Robust Face Recognition via Sparse Representation

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Abstract—We consider the problem of automatically recognizing human faces from frontal views with varying expression and illumination, as well as occlusion and disguise. We cast the recognition problem as one of classifying among multiple linear regression models and argue that new theory from sparse signal representation offers the key to addressing this problem. Based on a sparse representation computed by $\ell^1$-minimization, we propose a general classification algorithm for (image-based) object recognition. This new framework provides new insights into two crucial issues in face recognition: feature extraction and robustness to occlusion. For feature extraction, we show that if sparsity in the recognition problem is properly harnessed, the choice of features is no longer critical. What is critical, however, is whether the number of features is sufficiently large and whether the sparse representation is correctly computed. Unconventional features such as downsampled images and random projections perform just as well as conventional features such as Eigenfaces and Laplacianfaces, as long as the dimension of the feature space surpasses certain threshold, predicted by the theory of sparse representation. This framework can handle errors due to occlusion and corruption uniformly by exploiting the fact that these errors are often sparse with respect to the standard (pixel) basis. The theory of sparse representation helps predict how much occlusion the recognition algorithm can handle and how to choose the training images to maximize robustness to occlusion. We conduct extensive experiments on publicly available databases to verify the efficacy of the proposed algorithm and corroborate the above claims.

Index Terms—Face recognition, feature extraction, occlusion and corruption, sparse representation, compressed sensing, $\ell^1$-minimization, validation and outlier rejection.

1 INTRODUCTION

Parsimony has a rich history as a guiding principle for inference. One of its most celebrated instantiations, the principle of minimum description length in model selection [1], [2], stipulates that within a hierarchy of model classes, the model that yields the most compact representation should be preferred for decision-making tasks such as classification. A related, but simpler, measure of parsimony in high-dimensional data processing seeks models that depend on only a few of the observations, selecting a small subset of features for classification or visualization (e.g., Sparse PCA [3], [4] among others). Such sparse feature selection methods are, in a sense, dual to the support vector machine (SVM) approach in [5] and [6], which instead selects a small subset of relevant training examples to characterize the decision boundary between classes. While these works comprise only a small fraction of the literature on parsimony for inference, they do serve to illustrate a common theme: all of them use parsimony as a principle for choosing a limited subset of features or models from the training data, rather than directly using the data for representing or classifying an input (test) signal.

The role of parsimony in human perception has also been strongly supported by studies of human vision. Investigators have recently revealed that in both low-level and midlevel human vision [7], [8], many neurons in the visual pathway are selective for a variety of specific stimuli, such as color, texture, orientation, scale, and even view-tuned object images. Considering these neurons to form an overcomplete dictionary of base signal elements at each visual stage, the firing of the neurons with respect to a given input image is typically highly sparse.

In the statistical signal processing community, the algorithmic problem of computing sparse linear representations with respect to an overcomplete dictionary of base elements or signal atoms has seen a recent surge of interest [9], [10], [11], [12]. Much of this excitement centers around the discovery that whenever the optimal representation is sufficiently sparse, it can be efficiently computed by convex optimization [9], even though this problem can be extremely difficult in the general case [13]. The resulting optimization problem, similar to the Lasso in statistics...
Our use of sparsity for classification differs significantly from the various parsimony principles discussed above. Instead of using sparsity to identify a relevant model or relevant features that can later be used for classifying all test samples, it uses the sparse representation of each individual test sample directly for classification, adaptively selecting the training samples that give the most compact representation. The proposed classifier can be considered a generalization of popular classifiers such as nearest neighbor (NN) [18] and nearest subspace (NS) [19] (i.e., minimum distance to the subspace spanned all training samples from each object class). NN classifies the test sample based on the best representation in terms of a single training sample, whereas NS classifies based on the best linear representation in terms of all the training samples in each class. The nearest feature line (NFL) algorithm [20] strikes a balance between these two extremes, classifying based on the best affine representation in terms of a pair of training samples. Our method strikes a similar balance but considers all possible supports (within each class or across multiple classes) and adaptively chooses the minimal number of training samples needed to represent each test sample.

In this paper, we exploit the discriminative nature of sparse representation to perform classification. Instead of using the generic dictionaries discussed above, we represent the test sample in an overcomplete dictionary whose base elements are the training samples themselves. If sufficient training samples are available from each class, it will be possible to represent the test samples as a linear combination of just those training samples from the same class. This representation is naturally sparse, involving only a small fraction of the overall training database. We argue that in many problems of interest, it is actually the sparsest linear representation of the test sample in terms of this dictionary and can be recovered efficiently via \( \ell_1 \)-minimization. Seeking the sparsest representation therefore automatically discriminates between the various classes present in the training set. Fig. 1 illustrates this simple idea using face recognition as an example. Sparse representation also provides a simple and surprisingly effective means of rejecting invalid test samples not arising from any class in the training database: these samples’ sparsest representations tend to involve many dictionary elements, spanning multiple classes.

2. In contrast, methods such as that in [16] and [17] that utilize only a single training sample per class face a more difficult problem and generally incorporate more explicit prior knowledge about the types of variation that could occur in the test sample.

3. The relationship between our method and NN, NS, and NFL is explored more thoroughly in the supplementary appendix, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79.
high-dimensional test image into lower dimensional feature spaces: examples include Eigenfaces [23], Fisherfaces [24], Laplacianfaces [25], and a host of variants [26], [27]. With so many proposed features and so little consensus about which are better or worse, practitioners lack guidelines to decide which features to use. However, within our proposed framework, the theory of compressed sensing implies that the precise choice of feature space is no longer critical: Even random features contain enough information to recover the sparse representation and hence correctly classify any test image. What is critical is that the dimension of the feature space is sufficiently large and that the sparse representation is correctly computed.

Robustness to occlusion. Occlusion poses a significant obstacle to robust real-world face recognition [16], [28], [29]. This difficulty is mainly due to the unpredictable nature of the error incurred by occlusion; it may affect any part of the image and may be arbitrarily large in magnitude. Nevertheless, this error typically corrupts only a fraction of the image pixels and is therefore sparse in the standard basis given by individual pixels. When the error has such a sparse representation, it can be handled uniformly within our framework: the basis in which the error is sparse can be treated as a special class of training samples. The subsequent sparse representation of an occluded test image with respect to this expanded dictionary (training images plus error basis) naturally separates the component of the test image arising due to occlusion from the component arising from the identity of the test subject (see Fig. 1 for an example). In this context, the theory of sparse representation and compressed sensing characterizes when such source-and-error separation can take place and therefore how much occlusion the resulting recognition algorithm can tolerate.

Organization of this paper. In Section 2, we introduce a basic general framework for classification using sparse representation, applicable to a wide variety of problems in image-based object recognition. We will discuss why the sparse representation can take place and therefore how much error separation can take place and how much occlusion the resulting recognition algorithm can tolerate.

Classification based on sparse representation

A basic problem in object recognition is to use labeled training samples from $k$ distinct object classes to correctly determine the class to which a new test sample belongs. We arrange the given $n_i$ training samples from the $i$th class as columns of a matrix $A_i = [v_{i,1}, v_{i,2}, \ldots, v_{i,n_i}] \in \mathbb{R}^{m \times n_i}$. In the context of face recognition, we will identify a $w \times h$ grayscale image with the vector $v \in \mathbb{R}^m$ ($m = wh$) given by stacking its columns; the columns of $A_i$ are then the training face images of the $i$th subject.

2.1 Test Sample as a Sparse Linear Combination of Training Samples

An immense variety of statistical, generative, or discriminative models have been proposed for exploiting the structure of the $A_i$ for recognition. One particularly simple and effective approach models the samples from a single class as lying on a linear subspace. Subspace models are flexible enough to capture much of the variation in real data sets and are especially well motivated in the context of face recognition, where it has been observed that the images of faces under varying lighting and expression lie on a special low-dimensional subspace [24], [30], often called a face subspace. Although the proposed framework and algorithm can also apply to multimodal or nonlinear distributions (see the supplementary appendix for more detail, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79), for ease of presentation, we shall first assume that the training samples from a single class do lie on a subspace. This is the only prior knowledge about the training samples we will be using in our solution.

