers in analyzing and determining the best feature subspaces.

The rest of this paper is organized as follows. Sections 2 and 3 introduce in detail the proposed EDFE and bagging EDFE (BEDFE) algorithms. Section 4 shows the face recognition experimental results on the ORL and AR face databases. Section 5 gives some discussion on the relation between the proposed approach and relevant methods. Finally, Section 6 concludes the paper.

#### 2. Evolutionary Discriminant Feature Extraction (EDFE)

This section presents the proposed EDFE algorithm, which is based on GA and subspace analysis. Algorithm 1 shows the procedures of EDFE.

2.1. Data Preprocessing: Centralization and Whitening. All the data are firstly preprocessed by centralization, that is, the total mean is subtracted from them:

$$\overline{x}_i = x_i - M, \quad i = 1, 2, \dots, N.$$
 (1)

The centralization applies a translational transformation to the data so that their mean is moved to the origin. This helps to simplify subsequent processing without loss of accuracy.

Generally, the components of data could span various ranges of values and could be of high order of magnitude. If we calculate distance-based measures like scatterance directly on the data, the resulting values can be of various orders of magnitude. As a result, it will be difficult to combine such measures with others. This is particularly serious in defining fitness for GA-based methods. Therefore, we further whiten the centralized data to normalize their variance to unity. Step 1. Preprocess the data using whitened principal component analysis (WPCA). - Centralization - Whitening Step 2. Calculate a search space for the genetic algorithm (GA). - For example, the null space of  $S_w$  and the range space of  $S_b$ - Heuristic knowledge can be used in defining search spaces Step 3. Use GA to search for an optimal projection basis in the search space defined in Step 2. 3.1. Randomly generate a population of candidate projection bases. 3.2. Evaluate all individuals in the population using a predefined fitness function. 3.3. Generate a new population using selection, crossover and mutation according to the fitness of current individuals. 3.4. If the stopping criterion is met, retrieve the optimal projection basis from the fittest individual and proceed to Step 4; otherwise, go back to 3.2 to repeat the evolution loop. Step 4. Use a classifier to classify new samples in the feature subspace obtained in Step 3. - For example, Nearest Mean Classifier (NMC)

ALGORITHM 1: Procedures of the proposed EDFE algorithm.

This is done by the eigenvalue decomposition (EVD) on the covariance matrix of data. Let  $\lambda_1 \ge \lambda_2 \ge \cdots \ge \lambda_n (\ge 0)$  be the eigenvalues of  $S_t = (1/N) \sum_{i=1}^N (x_i - M)(x_i - M)^T$  and  $\alpha_1, \alpha_2, \ldots, \alpha_n$  the corresponding eigenvectors. The whitening transformation matrix is then

$$W_{\text{WPCA}} = \left[\frac{\alpha_1}{\sqrt{\lambda_1}} \frac{\alpha_2}{\sqrt{\lambda_2}} \cdots \frac{\alpha_{N-1}}{\sqrt{\lambda_{N-1}}}\right].$$
 (2)

Here, we set the dimensionality of the whitened space to (N-1), the rank of the covariance matrix. This means that we keep all the directions with nonzero variances, which ensures that no potential discriminative information is discarded from the data in whitening. Let  $\overline{X}$  and  $\widetilde{X}$  be the centralized and whitened data, then we have

$$\widetilde{X} = W_{\text{WPCA}}^T \overline{X}.$$
(3)

It can be easily proven that  $\tilde{S}_t = (1/N)\tilde{X}\tilde{X}^T = I_{N-1}$ , where  $I_{N-1}$  is the (N - 1) dimensional identity matrix. In addition to normalizing the data variance, this whitening process also decorrelates the data components. For simplicity, we denote the preprocessed data in the whitened space still by X, omitting the tildes.

2.2. Calculating the Constrained Search Space. Unlike existing GA-based feature extraction algorithms, the proposed EDFE algorithm imposes some constraints on the search space so that the GA can search more efficiently in the constrained space. This idea originates from the fact that guided search, given correct guidance, is always better than blind search. It is widely accepted that heuristic knowledge, if properly used, could significantly improve the performance of systems. Keeping this in mind, we combine the EDFE algorithm with a scheme of constraining the search space as follows.

According to the Fisher criterion

$$W_{\rm LDA} = \arg \max_{W} \left\{ J_{\rm LDA}(W) = \frac{W^T S_b W}{W^T S_w W} \right\},\tag{4}$$

the most discriminative directions are most probably lying in the subspaces generated from  $S_w$  and  $S_b$ . Researchers [16, 17] have investigated the null space of  $S_w$ , denoted by null( $S_w$ ), and the range space of  $S_b$ , denoted by range( $S_b$ ), using analytical methods. It can be proved that the solution to (4) lies in these subspaces. We will use the EDFE algorithm to search for discriminant projection directions in null( $S_w$ ), range( $S_w$ ), and range( $S_b$ ), respectively, and compare their discriminability in recognizing faces. In this section, we present a method to calculate these three spaces. If some other subspace is considered, it is only needed to take its basis as the original basis of the search space.