Given sufficient training samples of the $i$th object class, $A_i = [v_{i,1}, v_{i,2}, \ldots, v_{i,n_i}] \in \mathbb{R}^{m \times n_i}$, any new (test) sample $y \in \mathbb{R}^m$ from the same class will approximately lie in the linear span of the training samples$^5$ associated with object $i$:

$$y = \alpha_{i, 1} v_{i, 1} + \alpha_{i, 2} v_{i, 2} + \cdots + \alpha_{i, n_i} v_{i, n_i},$$

(1)

for some scalars, $\alpha_{i,j} \in \mathbb{R}$, $j = 1, 2, \ldots, n_i$.

Since the membership $i$ of the test sample is initially unknown, we define a new matrix $A$ for the entire training set as the concatenation of the $n$ training samples of all $k$ object classes:

$$A = [A_1, A_2, \ldots, A_k] = [v_{1,1}, v_{1,2}, \ldots, v_{k,n_k}].$$

(2)

Then, the linear representation of $y$ can be rewritten in terms of all training samples as

$$y = Ax_0 \in \mathbb{R}^m,$$

(3)

where $x_0 = [0, \ldots, 0, \alpha_{1,1}, \alpha_{1,2}, \ldots, \alpha_{1,n_1}, 0, \ldots, 0]^T \in \mathbb{R}^n$ is a coefficient vector whose entries are zero except those associated with the $i$th class.

4. In face recognition, we actually do not need to know whether the linear structure is due to varying illumination or expression, since we do not rely on domain-specific knowledge such as an illumination model [31] to eliminate the variability in the training and testing images.

5. One may refer to [32] for how to choose the training images to ensure this property for face recognition. Here, we assume that such a training set is given.
As the entries of the vector \( x_0 \) encode the identity of the test sample \( y \), it is tempting to attempt to obtain it by solving the linear system of equations \( y = Ax \). Notice, though, that using the entire training set to solve for \( x \) represents a significant departure from one sample or one class at a time methods such as NN and NS. We will later argue that one can obtain a more discriminative classifier from such a global representation. We will demonstrate its superiority over these local methods (NN or NS) both for identifying objects represented in the training set and for rejecting outlying samples that do not arise from any of the classes present in the training set. These advantages can come without an increase in the order of growth of the computation: As we will see, the complexity remains linear in the size of training set.

Obviously, if \( m > n \), the system of equations \( y = Ax \) is overdetermined, and the correct \( x_0 \) can usually be found as its unique solution. We will see in Section 3, however, that in robust face recognition, the system \( y = Ax \) is typically underdetermined, and so, its solution is not unique. Conventional, this difficulty is resolved by choosing the minimum \( \ell^2 \)-norm solution:

\[
(\ell^2) : \quad \hat{x}_2 = \arg \min \|x\|_2 \quad \text{subject to} \quad Ax = y. \tag{4}
\]

While this optimization problem can be easily solved (via the pseudoinverse of \( A \)), the solution \( \hat{x}_2 \) is not especially informative for recognizing the test sample \( y \). As shown in Example 1, \( \hat{x}_2 \) is generally dense, with large nonzero entries corresponding to training samples from many different classes. To resolve this difficulty, we instead exploit the following simple observation: A valid test sample \( y \) can be sufficiently represented using only the training samples from the same class. This representation is naturally sparse if the number of object classes \( k \) is reasonably large. For instance, if \( k = 20 \), only 5 percent of the entries of the desired \( x_0 \) should be nonzero. The more sparse the recovered \( x_0 \), the easier will it be to accurately determine the identity of the test sample \( y \).

This motivates us to seek the sparsest solution to \( y = Ax \), solving the following optimization problem:

\[
(\ell^0) : \quad \hat{x}_0 = \arg \min \|x\|_0 \quad \text{subject to} \quad Ax = y, \tag{5}
\]

where \( \| \cdot \|_0 \) denotes the \( \ell^0 \)-norm, which counts the number of nonzero entries in a vector. In fact, if the columns of \( A \) are in general position, then whenever \( y = Ax \) for some \( x \) with less than \( m/2 \) nonzeros, \( x \) is the unique sparsest solution: \( \hat{x}_0 = x \) [33]. However, the problem of finding the sparsest solution of an underdetermined system of linear equations is NP-hard and difficult even to approximate [13]; that is, in the general case, no known procedure for finding the sparsest solution is significantly more efficient than exhaustively testing all subsets of the entries for \( x \).

### 2.2 Sparse Solution via \( \ell^1 \)-Minimization

Recent development in the emerging theory of sparse representation and compressed sensing [9], [10], [11] reveals that if the solution \( x_0 \) sought is sparse enough, the solution of the \( \ell^0 \)-minimization problem (5) is equal to the solution to the following \( \ell^1 \)-minimization problem:

\[
(\ell^1) : \quad \hat{x}_1 = \arg \min \|x\|_1 \quad \text{subject to} \quad Ax = y. \tag{6}
\]

This problem can be solved in polynomial time by standard linear programming methods [34]. Even more efficient methods are available when the solution is known to be very sparse. For example, homotopy algorithms recover solutions with \( t \) nonzeros in \( O(t^4 + n) \) time, linear in the size of the training set [35].

#### 2.2.1 Geometric Interpretation

Fig. 2 gives a geometric interpretation (essentially due to [36]) of why minimizing the \( \ell^1 \)-norm correctly recovers sufficiently sparse solutions. Let \( P_0 \) denote the \( \ell^1 \)-ball (or crosspolytope) of radius \( \alpha \):

\[
P_0 = \{ x : \|x\|_1 \leq \alpha \} \subset \mathbb{R}^n. \tag{7}
\]

In Fig. 2, the unit \( \ell^1 \)-ball \( P_0 \) is mapped to the polytope \( P_0(A(P_0)) \subset \mathbb{R}^m \), consisting of all \( y \) that satisfy \( y = Ax \) for some \( x \) whose \( \ell^1 \)-norm is \( \leq 1 \).

The geometric relationship between \( P_0 \) and the polytope \( A(P_0) \) is invariant to scaling. That is, if we scale \( P_0 \), its image under multiplication by \( A \) is also scaled by the same amount. Geometrically, finding the minimum \( \ell^1 \)-norm solution \( \hat{x}_1 \) to (6) is equivalent to expanding the \( \ell^1 \)-ball \( P_0 \) until the polytope \( A(P_0) \) first touches \( y \). The value of \( \alpha \) at which this occurs is exactly \( \|\hat{x}_1\|_1 \).

Now, suppose that \( y = Ax_0 \) for some sparse \( x_0 \). We wish to know when solving (6) correctly recovers \( x_0 \). This question is easily resolved from the geometry of that in Fig. 2: Since \( \hat{x}_1 \) is found by expanding both \( P_0 \) and \( A(P_0) \) until a point of \( A(P_0) \) touches \( y \), the \( \ell^0 \)-minimizer \( \hat{x}_1 \) must generate a point \( A\hat{x}_1 \) on the boundary of \( P \).

Thus, \( \hat{x}_1 = x_0 \) if and only if the point \( A(x_0) \leq \alpha \) lies on the boundary of the polytope \( P \). For the example shown in Fig. 2, it is easy to see that the \( \ell^1 \)-minimization recovers all \( x_0 \) with only one nonzero entry. This equivalence holds because all of the vertices of \( P_1 \) map to points on the boundary of \( P \).