Before proceeding to the detailed method of calculating the basis for null( $S_w$ ), range( $S_w$ ), and range( $S_b$ ), we first give the definitions of these three subspaces as follows

$$\operatorname{null}(S_w) = \{ v \mid S_w v = 0, \ S_w \in \mathbb{R}^{n \times n}, \ v \in \mathbb{R}^n \},$$
(5)

$$\operatorname{range}(S_w) = \{ v \mid S_w v \neq 0, \ S_w \in \mathbb{R}^{n \times n}, \ v \in \mathbb{R}^n \}, \quad (6)$$

$$\operatorname{range}(S_b) = \{ v \mid S_b v \neq 0, \ S_b \in \mathbb{R}^{n \times n}, \ v \in \mathbb{R}^n \}.$$
(7)

According to the definitions of  $S_w$  and  $S_b$ , the ranks of them are, respectively,

$$\operatorname{rank}(S_w) = \min\{n, N - L\}, \quad \operatorname{rank}(S_b) = \min\{n, L - 1\}.$$
(8)

These ranks determine the numbers of vectors in the bases of range( $S_w$ ), null( $S_w$ ), and range( $S_b$ ). Next, we introduce an efficient method to calculate the basis.

To get a basis of  $range(S_w)$ , we use the EVD again. However, in real applications of image recognition, the dimensionality of data, *n*, is often very high. This makes it computationally infeasible to conduct EVD directly on  $S_w \in \mathbb{R}^{n \times n}$ . Instead, we calculate the eigenvectors of  $S_w$  from another  $N \times N$  matrix  $S'_w$  [9]. Let

$$H_w = [x_1 x_2 \cdots x_N] \in \mathbb{R}^{n \times N}, \qquad (9)$$

then

$$S_w = \frac{1}{N} H_w H_w^T.$$
(10)

Note that the data have already been centralized and whitened. Let

$$S'_w = \frac{1}{N} H^T_w H_w, \tag{11}$$

and suppose  $(\lambda, \alpha')$  to be an eigenvalue and the associated eigenvector of  $S'_w$ , that is,

$$S'_{w}\alpha' = \lambda\alpha'. \tag{12}$$

Substituting (7) into (8) gives

$$\frac{1}{N}H_{w}^{T}H_{w}\alpha' = \lambda\alpha'.$$
(13)

Multiplying both sides of (9) with  $H_w$ , we have

$$\frac{1}{N}H_{w}H_{w}^{T}H_{w}\alpha' = \lambda H_{w}\alpha'.$$
(14)

With (10) and (6), there is

$$S_{w} \cdot (H_{w}\alpha') = \lambda \cdot (H_{w}\alpha'), \qquad (15)$$

which proves that  $(\lambda, H_w \alpha')$  are the eigenvalue and eigenvector of  $S_w$ . Therefore, we first calculate the rank $(S_w)$  dominant eigenvectors of  $S'_w$ ,  $(\alpha'_1, \alpha'_2, \ldots, \alpha'_{rank}(S_w))$ , which have largest positive associated eigenvalues. A basis of range $(S_w)$  is then given by

$$B_{\text{range}}(S_w) = \{ \alpha_i = H_w \alpha'_i \mid i = 1, 2, \dots, \text{rank}(S_w) \}.$$
(16)

The basis of range( $S_b$ ) can be calculated in a similar way. Suppose that the *N* column vectors of  $H_b \in \mathbb{R}^{n \times N}$  consist of  $M_j$  (j = 1, 2, ..., L) with  $N_j$  entries, then  $S_b = (1/N)H_bH_b^T$ . Let  $S'_b = (1/N)H_b^TH_b$  and { $\beta'_i \mid i = 1, 2, ..., \operatorname{rank}(S_b)$ } be its rank( $S_b$ ) dominant eigenvectors. The basis of range( $S_b$ ) is then

$$B_{\text{range}}(S_b) = \{\beta_i = H_b \beta'_i \mid i = 1, 2, \dots, \text{rank}(S_b)\}.$$
 (17)

Based on the basis of range( $S_w$ ), it is easy to get the basis of null( $S_w$ ) through calculating the orthogonal complement space of range( $S_w$ ).

#### 2.3. Searching: An Evolutionary Approach

2.3.1. Encoding Individuals. Binary individuals are widely used owing to their simplicity; however, the specific definition is problem dependent. As for feature extraction,

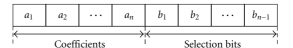


FIGURE 2: The individual defined in EDFE. Each coefficient is represented by 11 bits.

it depends on how the projection basis is constructed. The construction of projection basis vectors is to generate candidate transformation matrixes for the GA algorithm. Usually, the whole set of candidate projection basis vectors are encoded in an individual. This is the reason why the space complexity of existing GA-based feature extraction algorithms is so high. For example, in EP [18], one individual has  $(5n^2 - 4n)$  bits, where *n* is the dimensionality of the search space. In order to reduce the space complexity and make the algorithm more applicable for high dimensional data, we propose to construct projection basis vectors using the linear combination of the basis of search space and the orthogonal complement technique. As a result, only one vector is needed to encode for each individual.

First, we generate one vector via linearly combining the basis of the search space. Suppose that the search space is  $\mathbb{R}^n$  and let  $\{e_i \in \mathbb{R}^n \mid i = 1, 2, ..., n\}$  be a basis of it, and let  $\{a_i \in \mathbb{R} \mid i = 1, 2, ..., n\}$  be the linear combination coefficients. Then we can have a vector as follows:

$$v = \sum_{i=1}^{n} a_i e_i. \tag{18}$$

Second, we calculate a basis of the orthogonal complement space in  $\mathbb{R}^n$  of  $V = \operatorname{span}\{v\}$ , the space expanded by v. Let  $\{u_i \in \mathbb{R}^n \mid i = 1, 2, ..., n - 1\}$  be the basis, and  $U = \operatorname{span}\{u_1, u_2, ..., u_{n-1}\}$ , then

$$R^n = V \oplus U, \tag{19}$$

where " $\oplus$ " represents the direct sum of vector spaces, and

$$U = V^{\perp}, \tag{20}$$

where " $\perp$ " denotes the orthogonal complement space. Finally, we randomly choose part of this basis as the projection basis vectors.