In general, if \( A \) maps all \( t \)-dimensional facets of \( P_1 \) to facets of \( P \), the polytope \( P \) is referred to as (centrally) \( t \)-neighborly [36]. From the above, we see that the \( \ell^1 \)-minimization (6) correctly recovers all \( x_0 \) with \( \leq t + 1 \) nonzeros if and only if \( P \) is \( t \)-neighborly, in which case, it is

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6. Furthermore, even in the overdetermined case, such a linear equation may not be perfectly satisfied in the presence of data noise (see Section 2.2.2).

7. This intuition holds only when the size of the database is fixed. For example, if we are allowed to append additional irrelevant columns to \( A \), we can make the solution \( x_0 \) have a smaller fraction of nonzeros, but this does not make \( x_0 \) more informative for recognition.
equivalent to the $\ell^0$-minimization (5). This condition is surprisingly common: even polytopes $P$ given by random matrices (e.g., uniform, Gaussian, and partial Fourier) are highly neighborly [15], allowing correct recovery of sparse $x_0$ by $\ell^0$-minimization.

Unfortunately, there is no known algorithm for efficiently verifying the neighborliness of a given polytope $P$. The best known algorithm is combinatorial, and therefore, only practical when the dimension $m$ is moderate [37]. When $m$ is large, it is known that with overwhelming probability, the neighborliness of a randomly chosen polytope $P$ is loosely bounded between

$$c \cdot m < t < \frac{((m+1)/3)}{t},$$

for some small constant $c > 0$ (see [9] and [36]). Loosely speaking, as long as the number of nonzero entries of $x_0$ is a small fraction of the dimension $m$, $\ell^1$-minimization will recover $x_0$.

### 2.2.2 Dealing with Small Dense Noise

So far, we have assumed that (3) holds exactly. Since real data are noisy, it may not be possible to express the test sample exactly as a sparse superposition of the training samples. The model (3) can be modified to explicitly account for small possibly dense noise by writing

$$y = Ax_0 + z,$$

where $z \in \mathbb{R}^m$ is a noise term with bounded energy $\|z\|_2 < \varepsilon$. The sparse solution $x_0$ can still be approximately recovered by solving the following stable $\ell^1$-minimization problem:

$$\begin{align*}
(\ell^1) : \quad \hat{x}_1 &= \arg \min \|x\|_1 \quad \text{subject to} \quad \|Ax - y\|_2 \leq \varepsilon. \quad (10)
\end{align*}$$

This convex optimization problem can be efficiently solved via second-order cone programming [34] (see Section 4 for our algorithm of choice). The solution of $(\ell^1)$ is guaranteed to approximately recovery sparse solutions in ensembles of random matrices $A$ [38]: There are constants $\rho$ and $\zeta$ such that with overwhelming probability, if $\|x_0\|_0 < \rho m$ and $\|z\|_2 \leq \varepsilon$, then the computed $\hat{x}_1$ satisfies

$$\|\hat{x}_1 - x_0\|_2 \leq \zeta \varepsilon.$$  

8. Thus, neighborliness gives a necessary and sufficient condition for sparse recovery. The restricted isometry properties often used in analyzing the performance of $\ell^1$-minimization in random matrix ensembles (e.g., [15]) give sufficient, but not necessary, conditions.

### 2.3 Classification Based on Sparse Representation

Given a new test sample $y$ from one of the classes in the training set, we first compute its sparse representation $\hat{x}_1$ via (6) or (10). Ideally, the nonzero entries in the estimate $\hat{x}_1$ will all be associated with the columns of $A$ from a single object class $i$, and we can easily assign the test sample $y$ to that class. However, noise and modeling error may lead to small nonzero entries associated with multiple object classes (see Fig. 3). Based on the global sparse representation, one can design many possible classifiers to resolve this. For instance, we can simply assign $y$ to the object class with the single largest entry in $\hat{x}_1$. However, such heuristics do not harness the subspace structure associated with images in face recognition. To better harness such linear structure, we instead classify $y$ based on how well the coefficients associated with all training samples of each object reproduce $y$.

For each class $i$, let $\delta_i : \mathbb{R}^n \rightarrow \mathbb{R}^n$ be the characteristic function that selects the coefficients associated with the $i$th class. For $x \in \mathbb{R}^n$, $\delta_i(x) \in \mathbb{R}^n$ is a new vector whose only nonzero entries are the entries in $x$ that are associated with class $i$. Using only the coefficients associated with the $i$th class, one can approximate the given test sample $y$ as $\tilde{y}_i = A \delta_i(\hat{x}_1)$. We then classify $y$ based on these approximations by assigning it to the object class that minimizes the residual between $y$ and $\tilde{y}_i$:

$$\min_i r_i(y) = \|y - A \delta_i(\hat{x}_1)\|_2.$$  

Algorithm 1 below summarizes the complete recognition procedure. Our implementation minimizes the $\ell^1$-norm via a primal-dual algorithm for linear programming based on [39] and [40].

Algorithm 1: Sparse Representation-based Classification (SRC)

1: **Input:** a matrix of training samples $A = [A_1, A_2, \ldots, A_k] \in \mathbb{R}^{m \times n}$ for $k$ classes, a test sample $y \in \mathbb{R}^m$, (and an optional error tolerance $\varepsilon > 0$)
2: Normalize the columns of $A$ to have unit $\ell^2$-norm.
3: Solve the $\ell^1$-minimization problem:

$$\hat{x}_1 = \arg \min_{x} \|x\|_1 \quad \text{subject to} \quad Ax = y.$$  

(Or alternatively, solve

$$\hat{x}_1 = \arg \min_{x} \|x\|_1 \quad \text{subject to} \quad Ax = y \leq \varepsilon.)$$

4: Compute the residuals $r_i(y) = \|y - A \delta_i(\hat{x}_1)\|_2$ for $i = 1, \ldots, k$.
5: **Output:** identity($y$) = arg min$_i$ $r_i(y)$. 

![Fig. 3. A valid test image. (a) Recognition with $12 \times 10$ downscaled images as features. The test image $y$ belongs to subject 1. The values of the sparse coefficients recovered from Algorithm 1 are plotted on the right together with the two training examples that correspond to the two largest sparse coefficients. (b) The residuals $r_i(y)$ of a test image of subject 1 with respect to the projected sparse coefficients $\delta_i(x)$ by $\ell^1$-minimization. The ratio between the two smallest residuals is about 1:8.6.](image-url)
**Example 1 (ℓ2-minimization versus ℓ1-minimization).** To illustrate how Algorithm 1 works, we randomly select half of the 2,414 images in the Extended Yale B database as the training set and the rest for testing. In this example, we subsample the images from the original 192 × 168 to size 12 × 10. The pixel values of the downsampled image are used as 120-D features—stacked as columns of the matrix A in the algorithm. Hence, matrix A has size 120 × 1,207, and the system \( y = Ax \) is underdetermined. Fig. 3a illustrates the sparse coefficients recovered by Algorithm 1 for a test image from the first subject. The figure also shows the features and the original images that correspond to the two largest coefficients. The two largest coefficients are both associated with training samples from subject 1. Fig. 3b shows the residuals with respect to the 38 projected coefficients \( \delta_i(\hat{x}) \), \( i = 1, 2, \ldots, 38 \). With 12 × 10 downsampled images as features, Algorithm 1 achieves an overall recognition rate of 92.1 percent across the Extended Yale B database. (See Section 4 for details and performance with other features such as Eigenfaces and Fisherfaces, as well as comparison with other methods.) Whereas the more conventional minimum \( \ell^2 \)-norm solution to the underdetermined system \( y = Ax \) is typically quite dense, minimizing the \( \ell^1 \)-norm favors sparse solutions and provably recovers the sparsest solution when this solution is sufficiently sparse. To illustrate this contrast, Fig. 4a shows the coefficients of the same test image given by the conventional \( \ell^2 \)-minimization (4), and Fig. 4b shows the corresponding residuals with respect to the 38 subjects. The coefficients are much less sparse than those given by \( \ell^1 \)-minimization (in Fig. 3), and the dominant coefficients are not associated with subject 1. As a result, the smallest residual in Fig. 4 does not correspond to the correct subject (subject 1).