According to the above method of generating projection basis vectors, the information encoded in an individual includes the *n* combination coefficients and (n - 1) selection bits. Each coefficient is represented by 11 bits with the leftmost bit denoting its sign ("0" means negative and "1" positive) and the remaining 10 bits representing its value as a binary decimal. Figure 2 shows such an individual, in which the selection bits  $b_1, b_2, \ldots, b_{n-1}$ , taking a value of "0" or "1," indicate whether the corresponding basis vector is chosen as a projection basis vector or not. The individual under such definition has (12n - 1) bits. Apparently, it is much shorter than that by existing GA-based feature extraction algorithms (such as EP), and consequently the proposed EDFE algorithm has a much lower space complexity. 2.3.2. Evaluating Individuals. We evaluate individuals from two perspectives, pattern recognition and machine learning. Our ultimate goal is to accurately classify data. Therefore, from the perspective of pattern recognition, an obvious measure is the classification accuracy in the obtained feature subspace. In fact, almost all existing GA-based feature extraction algorithms use this measure in their fitness functions. They calculate this measure based on the training samples or a subset of them. However, after preprocessing the data using WPCA, the classification accuracy on the training samples is always almost 100%. In [26], Zheng et al. also pointed this out when they used PCA to process the training data. They then simply ignored its role in evaluating individuals. Different from their method, we keep this classification term in the fitness function but use a validation set, instead of the training set. Specifically, we randomly choose from the  $N_{\rm va}$  samples to create a validation set  $\Omega_{\rm va}$  and use the remaining  $N_{\rm tr} = (N - L \times N_{\rm va})$  samples as the training set  $\Omega_{\rm tr}$ . Assume that  $N_{\rm va}^c(D)$  samples in the validation set are correctly classified in the feature subspace defined by the individual D on the training set  $\Omega_{tr}$ ; the classification accuracy term for this individual is then defined as

$$\zeta_a(D) = \frac{N_{\rm va}^c(D)}{N_{\rm va}}.$$
(21)

From the machine learning perspective, the generalization ability is an important index of machine learning systems. In previous methods, the between-class scatter is widely used in fitness functions. However, according to the Fisher criterion, it is better to simultaneously minimize the within-class scatter and maximize the between-class scatter. Thus, we use the following between-class and within-class scatter distances of samples in the feature subspace:

$$d_{b}(D) = \frac{1}{N} \sum_{j=1}^{L} N_{j} \left( M_{j} - M \right)^{T} \left( M_{j} - M \right),$$

$$d_{w}(D) = \frac{1}{L} \sum_{j=1}^{L} \frac{1}{N_{j}} \sum_{i \in I_{j}} (y_{i} - M_{j})^{T} \left( y_{i} - M_{j} \right)$$
(22)

to measure the generalization ability as

$$\zeta_g(D) = d_b(D) - d_w(D). \tag{23}$$

Here, *M* and  $M_j$ , j = 1, 2, ..., L, are calculated based on  $\{y_i \mid i = 1, 2, ..., N\}$  in the feature subspace.

Finally we define the fitness function as the weighted sum of the above two terms:

$$\zeta(D) = \pi_a \zeta_a(D) + (1 - \pi_a) \zeta_g(D), \qquad (24)$$

where  $\pi_a \in [0, 1]$  is the weight. The accuracy term  $\zeta_a$  in this fitness function lies in interval [0, 1]. Thus, it is reasonable to make the value of the second generalization term  $\zeta_g$  be of a similar magnitude order to  $\zeta_a$ . This verifies the motivation of data preprocessing by centralizing and whitening.

2.3.3. Generating New Individuals. To generate new individuals from the current generation, we use three genetic operators, selection, crossover, and mutation. The selection is based on the relative fitness of individuals. Specifically, the ratio of the fitness of an individual to the total fitness of the population determines how many times the individual will be selected as parent individuals. After evaluating all individuals in the current population, we select (S - 1) pairs of parent individuals from them, where *S* is the size of the GA population. Then the population of the next generation consists of the individual with the highest fitness in the current generation and the (S - 1) new individuals generated from these parent individuals.

The crossover operator is conducted under a given probability. If two parent individuals are not subjected to crossover, the one having higher fitness will be chosen into the next generation. Otherwise, two crossover points are randomly chosen, one of which is within the coefficient bits and the other is within the selection bits. These two points divide both parent individuals into three parts, and the second part is then exchanged between them to form two new individuals, one of which is randomly chosen as an individual in the next generation.

At last, each bit in the (S-1) new individuals is subjected to mutation from "0" to "1" or reversely under a specific probability. After applying all the three genetic operators, we have a new population for the next GA iteration.

2.3.4. Imposing Constraints on Searching. As discussed before, to further improve the search efficiency and the performance of the obtained projection basis vectors, some constraints are necessary for the search space. Thanks to the linear combination mechanism used by the proposed EDFE algorithm, it is very easy to force the GA to search in a constrained space. Our method is to construct vectors by linearly combining the basis of the constrained search space, instead of the original space. Take  $null(S_w)$ , the null space of  $S_w$ , as an example. Suppose that we want to constrain the GA to search in null( $S_w$ ). Let { $\alpha_i \in \mathbb{R}^n \mid i = 1, 2, ..., m$ } be the eigenvectors of  $S_w$  associated with zero eigenvalues. They form a basis of  $null(S_w)$ . After obtaining a vector v via linearly combining the above basis, we have to calculate the basis of the orthogonal complement space of V =span{v} in the constrained search space null( $S_w$ ), but not the original space  $R^n$  (referring to Section 2.1). For this purpose, we first calculate the isomorphic space of V in  $\mathbb{R}^m$ , denoted by  $\hat{V} = \text{span}\{P^T v\}$ , where  $P = [\alpha_1 \alpha_2 \cdots \alpha_m]$  is an isomorphic mapping. We then calculate a basis of the orthogonal complement space of  $\hat{V}$  in  $\mathbb{R}^m$ . Let  $\{\hat{\beta}_i \in \mathbb{R}^m \mid$  $i = 1, 2, \dots, m - 1$ } be the obtained basis. Finally, we map this basis back into null( $S_w$ ) through  $\{\beta_i = P\hat{\beta}_i \in \mathbb{R}^n \mid i = i\}$  $1, 2, \ldots, m-1$ .

The following theorem demonstrates that  $\{\beta_i \mid i = 1, 2, ..., m - 1\}$  comprise a basis of the orthogonal complement space of *V* in null(*S*<sub>*w*</sub>).

**Theorem 1.** Assume that  $A 
ightharpoonrightarrow R^n$  is an m-dimensional space, and  $P = [\alpha_1 \alpha_2 \cdots \alpha_m]$  is an identity orthogonal basis of A, where  $\alpha_i \in R^n$ , i = 1, 2, ..., m. For any  $v \in A$ , suppose that  $\hat{V} = span\{P^Tv\} \subset R^m$ . Let  $\{\hat{\beta}_i \in R^m \mid i = 1, 2, ..., m - 1\}$  be an identity orthogonal basis of the orthogonally complement space of  $\hat{V}$  in  $\mathbb{R}^m$ , then  $\{\beta_i = P\hat{\beta}_i \in \mathbb{R}^n \mid i = 1, 2, ..., m-1\}$  is a basis of the orthogonally complement space of  $V = \text{span}\{v\}$ in A.

Proof. See Appendix 6.

#### **3. Bagging EDFE**

The EDFE algorithm proposed above is very applicable to high-dimensional data because of its low space complexity. However, since it is based on the idea of subspace methods like LDA, it could suffer from the outlier and over-fitting problems when the training set is large. Moreover, when there are many training samples, the null( $S_w$ ) becomes small, resulting in poor discrimination performance in the space. Wang and Tang [33] proposed to solve this problem using two random sampling techniques, random subspace and bagging. To improve the performance of the EDFE algorithm on large scale datasets, we propose to incorporate the bagging technique into the EDFE algorithm and hence develop the bagging evolutionary discriminant feature extraction (BEDFE) algorithm.

Bagging (acronym for Bootstrap AGGregatING), proposed by Breiman [34], uses resampling to generate several random subsets (called random bootstrap replicates) from the whole training set. From each replicate, one classifier is constructed. The results by these classifiers are integrated using some fusion scheme to give the final result. Since these classifiers are trained from relatively small bootstrap replicates, the outlier and over-fitting problems for them are expected to be alleviated. In addition, the stability of the overall classifier system can be improved by integration of several (weak) classifiers.

Like Wang and Tang's method, we randomly choose some classes from all the classes in the training set. The training samples belonging to these classes compose a bootstrap replicate. Usually, the unchosen samples become useless in the learning process. Instead, we do not overlook these data, but rather use them for validation and calculate the classification accuracy term in the fitness function. Below are the primary steps of the BEDFE algorithm.

- (1) Preprocess the data using centralizing and whitening.
- (2) Randomly choose some classes, say  $\hat{L}$  classes, from all the *L* classes in the training set. The samples belonging to the  $\hat{L}$  classes compose a bootstrap replicate used for training, and those belonging to the other  $(L \hat{L})$  classes are used for validation. Totally, *K* replicates are created (different replicates could have different classes).
- (3) Run the EDFE algorithm on each replicate to learn a feature subspace. In all, *K* feature subspaces are obtained.
- (4) Classify each new sample using a classifier in the K feature subspaces, respectively. The resulting K results are combined by a fusion scheme to give the final result.

There are two key steps in the BEDFE algorithm: how to do validation and classification, and how to fuse the results from different replicates. In the following we present our solutions to these two problems.

3.1. Validation and Classification. As shown above, a training replicate is created from the chosen  $\hat{L}$  classes. Based on this training replicate, an individual in the EDFE population generates a candidate projection basis of feature subspace. All the samples in the training replicate are projected into this feature subspace. The generalization term in the fitness function is then calculated from these projections. To obtain the value of the classification accuracy term, we again randomly choose some samples from all the samples of each class in the  $(L - \hat{L})$  validation classes to form the validation set. We then project the remaining samples in these classes to the feature subspace and calculate the mean as the prototype for each validation class according to the projections. Finally, the chosen samples are classified based on these prototypes using a classifier. The classification rate is used as the value of the classification accuracy term in the fitness function.

After running the EDFE algorithm on all the replicates, we get K feature subspaces as well as one projection basis for each of them. For each feature subspace, all the training samples (including the samples in training replicates and validation classes) are projected into the feature subspace, and the means of all classes are calculated as the prototypes of them. To classify a new sample, we first classify it in each of the K feature subspaces based on the class prototypes in that space and then fuse the K results to give the final decision, which is introduced in the following part.