2.4 Validation Based on Sparse Representation

Before classifying a given test sample, we must first decide if it is a valid sample from one of the classes in the data set. The ability to detect and then reject invalid test samples, or “outliers,” is crucial for recognition systems to work in real-world situations. A face recognition system, for example, could be given a face image of a subject that is not in the database or an image that is not a face at all.

Systems based on conventional classifiers such as NN or NS, often use the residuals \( r_i(y) \) for validation, in addition to identification. That is, the algorithm accepts or rejects a test sample based on how small the smallest residual is. However, each residual \( r_i(y) \) is computed without any knowledge of images of other object classes in the training data set and only measures similarity between the test sample and each individual class.

In the sparse representation paradigm, the coefficients \( \hat{x} \) are computed globally, in terms of images of all classes. In a sense, it can harness the joint distribution of all classes for validation. We contend that the coefficients \( \hat{x} \) are better statistics for validation than the residuals. Let us first see this through an example.

**Example 2 (concentration of sparse coefficients).** We randomly select an irrelevant image from Google and downsample it to 12 × 10. We then compute the sparse representation of the image against the same Extended Yale B training data, as in Example 1. Fig. 5a plots the obtained coefficients, and Fig. 5b plots the corresponding residuals. Compared to the coefficients of a valid test image in Fig. 3, notice that the coefficients \( \hat{x} \) here are not concentrated on any one subject and instead spread widely across the entire training set. Thus, the distribution of the estimated sparse coefficients \( \hat{x} \) contains important information about the validity of the test image: a valid test image should have a sparse representation whose nonzero entries concentrate mostly on one subject, whereas an invalid image has sparse coefficients spread widely among multiple subjects.

To quantify this observation, we define the following measure of how concentrated the coefficients are on a single class in the data set:

**Definition 1 (sparsity concentration index (SCI)).** The SCI of a coefficient vector \( x \in \mathbb{R}^n \) is defined as

\[
SCI(x) \triangleq \frac{k \cdot \max_i ||\delta_i(x)||_1/||x||_1 - 1}{k - 1} \in [0, 1].
\] (14)

For a solution \( \hat{x} \) found by Algorithm 1, if SCI(\( \hat{x} \)) = 1, the test image is represented using only images from a single object, and if SCI(\( \hat{x} \)) = 0, the sparse coefficients are spread evenly over all classes.\(^9\) We choose a threshold \( \tau \in (0, 1) \) and accept a test image as valid if

\(^9\) Directly choosing \( x \) to minimize the SCI might produce more concentrated coefficients; however, the SCI is highly nonconvex and difficult to optimize. For valid test images, minimizing the \( \ell^0 \)-norm already produces representations that are well-concentrated on the correct subject class.
and otherwise reject as invalid. In step 5 of Algorithm 1, one may choose to output the identity of \( y \) only if it passes this criterion.

Unlike NN or NS, this new rule avoids the use of the residuals \( r_i(y) \) for validation. Notice that in Fig. 5, even for a nonface image, with a large training set, the smallest residual of the invalid test image is not so large. Rather than relying on a single statistic for both validation and identification, our approach separates the information required for these tasks: the residuals for identification and the sparse coefficients for validation.\(^{10}\) In a sense, the residual measures how well the representation approximates the test image; and the sparsity concentration index measures how good the representation itself is, in terms of localization.

One benefit to this approach to validation is improved performance against generic objects that are similar to multiple object classes. For example, in face recognition, a generic face might be rather similar to some of the subjects in the data set and may have small residuals with respect to their training images. Using residuals for validation more likely leads to a false positive. However, a generic face is unlikely to pass the new validation rule as a good representation of it typically requires contribution from images of multiple subjects in the data set. Thus, the new rule can better judge whether the test image is a generic face or the face of one particular subject in the data set. In Section 4.7, we will demonstrate that the new validation rule outperforms the NN and NS methods, with as much as 10-20 percent improvement in verification rate for a given false accept rate (see Fig. 14 in Section 4 or Fig. 18 in the supplementary appendix, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79).

### 3 Two Fundamental Issues in Face Recognition

In this section, we study the implications of the above general classification framework for two critical issues in face recognition: 1) the choice of feature transformation, and 2) robustness to corruption, occlusion, and disguise.

\(^{10}\) We find empirically that this separation works well enough in our experiments with face images. However, it is possible that better validation and identification rules can be contrived from using the residual and the sparsity together.

#### 3.1 The Role of Feature Extraction

In the computer vision literature, numerous feature extraction schemes have been investigated for finding projections that better separate the classes in lower dimensional spaces, which are often referred to as feature spaces. One class of methods extracts holistic face features such as Eigenfaces\(^{23}\), Fisherfaces \(^{24}\), and Laplacianfaces \(^{25}\). Another class of methods tries to extract meaningful partial facial features (e.g., patches around eyes or nose) \(^{21},^{41}\) (see Fig. 6 for some examples). Traditionally, when feature extraction is used in conjunction with simple classifiers such as NN and NS, the choice of feature transformation is considered critical to the success of the algorithm. This has led to the development of a wide variety of increasingly complex feature extraction methods, including nonlinear and kernel features \(^{42},^{43}\). In this section, we reexamine the role of feature extraction within the new sparse representation framework for face recognition.

One benefit of feature extraction, which carries over to the proposed sparse representation framework, is reduced data dimension and computational cost. For raw face images, the corresponding linear system \( y = Ax \) is very large. For instance, if the face images are given at the typical resolution, \( 640 \times 480 \) pixels, the dimension \( m \) is in the order of \( 10^6 \). Although Algorithm 1 relies on scalable methods such as linear programming, directly applying it to such high-resolution images is still beyond the capability of regular computers.

Since most feature transformations involve only linear operations (or approximately so), the projection from the image space to the feature space can be represented as a matrix \( R \in \mathbb{R}^{d \times m} \) with \( d \ll m \). Applying \( R \) to both sides of (3) yields

\[
\hat{y} = Ry = RAx_0 \in \mathbb{R}^d.
\]

In practice, the dimension \( d \) of the feature space is typically chosen to be much smaller than \( n \). In this case, the system of equations \( \hat{y} = RAx \in \mathbb{R}^d \) is underdetermined in the unknown \( x \in \mathbb{R}^n \). Nevertheless, as the desired solution \( x_0 \) is sparse, we can hope to recover it by solving the following reduced \( \ell^1 \)-minimization problem:

\[
(\ell_1^1) : \quad \hat{x}_1 = \arg \min \| x \|_1 \quad \text{subject to} \quad \| RAx - \hat{y} \|_2 \leq \varepsilon,
\]

for a given error tolerance \( \varepsilon > 0 \). Thus, in Algorithm 1, the matrix \( A \) of training images is now replaced by the matrix...
distribution of features space generally improves the recognition rate, as long as the have shown that increasing the dimension replaced by its features.

Fig. 6. Examples of feature extraction. (a) Original face image. (b) 120D representations in terms of four different features (from left to right): Eigenfaces, Laplacianfaces, downsampled (12 x 10 pixel) image, and random projection. We will demonstrate that all these features contain almost the same information about the identity of the subject and give similarly good recognition performance. (c) The eye is a popular choice of feature for face recognition. In this case, the feature matrix $R$ is simply a binary mask. (a) Original $y$. (b) 120D features $\tilde{y} = ry$. (c) Eye feature $y$.  

$RA \in \mathbb{R}^{d \times n}$ of $d$-dimensional features; the test image $y$ is replaced by its features $\tilde{y}$.