3.2. The Majority Voting Fusion Scheme. A number of fusion schemes [35, 36] have been proposed in literature of multiple classifiers and information fusion. In the present paper, we only focus on Majority Voting for its intuitiveness and simplicity. Let  $\{M_i^k \in \mathbb{R}^{l_k} \mid j = 1, 2, ..., L; k = 1, 2, ..., K\}$ be the prototype of class j in the kth feature subspace, whose dimensionality is  $l_k$ . Given a new sample (represented as a vector), we first preprocess it by centralization and whitening; that is, the mean of all the training samples is subtracted from it, and the resulting vector is projected into the whitened PCA space learned from the training samples. Denote by  $x_t$  the preprocessed sample. It is projected into each of the K feature subspaces, resulting in  $y_t^k$  in the kth feature subspace, and classified in these feature subspaces, respectively. Finally, the Majority Voting scheme is employed to fuse the classification results obtained in the K feature subspaces.

Majority Voting is one of the simplest and most popular classifier fusion schemes. Take the Nearest Mean Classifier (NMC) and the *k*th feature subspace as an example. The NMC assigns  $x_t$  to the class  $c^k \in \{1, 2, ..., L\}$  such that

$$\left\| y_{t}^{k} - M_{c^{k}}^{k} \right\| = \min_{j \in \{1, 2, \dots, L\}} \left\| y_{t}^{k} - M_{j}^{k} \right\|.$$
(25)

In other words, it votes for the class whose prototype is closest to  $y_t^k$ . After classifying  $x_t$  in all the *K* feature

TABLE 1: General information and settings of the used databases.

Database	Sub number	Size number	Image/Sub number	Train number	Validation number	Test number
ORL	40	$92 \times 112$	10	4(2)	1(3)	5
AR	120	80  imes 100	14	6(3)	1(4)	7

<sup>a</sup> From the first column to the last column: the name of the database, the number of subjects, the size of images, the number of images per subject, the number of training samples per subject, the number of validation samples per subject, and the number of test samples per subject.

<sup>b</sup>The numbers in parentheses are the numbers of samples per validation subject used by BEDFE to calculate the class prototypes and to evaluate the training performance.

subspaces, we get *K* results  $\{c^k | k = 1, 2, ..., K\}$ . Let Votes(i) be the number of votes obtained by class *i*, that is,

Votes(i) = #
$$\{c^k = i \mid k = 1, 2, ..., K\},$$
 (26)

where "#" denotes the cardinality of a set. The final class label of  $x_t$  is determined to be c if

$$Votes(c) = \max_{i \in \{1, 2, \dots, L\}} Votes(i).$$
(27)

#### 4. Face Recognition Experiments

Face recognition experiments have been performed on the ORL and AR face databases. Due to the high dimensionality of the data, conventional GA-based feature extraction methods like EP [18] and EDA [37] cannot be directly applied to these two databases unless reducing the dimensionality in advance. By contraries, the EDFE and BEDFE algorithms proposed in this paper can still work very well with them. As an application of the algorithms, we will use them to investigate the discriminative ability of the three subspaces, null( $S_w$ ), range( $S_w$ ), and range( $S_b$ ). We will experimentally demonstrate the necessity of carefully choosing the dimension of feature subspace. Finally, we will compare the proposed algorithms with some state-of-the-art methods in the literature, that is, Eigenfaces [9], Fisherfaces [10], Null-space LDA [16], EP [18], and EDA+Full-space LDA [32].

4.1. The Face Databases and Parameter Settings. The ORL database of faces [38] contains 400 face images of 40 distinct subjects. Each subject has 10 different images, which were taken at different times. These face images have variant lighting, facial expressions (open/closed eyes, smiling/not smiling) and facial details (glasses/no glasses). They also display slight pose changes. The size of each image is 92 imes112 pixels, with 256 gray levels per pixel. The AR face database [39] has much larger scale than the ORL database. It has over 4000 color images of 126 people (70 men and 56 women), which have different facial expressions, illumination conditions, and occlusions (wearing sun-glasses and scarf). In our experiments, we randomly chose some images of 120 subjects and discarded the samples of wearing sun-glasses and scarf. In the resulting dataset, there are 14 face images for each chosen subject, totally 1680 images. All these images were converted to gray scale images, and the face portion on them was manually cropped and resized to  $80 \times 100$  pixels. For both databases, all images were preprocessed by histogram equalization. Table 1 lists some



FIGURE 3: Some sample images in the (a) ORL and (b) AR face databases.

(b)

general information of the two databases, and Figure 3 shows some sample images of them.

In the GA algorithm, we set the probability of crossover to 0.8, the probability of mutation to 0.01, the size of population to 50, and the number of generations to 100. For the weight of the classification accuracy term in the fitness function, we considered the following choices for EDFE: {0.0, 0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, 0.9, 1.0}. After finding the weight which gives the best classification accuracy for a dataset, we adopted it in BEDFE on the dataset. Regarding the number of bagging replicates in BEDFE, we conducted experiments for four cases, that is, using 3, 5, 7, and 9 replicates, and then chose the best one among them. The results will be presented in the following parts.

To create an evaluation face image set, all the sample images in each database were randomly divided into three parts: the training set, the validation set, and the test set. In the experiments on the ORL database, four images were randomly chosen from the samples of each subject for training, one image for validation and the remaining five images for test. In the experiments on the AR database, six images were randomly chosen for training from the samples of each subject, one image for validation, and the rest seven 8

TABLE 2: Recognition accuracy of EDFE in different subspaces.

Database	$\operatorname{Null}(S_w)$	Range( $S_w$ )	Range( $S_b$ )
ORL	90.3%	78.1%	79.5%
AR	95.33%	80.95%	81.67%

TABLE 3: Recognition accuracy of BEDFE in different subspaces.

Database	$\operatorname{Null}(S_w)$	$\operatorname{Range}(S_w)$	Range( $S_b$ )
ORL	91.3%	80.02%	81.14%
AR	96.86%	83.1%	83.38%

TABLE 4: The mean and standard deviation of recognition accuracy (%) of the proposed EDFE and BEDFE methods and some other state-of-the-art methods on the ORL and AR face databases.

Method	ORL face database	AR face database
Eigenfaces	90.15 ± 3.2	$82.68 \pm 0.9$
Fisherfaces	$91.6 \pm 1.51$	$96.99 \pm 0.7$
Null-space LDA	$89.75 \pm 1.21$	$96.71 \pm 0.59$
EP	$80.31\pm3.5$	N/A
EDA+Full-space LDA	$92.5 \pm 2.1$	$97.02\pm0.8$
$EDFE+Full-space(S_w)$	$93 \pm 1.8$	$97.9\pm0.7$
BEDFE+Full-space( $S_w$ )	$95.5 \pm 1.12$	$98.55\pm0.46$

images for test. For the methods Eigenfaces, Fisherfaces, Null-space LDA, and EP, no validation set is required. Thus we combined the training images and validation images to form the training set for them. The case was a little bit different for the experiments with BEDFE, where the division of samples is on the class level (each subject is a class). Specifically, we first randomly chose five (seven) images from each subject to compose the test set of the ORL (AR) database. Among the remaining images, a subset of classes was randomly chosen. The samples belonging to these classes were used for training whereas those belonging to the other classes composed the validation set. From each class in the validation set, some samples were randomly chosen to calculate the prototype of the class, and the remaining ones were used for evaluation. On the ORL database, two images of each validation class were randomly chosen for class prototype calculation, and the other three images of the class were used to evaluate the training performance. On the AR database, three images were randomly chosen from each validation class for computing the class prototype, and the other four images of the class were used for training performance evaluation. The last three columns in Table 1 summarize these settings.

Totally, we created 10 evaluation sets from each of the two databases and ran algorithms over them one by one. We will use the mean and standard deviation of recognition accuracy over the 10 evaluation sets to evaluate the performance of different methods. When applying EP to the ORL databases, the dimensionality of the data should be reduced in advance due to the high space complexity of EP. We reduced the data to a dimension of the number of training samples minus one using WPCA (note that the role of WPCA here is different from that in the proposed EDFE algorithm). To evaluate the performance of Eigenfaces on the databases, we tested all possible dimensions of PCA-transformed subspace (between 1 and N-1) and found out the one with the best classification accuracy. As for Fisherfaces, we set the dimension of PCA-transformed subspace to the rank of  $S_t$  and tried all possible dimensions of LDA-transformed subspace (between 1 and L-1).

4.2. Investigation on Different Subspaces. Three subspaces, null( $S_w$ ), range( $S_w$ ), and range( $S_b$ ), are thought to contain rich discriminative information within data [16, 17]. As mentioned above, the algorithms proposed in this paper provide a method to constrain the search in a specific subspace. Hence, we can restrict the algorithms to search for a solution within that subspace. Here we report our experimental results in investigating the above three subspaces using the EDFE and BEDFE algorithms on the ORL and AR databases.

Table 2 shows the average recognition accuracy of EDFE in the three different subspaces on the ten evaluation sets of ORL and AR databases. The presented classification accuracies are the best ones among those obtained using different weights. On all the ten evaluation sets of both ORL and AR databases,  $null(S_w)$  gives the best results, which are significant better than the other two subspaces. On the other hand, there is no big difference between the performance of  $range(S_w)$  and  $range(S_b)$ . This is not surprising because in the null space of  $S_w$ , if exists, samples in the same class will be condensed to one point. Then if a projection basis in it can be found to make the samples of different classes separable from each other, the classification performance on these samples will be surely the best. However, for new samples unseen in the training set, the classification accuracy on them depends on the accuracy of the estimation of  $S_w$ . Another problem with  $null(S_w)$  is that its dimensionality is bounded by the minimum of the dimensionality of the data and the difference between the number of samples and the number of classes. Consequently, as the number of training samples increases, this null space could become too small to contain sufficient discriminant information. In this case, we propose to incorporate the bagging technique to the EDFE algorithm to enhance its performance. The results of BEDFE are given in Table 3, from which similar conclusion can be drawn.

4.3. Investigation on Dimensionality of Feature Subspaces. In order to show the importance of carefully choosing the dimensionality of feature subspaces, we calculated the average recognition accuracy of Eigenfaces and Fisherfaces on the ten evaluation sets taken from the ORL and AR face databases when different numbers of features were chosen for the feature subspaces. The possible dimension of the feature subspace obtained by Eigenfaces on the ORL evaluation sets is between 1 and 199 (i.e., the number of samples minus one), whereas that on the AR evaluation sets is between 1 and 839. As for Fisherfaces, we set the dimension of PCA-reduced feature subspace to 720 (i.e., the number of samples minus the number of classes) and tested all the possible dimension