For extant face recognition methods, empirical studies have shown that increasing the dimension $d$ of the feature space generally improves the recognition rate, as long as the distribution of features $RA$, does not become degenerate [42]. Degeneracy is not an issue for $\ell^1$-minimization, since it merely requires that $\tilde{y}$ be in or near the range of $RA$,—it does not depend on the covariance $\Sigma_{1} = A A^{T} RA$, being nonsingular as in classical discriminant analysis. The stable version of $\ell^1$-minimization (10) or (17) is known in statistical literature as the Lasso [14].

It effectively regularizes highly underdetermined linear regression when the desired solution is sparse and has also been proven consistent in some noisy overdetermined settings [12].

For our sparse representation approach to recognition, we would like to understand how the choice of the feature extraction $R$ affects the ability of the $\ell^1$-minimization (17) to recover the correct sparse solution $x_0$. From the geometric interpretation of $\ell^1$-minimization given in Section 2.2.1, the answer to this depends on whether the associated new polytope $P = RA(P)$ remains sufficiently neighborly. It is easy to show that the neighborliness of the polytope $P = RA(P)$ increases with $d$ [11], [15]. In Section 4, our experimental results will verify the ability of $\ell^1$-minimization, in particular, the stable version (17), to recover sparse representations for face recognition using a variety of features. This suggests that most data-dependent features popular in face recognition (e.g., eigenfaces and Laplacianfaces) may indeed give highly neighborly polytopes $P$.

Further analysis of high-dimensional polytope geometry has revealed a somewhat surprising phenomenon: if the solution $x_0$ is sparse enough, then with overwhelming probability, it can be correctly recovered via $\ell^1$-minimization from any sufficiently large number $d$ of linear measurements $\tilde{y} = RAx_0$. More precisely, if $x_0$ has $t \ll n$ nonzeros, then with overwhelming probability

$$d \geq 2t \log(n/d)$$

11. Classically, the Lasso solution is defined as the minimizer of $\|y - Ax\|^2 + \lambda \|x\|_1$. Here, $\lambda$ can be viewed as inverse of the Lagrange multiplier associated with a constraint $\|y - Ax\|_2^2 \leq \varepsilon$. For every $\lambda$, there is an $\varepsilon$ such that the two problems have the same solution. However, $\varepsilon$ can be interpreted as a pixel noise level and fixed across various instances of the problem, whereas $\lambda$ cannot. One should distinguish the Lasso optimization problem from the LARS algorithm, which provably solves some instances of Lasso with very sparse optimizers [35].

random linear measurements are sufficient for $\ell^1$-minimization (17) to recover the correct sparse solution $x_0$. This surprising phenomenon has been dubbed the “blessing of dimensionality” [15], [46]. Random features can be viewed as a less-structured counterpart to classical face features such as Eigenfaces or Fisherfaces. Accordingly, we call the linear projection generated by a Gaussian random matrix Randomfaces.

**Definition 2 (Randomfaces).** Consider a transform matrix $R \in \mathbb{R}^{d \times n}$ whose entries are independently sampled from a zero-mean normal distribution, and each row is normalized to unit length. The row vectors of $R$ can be viewed as $d$ random faces in $\mathbb{R}^n$.

One major advantage of Randomfaces is that they are extremely efficient to generate, as the transformation $R$ is independent of the training data set. This advantage can be important for a face recognition system, where we may not be able to acquire a complete database of all subjects of interest to precompute data-dependent transformations such as Eigenfaces, or the subjects in the database may change over time. In such cases, there is no need for recomputing the random transformation $R$.

As long as the correct sparse solution $x_0$ can be recovered, Algorithm 1 will always give the same classification result, regardless of the feature actually used. Thus, when the dimension of feature $d$ exceeds the above bound (18), one should expect that the recognition performance of Algorithm 1 with different features quickly converges, and the choice of an “optimal” feature transformation is no longer critical: even random projections or downsampling images should perform as well as any other carefully engineered features. This will be corroborated by the experimental results in Section 4.

### 3.2 Robustness to Occlusion or Corruption

In many practical face recognition scenarios, the test image $y$ could be partially corrupted or occluded. In this case, the above linear model (3) should be modified as

$$y = y_0 + e_0 = Ax_0 + e_0,$$

12. Strictly speaking, this threshold holds when random measurements are computed directly from $x_0$, i.e., $\tilde{y} = RAx_0$. Nevertheless, our experiments roughly agree with the bound given by (18). The case where $x_0$ is instead sparse in some overcomplete basis $A$, and we observe that random measurements $\tilde{y} = RAx_0$ has also been studied in [43]. While conditions for correct recovery have been given, the bounds are not yet as sharp as (18) above.

13. Random projection has been previously studied as a general dimensionality-reduction method for numerous clustering problems [47], [48], [49], as well as for learning nonlinear manifolds [50], [51].
where $e_0 \in \mathbb{R}^m$ is a vector of errors—a fraction, $\rho$, of its entries are nonzero. The nonzero entries of $e_0$ model which pixels in $y$ are corrupted or occluded. The locations of corruption can differ for different test images and are not known to the computer. The errors may have arbitrary magnitude and therefore cannot be ignored or treated with techniques designed for small noise such as the one given in Section 2.2.2.

A fundamental principle of coding theory [52] is that redundancy in the measurement is essential to detecting and correcting gross errors. Redundancy arises in object recognition because the number of image pixels is typically far greater than the number of subjects that have generated the images. In this case, even if a fraction of the pixels are completely corrupted by occlusion, recognition may still be possible based on the remaining pixels. On the other hand, feature extraction schemes discussed in the previous section would discard useful information that could help compensate for the occlusion. In this sense, no representation is more redundant, robust, or informative than the original images. Thus, when dealing with occlusion and corruption, we should always work with the highest possible resolution, performing downsampling or feature extraction only if the resolution of the original images is too high to process.

Of course, redundancy would be of no use without efficient computational tools for exploiting the information encoded in the redundant data. The difficulty in directly harnessing the redundancy in corrupted raw images has led researchers to instead focus on spatial locality as a guiding principle for robust recognition. Local features computed from only a small fraction of the image pixels are clearly less likely to be corrupted by occlusion than holistic features. In face recognition, methods such as ICA [53] and LNMF [54] exploit this observation by adaptively choosing filter bases that are locally concentrated. Local Binary Patterns [55] and Gabor wavelets [56] exhibit similar properties, since they are also computed from local image regions. A related approach partitions the image into fixed regions and computes features for each region [16], [57]. Notice, though, that projecting onto locally concentrated bases transforms the domain of the occlusion problem, rather than eliminating the occlusion. Errors on the original pixels become errors in the transformed domain and may even become less local. The role of feature extraction in achieving spatial locality is therefore questionable, since no bases or features are more spatially localized than the original image pixels themselves. In fact, the most popular approach to robustifying feature-based methods is based on randomly sampling individual pixels [28], sometimes in conjunction with statistical techniques such as multivariate trimming [29].

Now, let us show how the proposed sparse representation classification framework can be extended to deal with occlusion. Let us assume that the corrupted pixels are a relatively small portion of the image. The error vector $e_0$, like the vector $x_0$, then has sparse14 nonzero entries. Since $y_i = Ax_0$, we can rewrite (19) as

$$y = [A, I][x_0 \; e_0] = Bw_0.$$ (20)

Here, $B = [A, I] \in \mathbb{R}^{m \times (n+m)}$, so the system $y = Bw$ is always underdetermined and does not have a unique solution for $w$. However, from the above discussion about the sparsity of $x_0$ and $e_0$, the correct generating $w_0 = [x_0, e_0]$ has at most $n_1 + \rho n_0$ nonzeros. We might therefore hope to recover $w_0$ as the sparsest solution to the system $y = Bw$. In fact, if the matrix $B$ is in general position, then as long as $y = B\tilde{w}$ for some $\tilde{w}$ with less than $n_1 + \rho n_0$ nonzeros, $\tilde{w}$ is the unique sparsest solution. Thus, if the occlusion $e$ covers less than $\frac{n_1 + \rho n_0}{2}$ pixels, $\approx 50$ percent of the image, the sparsest solution $\hat{w}$ to $y = Bw$ is the true generator, $w_0 = [x_0, e_0]$.