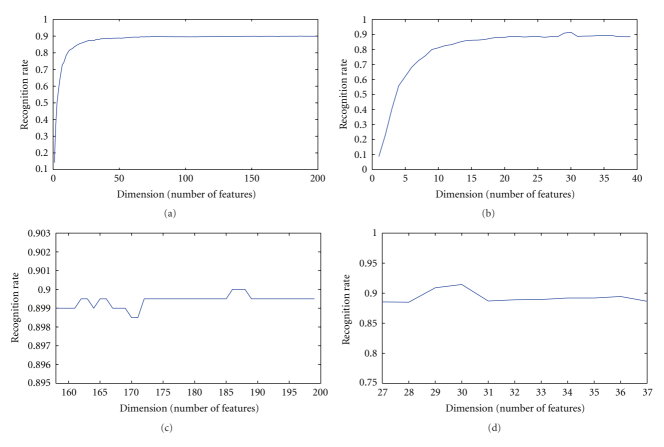


FIGURE 4: The curves of the average recognition accuracy of (a) Eigenfaces and (b) Fisherfaces on the ORL face database versus the number of features or the dimension of feature subspaces. (c) and (d) are the corresponding enlarged last parts of the curves.

of LDA-reduced feature subspace from 1 to 119 (i.e., the number of classes minus one). According to the experimental results, the overall trend of recognition accuracy is increasing as the number of features (i.e., the dimension of feature subspace) increases. However, the best accuracy is often obtained not at the largest possible dimension (i.e., the number of samples minus one in case of Eigenfaces and the number of classes minus one in case of Fisherfaces). Figures 4 and 5 show the curves of the average recognition accuracy of Eigenfaces and Fisherfaces on ORL and AR face databases versus the dimension of feature subspaces (to clearly show that the best accuracy is achieved not necessarily at the largest possible dimension, we also display the last part of the curves in an enlarged view). From these results, we can see that the dimension at which the best recognition accuracy is achieved varies with respect to the datasets. Therefore, using a systematic method like the ones proposed in this paper to automatically determine the dimension of feature subspaces is very helpful to a subspace-based recognition system.

4.4. Performance Comparison. Finally, we compared the proposed algorithms with some state-of-the-art methods in literature, including Eigenfaces [9], Fisherfaces [10], Null-space LDA [16], EP [18], and EDA+Full-space LDA [32]. Considering that both null( $S_w$ ) and range( $S_w$ ) have useful

discriminative information, we ran our proposed EDFE and BEDFE methods in both  $null(S_w)$  and  $range(S_w)$  and then employed the same fusion method used by [32] to fuse the results obtained in these two subspaces. We called them EDFE+Full-space( $S_w$ ) and BEDFE+Full-space( $S_w$ ). We implemented these methods by using Matlab and evaluated their performance on the ten evaluation sets of ORL and AR face databases. But as for the EP method, it is too computationally complex to be applicable (N/A) on the AR face database (in Matlab an error of 'out of memory' will be reported to the EP method). We calculated the mean and standard deviation of the recognition rates for all the methods. The results are listed in Table 4 (the results of Eigenfaces and Fisherfaces are according to the best results obtained in the last subsection).

It can be seen from the results that the proposed EDFE and BEDFE methods overwhelm their counterpart methods in the average recognition accuracy. Moreover, by using the bagging technique, the BEDFE method performs much more stable than EDFE, and it has the smallest deviation of recognition accuracy among all the methods. A possible reason for such improvement on the stability is that by using smaller training sets and multiple feature subspace fusion, the outlier and over-fitting problems of conventional machine learning and pattern recognition

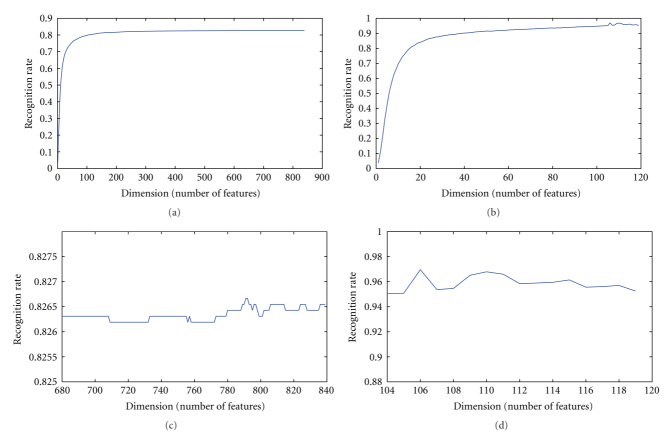


FIGURE 5: The curves of the average recognition accuracy of (a) Eigenfaces and (b) Fisherfaces on the AR face database versus the number of features or the dimension of feature subspaces. (c) and (d) are the corresponding enlarged last parts of the curves.

systems could be alleviated. Moreover, the improvement on recognition accuracy made by the proposed EDFE and BEDFE compared with the other methods could be due to their better generalization ability. In Eigenfaces, Fisherfaces, Null-space LDA, and EDA+Full-space LDA, the projection basis used for dimension reduction is directly calculated from certain covariance or scatter matrix of the training data. Instead, the methods proposed in this paper begin the search of optimal projection basis from these directly calculated ones and iteratively approach the best one via the linear combination of them. The linear combination not only ensures that the resulting projection basis still lies in the feature subspace but also enhances the generalization ability of the obtained projection basis by adjusting them according to the recognition accuracy on some validation data.

## 5. Discussion

In the proposed EDFE and BEDFE algorithms, we take the classification accuracy term as a part of the fitness function of the GA. It is then naturally optimized as the GA population evolves. Unlike existing evolutionary computation-based feature extraction methods like EP [18], we define this term on a randomly chosen validation sample set, but not the training set. Since the validation set's role is to simulate

new test samples, the performance of the resulting feature subspace is supposed to be more reliable. We also set up a Fisher criterion-like term as another part of the GA's fitness function and optimize it in an iterative way, avoiding the matrix inverse operation required by the conventional LDA method. As a result, the proposed algorithms could alleviate the small sample size (SSS) problem of LDA.