More generally, one can assume that the corrupting error $e_0$ has a sparse representation with respect to some basis $A_e \in \mathbb{R}^{m \times n_e}$. That is, $e_0 = A_e u_0$ for some sparse vector $u_0 \in \mathbb{R}^{n_e}$. Here, we have chosen the special case $A_e = I \in \mathbb{R}^{m \times n_e}$ as $e_0$ is assumed to be sparse with respect to the natural pixel coordinates. If the error $e_0$ is instead more sparse with respect to another basis, e.g., Fourier or Haar, we can simply redefine the matrix $B$ by appending $A_e$ (instead of the identity $I$) to $A$ and instead seek the sparsest solution $w_0$ to the equation:

$$y = Bw$$  with  $$B = [A, A_e] \in \mathbb{R}^{m \times (n+n_e)}.$$ (21)

In this way, the same formulation can handle more general classes of (sparse) corruption.

As before, we attempt to recover the sparsest solution $w_0$ from solving the following extended $\ell^1$-minimization problem:

$$\hat{w}_1 = \arg \min \|w\|_1 \text{ subject to } Bw = y.$$ (22)

That is, in Algorithm 1, we now replace the image matrix $A$ with the extended matrix $B = [A, I]$ and $x$ with $w = [x, e]$.

Clearly, whether the sparse solution $w_0$ can be recovered from the above $\ell^1$-minimization depends on the neighborhood of the new polytope $P = B(P_i) = [A, I](P_i)$. This polytope contains vertices from both the training images $A$ and the identity matrix $I$, as illustrated in Fig. 7. The bounds given in (8) imply that if $y$ is an image of subject $i$, the $\ell^1$-minimization (22) cannot guarantee to correctly recover $w_0 = [x_0, e_0]$ if

$$n_i + |\text{support}(e_0)| > d/3.$$ (23)

Generally, $d \gg n_i$, so, (8) implies that the largest fraction of occlusion under which we can hope to still achieve perfect reconstruction is 33 percent. This bound is corroborated by our experimental results, see Fig. 12.

To know exactly how much occlusion can be tolerated, we need more accurate information about the neighborhood of the polytope $P$ than a loose upper bound given by (8). For instance, we would like to know for a given set of training images, what is the largest amount of (worst possible) occlusion it can handle. While the best known algorithms for exactly computing the neighborhood of a polytope are combinatorial in nature, tighter upper bounds can be obtained by restricting the search for intersections between the nullspace of $B$ and the $\ell^1$-ball to a random subset of the $t$-faces of the $\ell^1$-ball (see [37] for details). We
contribute to the representation; all other vertices have no contribution. The facet with the fewest possible vertices. Only vertices of that facet contribute to the representation; all other vertices have no contribution.

We test our SRC algorithm using several conventional classifiers. We will then demonstrate the robustness of the proposed algorithm to corruption and occlusion.

Finally, we demonstrate (using ROC curves) the effectiveness of sparsity as a means of validating test images and examine how to choose training sets to maximize robustness to occlusion.

4.1.1 Extended Yale B Database
The Extended Yale B database consists of 2,414 frontal-face images of 38 individuals [58]. The cropped and normalized 192 × 168 face images were captured under various laboratory-controlled lighting conditions [59]. For each subject, we randomly select half of the images for training (i.e., about 32 images per subject) and the other half for testing. Randomly choosing the training set ensures that our results and conclusions will not depend on any special choice of the training data.

We compute the recognition rates with the feature space dimensions 30, 56, 120, and 504. Those numbers correspond to downsampling ratios of 1/32, 1/24, 1/16, and 1/8, respectively. Notice that Fisherfaces are different from the other features because the maximal number of valid facial images is one less than the number of classes k [24], 38 in this case. As a result, the recognition result for Fisherfaces is only available at dimension 30 in our experiment.

The subspace dimension for the NS algorithm is 9, which has been mostly agreed upon in the literature for processing facial images with only illumination change. Fig. 8 shows the recognition performance for the various features, in conjunction with four different classifiers: SRC, NN, NS, and SVM.

SRC achieves recognition rates between 92.1 percent and 95.6 percent for all 120D feature spaces and a maximum rate of 98.1 percent with 504D randomfaces. The maximum recognition rates for NN, NS, and SVM are 90.7 percent, 94.1 percent, and 97.7 percent, respectively. Tables with all the recognition rates are available in the supplementary appendix, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79. The recognition rates shown in Fig. 8 are consistent with those that have been reported in the literature, although some reported on different databases or with different training subsets. For example, He et al. [25] reported the best recognition rate of 75 percent using Eigenfaces at 33D, and 89 percent using Laplacianfaces at unconventional features: randomfaces and downsamples images. We compare our algorithm with three classical algorithms, namely, NN, and NS, discussed in the previous section, as well as linear SVM.

Due to the subspace structure of face images, linear SVM is already appropriate for separating features from different faces. The use of a linear kernel (as opposed to more complicated nonlinear transformations) also makes it possible to directly compare between different algorithms working in the same feature space. Nevertheless, better performance might be achieved by using nonlinear kernels in addition to feature transformations.

We cut off the dimension at 504 as the computation of Eigenfaces and Laplacianfaces reaches the memory limit of Matlab. Although our algorithm persists to work far beyond on the same computer, 504 is already sufficient to reach all our conclusions.

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16. We cut off the dimension at 504 as the computation of Eigenfaces and Laplacianfaces reaches the memory limit of Matlab. Although our algorithm persists to work far beyond on the same computer, 504 is already sufficient to reach all our conclusions.

17. Subspace dimensions significantly greater or less than 9 eventually led to a decrease in performance.

18. We also experimented with replacing the constrained ℓ1-minimization in the SRC algorithm with the Lasso. For appropriate choice of regularization λ, the results are similar. For example, with downsampled faces as features and λ = 1,000, the recognition rates are 73.7 percent, 86.2 percent, 91.9 percent, 97.3 percent, at dimensions 30, 56, 120, and 504 (within 1 percent of the results in Fig. 8).
28D on the Yale face database, both using NN. In [32], Lee et al. reported 95.4 percent accuracy using the NS method on the Yale B database.

4.1.2 AR Database

The AR database consists of over 4,000 frontal images for 126 individuals. For each individual, 26 pictures were taken in two separate sessions [60]. These images include more facial variations, including illumination change, expressions, and facial disguises comparing to the Extended Yale B database. In the experiment, we chose a subset of the data set consisting of 50 male subjects and 50 female subjects. For each subject, 14 images with only illumination change and expressions were selected: the seven images from Session 1 for training, and the other seven from Session 2 for testing. The images are cropped with dimension 165×120 and converted to gray scale. We selected four feature space dimensions: 30, 54, 130, and 540, which correspond to the downsample ratios 1/24, 1/18, 1/12, and 1/6, respectively.

Because the number of subjects is 100, results for Fisherfaces are only given at dimension 30 and 54.

This database is substantially more challenging than the Yale database, since the number of subjects is now 100, but the training images is reduced to seven per subject: four neutral faces with different lighting conditions and three faces with different expressions. For NS, since the number of training images per subject is seven, any estimate of the face subspace cannot have dimension higher than 7. We chose to keep all seven dimensions for NS in this case.

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Fig. 9 shows the recognition rates for this experiment. With 540D features, SRC achieves a recognition rate between 92.0 percent and 94.7 percent. On the other hand, the best rates achieved by NN and NS are 89.7 percent and 90.3 percent, respectively. SVM slightly outperforms SRC on this data set, achieving a maximum recognition rate of 95.7 percent. However, the performance of SVM varies more with the choice of feature space—the recognition rate using random features is just 88.8 percent. The supplementary appendix, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79, contains a table of detailed numerical results.

Based on the results on the Extended Yale B database and the AR database, we draw the following conclusions:

1. For both the Yale database and AR database, the best performances of SRC and SVM consistently exceed the best performances of the two classical methods NN and NS at each individual feature dimension. More specifically, the best recognition rate for SRC on the Yale database is 98.1 percent, compared to 97.7 percent for SVM, 94.0 percent for NS, and 90.7 percent for NN; the best rate for SRC on the AR database is 94.7 percent, compared to 95.7 percent for SVM, 90.3 percent for NS, and 89.7 percent for NN.

2. The performances of the other three classifiers depends strongly on a good choice of “optimal” features—Fisherfaces for lower feature space dimension and Laplacianfaces for higher feature space dimension. With NN and SVM, the performance of the various features does not converge as the dimension of the feature space increases.

3. The results corroborate the theory of compressed sensing: (18) suggests that $d \approx 128$ random linear measurements should suffice for sparse recovery in the Yale database, while $d \approx 88$ random linear measurements should suffice for sparse recovery in the AR database [44]. Beyond these dimensions, the performances of various features in conjunction with $\ell^1$-minimization converge, with conventional and unconventional features (e.g., Randomfaces and downsampled images) performing similarly. When
the feature dimension is large, a single random projection performs the best (98.1 percent recognition rate on Yale, 94.7 percent on AR).

### 4.2 Partial Face Features

Here have been extensive studies in both the human and computer vision literature about the effectiveness of partial features in recovering the identity of a human face, e.g., see [21] and [41]. As a second set of experiments, we test our algorithm on the following three partial facial features: nose, right eye, and mouth and chin. We use the Extended Yale B database for the experiment, with the same training and test sets, as in Section 4.1.1. See Fig. 10 for a typical example of the extracted features.

For each of the three features, the dimension \(d\) is larger than the number of training samples \(n = 1,207\), and the linear system (16) to be solved becomes overdetermined. Nevertheless, sparse approximate solutions \(x\) can still be obtained by solving the \(\epsilon\)-relaxed \(\ell^1\)-minimization problem (17) (here, again, \(\epsilon = 0.05\)). The results in Fig. 10 right again show that the proposed SRC algorithm achieves better recognition rates than NN, NS, and SVM. These experiments also show the scalability of the proposed algorithm in working with more than \(10^4\)-dimensional features.

### 4.3 Recognition Despite Random Pixel Corruption

For this experiment, we test the robust version of SRC, which solves the extended \(\ell^1\)-minimization problem (22) using the Extended Yale B Face Database. We choose Subsets 1 and 2 (717 images, normal-to-moderate lighting conditions) for training and Subset 3 (453 images, more extreme lighting conditions) for testing. Without occlusion, this is a relatively easy recognition problem. This choice is deliberate, in order to isolate the effect of occlusion. The images are resized to 96 \times 84 pixels, so in this case, \(B = [A, I]\) is an 8,064 \times 8,761 matrix. For this data set, we have estimated that the polytope \(P = \text{conv}(\pm B)\) is approximately 1,185 neighborly (using the method given in [37]), suggesting that perfect reconstruction can be achieved up to 13.3 percent (worst possible) occlusion.

We corrupt a percentage of randomly chosen pixels from each of the test images, replacing their values with independent and identically distributed samples from a uniform distribution. The corrupted pixels are randomly chosen for each test image, and the locations are unknown to the algorithm. We vary the percentage of corrupted pixels from 0 percent to 90 percent. Figs. 11a, 11b, 11c, and 11d shows several example test images. To the human eye, beyond 50 percent corruption, the corrupted images (Fig. 11a second and third rows) are

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#### Table: Recognition Rates

<table>
<thead>
<tr>
<th>Features</th>
<th>Nose</th>
<th>Right Eye</th>
<th>Mouth &amp; Chin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimension ((d))</td>
<td>4,270</td>
<td>5,040</td>
<td>12,936</td>
</tr>
<tr>
<td>SRC</td>
<td>87.3%</td>
<td>93.7%</td>
<td>98.3%</td>
</tr>
<tr>
<td>NN</td>
<td>49.2%</td>
<td>68.8%</td>
<td>72.7%</td>
</tr>
<tr>
<td>NS</td>
<td>83.7%</td>
<td>78.6%</td>
<td>94.4%</td>
</tr>
<tr>
<td>SVM</td>
<td>70.8%</td>
<td>85.8%</td>
<td>95.3%</td>
</tr>
</tbody>
</table>

---

19. The only reason for resizing the images is to be able to run all the experiments within the memory size of Matlab on a typical PC. The algorithm relies on linear programming and is scalable in the image size.

20. Uniform over \([0, y_{\text{max}}]\), where \(y_{\text{max}}\) is the largest possible pixel value.
barely recognizable as face images; determining their identity seems out of the question. Nevertheless, even in this extreme circumstance, SRC correctly recovers the identity of the subjects.

We quantitatively compare our method to four popular techniques for face recognition in the vision literature. The Principal Component Analysis (PCA) approach in [23] is not robust to occlusion. There are many variations to make PCA robust to corruption or incomplete data, and some have been applied to robust face recognition, e.g., [29]. We will later discuss their performance against ours on more realistic conditions. Here, we use the basic PCA to provide a standard baseline for comparison. The remaining three techniques are designed to be more robust to occlusion. Independent Component Analysis (ICA) architecture I [53] attempts to express the training set as a linear combination of statistically independent basis images. Local Nonnegative Matrix Factorization (LNMF) [54] approximates the training set as an additive combination of basis images, computed with a bias toward sparse bases. Finally, to demonstrate that the improved robustness is really due to the $\ell^1$-norm, we compare to a least-squares technique that first projects the test image onto the subspace spanned by all face images and then performs NS.

21. Following [58], we normalize the image pixels to have zero mean and unit variance before applying PCA.

22. For PCA, ICA, and LNMF, the number of basis components is chosen to give the optimal test performance over the range $\{100, 200, 300, 400, 500, 600\}$. This greatly surpasses the theoretical bound of the worst-case corruption (13.3 percent) that the algorithm is ensured to tolerate. Clearly, the worst-case analysis is too conservative for random corruption.

### 4.4 Recognition Despite Random Block Occlusion

We next simulate various levels of contiguous occlusion, from 0 percent to 50 percent, by replacing a randomly located square block of each test image with an unrelated image, as in Fig. 12a. Again, the location of occlusion is randomly chosen for each image and is unknown to the computer. Methods that select fixed facial features or blocks of the image (e.g., [16] and [57]) are less likely to succeed here due to the unpredictable location of the occlusion. The top two rows in Figs. 12a, 12b, 12c, and 12d shows the two representative results of Algorithm 1 with 30 percent occlusion. Fig. 12a is the occluded image. In the second row, the entire center of the face is occluded; this is a difficult recognition task even for humans. Fig. 12b shows the magnitude of the estimated error $\hat{e}_1$. Notice that $\hat{e}_1$ compensates not only for occlusion due to the baboon but also for the violation of the linear subspace model caused by the shadow under the nose. Fig. 12c plots the estimated coefficient vector $\hat{x}_1$. The red entries are coefficients corresponding to test image’s true class. In both examples, the estimated coefficients are indeed sparse and have large magnitude only for training images of the same person. In both cases, the SRC algorithm correctly classifies the occluded image. For this data set, our Matlab implementation requires 90 seconds per test image on a PowerMac G5.

The graph in Fig. 12e shows the recognition rates of all six algorithms. SRC again significantly outperforms the other five methods for all levels of occlusion.upto 30 percent occlusion, Algorithm 1 performs almost perfectly, correctly identifying over 98 percent of test subjects. Even at 40 percent occlusion, only 9.7 percent of subjects are misclassified. Compared to the random pixel corruption, contiguous
Recall that the problem is not to recover the corrupted image but rather to express an occluded image in terms of a holistic basis such as the training images themselves, all of the coefficients may be corrupted (as in Fig. 12 third row). The implication here is that proper harnessing redundancy and sparsity is the key to error correction and robustness. Extracting local or disjoint features can only reduce redundancy, resulting in inferior robustness.