Current PCA- and LDA-based subspace methods such as Eigenfaces and Fisherfaces require setting the dimensionality for the feature subspace in advance. They fail to provide systematic way to automatically determine the dimensionality from the classification viewpoint. Since the optimal dimensionality of feature subspace in terms of recognition rates will vary across datasets, it is desired to select automatically the optimal dimensionality for specific datasets, instead of using a predefined one. The proposed EDFE and BEDFE algorithms provide such a way by employing the stochastic optimization scheme of GA.

Some other GA-based feature selection/extraction methods have been also proposed in literature. Although these GA-based feature selection methods, such as EDA+Fullspace LDA [32], GA-PCA and GA-Fisher [26], have the advantage in lower space and time requirement, they are limited in the ability of searching discriminative features. On the other hand, those GA-based feature extraction methods have high space complexity and are thus not applicable to high dimensional and large scale datasets. For example, in EP [18], an individual has  $(5n^2 - 4n)$  bits. In the recently proposed EDA algorithm [37], the individual has to encode  $(n \times m)$  weights, which are between -0.5 and 0.5 (here, n is the dimension of the original data space, and m is the dimension of the feature subspace). On the contrary, the individual in the proposed algorithms has only (12n - 1)bits (note that in face recognition applications, *m* is usually much larger than 12 and a number of bits have to be used to represent a decimal weight used by EDA). As a result, the space complexity is significantly reduced in our proposed methods. After incorporating the bagging technique with the proposed EDFE algorithm, it becomes more stable by eliminating possible outliers and fusing different feature subspaces, and hence more suitable for high-dimensional and large scale datasets.

Another problem with existing GA-based feature extraction methods lies in their blind search strategy. Using the linear combination and orthogonal complement techniques, the EDFE successfully provides a way to impose constraints on the search space of GA. This also enables EDFE to effectively make use of the heuristic information of the discriminative feature subspace to improve its search efficiency and classification performance. In addition, the proposed algorithms make it possible to investigate the discriminative ability of different feature subspaces.

#### 6. Conclusions

In this paper, we proposed an evolutionary approach to extracting discriminative features, namely, evolutionary discriminant feature extraction (EDFE) as well as its bagging version (BEDFE). The basic idea underlying the EDFE algorithm is to use the genetic algorithm (GA) to search for an optimal discriminative projection basis in a constrained subspace with the goal of making the data in different clusters much easier to be separated. The primary contribution of this paper includes (1) reducing the space complexity of GA-based feature extraction algorithms; (2) enhancing the search efficiency, stability, as well as recognition accuracy; (3) providing an effective way to investigate different feature subspaces. Experiments on the ORL and AR databases have been performed to validate the proposed methods.

There are still some issues worthy further study on the proposed approach. Firstly, it is a supervised linear feature extraction method. Therefore, how to extend it to nonlinear cases deserves further study. Secondly, the latest progress in the research on GA, for example, how to set up an initial population and how to choose proper GA parameters, could give us some useful hints on further improving the proposed methods. Thirdly, more promising results could be obtained by exploring other criteria to evaluate the feature subspaces and incorporating them into the evolutionary approach, for instance, those of recently proposed manifold learning algorithms [40, 41]. Finally, it could be very interesting to investigate the discriminability of other subspaces using the proposed EDFE and BEDFE algorithms.

#### Appendix

## **Proof of Theorem 1**

*Proof.* First it can be easily proved that for all  $i \in \{1, 2, ..., m-1\}$ ,  $\beta_i \in A$ . Since  $\beta_i = P\hat{\beta}_i = \sum_{j=1}^m \beta_{ij}\alpha_j$ ,  $\beta_i$  can be represented by a basis of *A*. Thus  $\beta_i \in A$ .

Secondly, let us prove  $A = U \oplus V$ , where  $U = \text{span}\{\beta_1, \beta_2, \dots, \beta_{m-1}\}$ . This is to prove that  $\{\beta_1, \beta_2, \dots, \beta_{m-1}, \nu\}$  is a linear independent bundle. Since  $\beta_i = P\hat{\beta}_i$  and  $P^TP = E_m$ , which is an *m*-dimensional identity matrix, we have  $\beta_i^T\beta_j = \hat{\beta}_i^T\hat{\beta}_j$ . However,  $\{\hat{\beta}_i \mid i = 1, 2, \dots, m-1\}$  is an identity orthogonal basis. Thus,  $\beta_1, \beta_2, \dots, \beta_{m-1}$  are orthogonal to each other.

Furthermore, for all  $i \in \{1, 2, ..., m - 1\}$ ,  $\beta_i^T v = (P\hat{\beta}_i)^T v = \hat{\beta}_i^T P^T v$ . Because  $\{\hat{\beta}_i \mid i = 1, 2, ..., m - 1\}$  is an identity orthogonal basis of the orthogonal complement space of  $\hat{V} = \text{span}\{P^T v\}$  in  $R^m$ ,  $\hat{\beta}_i^T(P^T v)$  should be zero. Therefore,  $\beta_i^T v = 0$ , that is,  $\beta_i$  is also orthogonal to v.

To sum up, we get that  $\{\beta_1, \beta_2, \dots, \beta_{m-1}, \nu\}$  is a linear independent bundle containing *m* orthogonal vectors in the *m* dimensional space *A*. Thus  $A = U \oplus V$ .

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