4.5 Recognition Despite Disguise

We test SRC’s ability to cope with real possibly malicious occlusions using a subset of the AR Face Database. The chosen subset consists of 1,399 images (14 each, except for a corrupted image w-027-14.jpg) of 100 subjects, 50 male and 50 female. For training, we use 799 images (about 8 per subject) of unoccluded frontal views with varying facial expression, giving a matrix $B$ of size $4,980 \times 5,779$. We estimate $P = \text{conv}(\pm B)$ is approximately 577 neighborly, indicating that perfect reconstruction is possible up to 11.6 percent occlusion. Our Matlab implementation requires about 75 seconds per test image on a PowerMac G5.

We consider two separate test sets of 200 images. The first test set contains images of the subjects wearing sunglasses, which occlude roughly 20 percent of the image. Fig. 1a shows a successful example from this test set. Notice that $\hat{e}_1$ compensates for small misalignment of the image edges, as well as occlusion due to sunglasses. The second test set considered contains images of the subjects wearing a scarf, which occludes roughly 40 percent of the image. Since the occlusion level is more than three times the maximum worst case occlusion given by the neighborliness of $\text{conv}(\pm B)$, our approach is unlikely to succeed in this domain. Fig. 13a shows one such failure. Notice that the largest coefficient corresponds to an image of a bearded man whose mouth region resembles the scarf.

The table in Fig. 13 left compares SRC to the other five algorithms described in the previous section. On faces occluded by sunglasses, SRC achieves a recognition rate of 87 percent, more than 17 percent better than the nearest competitor. For occlusion by scarves, its recognition rate is 59.5 percent, more than double its nearest competitor but still quite poor. This confirms that although the algorithm is provably robust to arbitrary occlusions up to the breakdown point determined by the neighborliness of the training set, beyond that point, it is sensitive to occlusions that resemble regions of a training image from a different individual. Because the amount of occlusion exceeds this breakdown point, additional assumptions, such as the disguise is likely to be contiguous, are needed to achieve higher recognition performance.

4.6 Improving Recognition by Block Partitioning

Thus far, we have not exploited the fact that in many real recognition scenarios, the occlusion falls on some patch of image pixels which is a priori unknown but is known to be connected. A somewhat traditional approach (explored in [57] among others) to exploiting this information in face recognition is to partition the image into blocks and process each block independently. The results for individual blocks are then aggregated, for example, by voting, while discarding blocks believed to be occluded (using, say, the outlier
We apply the classifier from Algorithm 1 within each block. We verify the efficacy of this scheme on the AR database for faces disguised with sunglasses or scarves. We partition the images into eight \((4 \times 2)\) blocks of size \(20 \times 30\) pixels. Partitioning increases the recognition rate on scarves from 59.5 percent to 93.5 percent and also improves the recognition rate on sunglasses from 87.0 percent to 97.5 percent. This performance exceeds the best known results on the AR data set [29] to date. That work obtains 84 percent on the sunglasses and 93 percent on the scarfs, on only 50 subjects, using more sophisticated random sampling techniques. Also noteworthy is [16], which aims to recognize occluded faces from only a single training sample per subject. On the AR database, that method achieves a lower combined recognition rate of 80 percent.\footnote{From our own implementation and experiments, we find that this does not significantly increase the recognition rate.}

### 4.7 Rejecting Invalid Test Images

We next demonstrate the relevance of sparsity for rejecting invalid test images, with or without occlusion. We test the outlier rejection rule (15) based on the Sparsity Concentration Index (14) on the Extended Yale B database, using Subsets 1 and 2 for training and Subset 3 for testing as before. We again simulate varying levels of occlusion (10 percent, 30 percent, and 50 percent) by replacing a randomly chosen block of each test image with an unrelated image. However, in this experiment, we include only half of the subjects in the training set. Thus, half of the subjects in the testing set are new to the algorithm. We test the system’s ability to determine whether a given test subject is in the training database or not by sweeping the threshold \(\tau\) through a range of values in \([0, 1]\), generating the receiver operator characteristic (ROC) curves in Fig. 14. For comparison, we also considered outlier rejection by thresholding the euclidean distance between (features of) the test image and (features of) the nearest training images within the PCA, ICA, and LNMF feature spaces. These curves are also displayed in Fig. 14. Notice that the simple rejection rule (15) performs nearly perfectly at 10 percent and 30 percent occlusion. At 50 percent occlusion, it still significantly outperforms the other three algorithms and is the only one of the four algorithms that performs significantly better than chance. The supplementary appendix, which can be found on the Computer Society Digital Library at http://doi.ieeecomputersociety.org/10.1109/TPAMI.2008.79, contains more validation results on the AR database using Eigenfaces, again demonstrating significant improvement in the ROC.

### 4.8 Designing the Training Set for Robustness

An important consideration in designing recognition systems is selecting the number of training images, as well as the conditions (lighting, expression, viewpoint, etc.) under which they are to be taken. The training images should be extensive enough to span the conditions that might occur in the test set; they should be “sufficient” from a pattern recognition standpoint. For instance, Lee et al. [59] shows how to choose the fewest representative images to well approximate the illumination cone of each face. The notion of neighborliness discussed in Section 2 provides a different quantitative measure for how “robust” the training set is: the amount of worst case occlusion the algorithm can tolerate is directly determined by how neighborly the associated polytope is. The worst case is relevant in visual recognition, and the polytope is. The worst case is relevant in visual recognition,
since the occluding object could potentially be quite similar
to one of the other training classes. However, if the occlusion
is random and uncorrelated with the training images, as in
Section 4.3, the average behavior may also be of interest.

In fact, these two concerns, sufficiency and robustness,
are complementary. Fig. 15a shows the estimated neighbor-
liness for the four subsets of the Extended Yale B database.
Notice that the highest neighborliness, \( \frac{1}{C} \mathbf{25} \) \( \mathbf{1}, \mathbf{3}, \mathbf{3} \mathbf{0} \), is achieved
with Subset 4, the most extreme lighting conditions. Fig. 15b
shows the breakdown point for subsets of the AR database
with different facial expressions. The data set contains four
facial expressions, Neutral, Happy, Angry, and Scream,
pictured in Fig. 15b. We generate training sets from all pairs
of expressions and compute the neighborliness of each of
the corresponding polytopes. The most robust training sets
are achieved by the Neutral+Happy and Happy+Scream
combinations, while the least robustness comes from
Neutral+Angry. Notice that the Neutral and Angry images
are quite similar in appearance, while (for example) Happy
and Scream are very dissimilar.

Thus, both for varying lighting (Fig. 15a) and expression
(Fig. 15b), training sets with wider variation in the images
allow greater robustness to occlusion. Designing a training
set that allows recognition under widely varying conditions
does not hinder our algorithm; in fact, it helps it. However,
the training set should not contain too many similar images,
as in the Neutral+Angry example in Fig. 15b. In the
language of signal representation, the training images
should form an incoherent dictionary \([9]\).

5 CONCLUSIONS AND DISCUSSIONS

In this paper, we have contended both theoretically and
experimentally that exploiting sparsity is critical for the
high-performance classification of high-dimensional data
such as face images. With sparsity properly harnessed, the
choice of features becomes less important than the number
of features used (in our face recognition example, approxi-
mately 100 are sufficient to make the difference negligible).
Moreover, occlusion and corruption can be handled
uniformly and robustly within the same classification
framework. One can achieve a striking recognition perfor-
mance for severely occluded or corrupted images by a
simple algorithm with no special engineering.

An intriguing question for future work is whether this
framework can be useful for object detection, in addition to
recognition. The usefulness of sparsity in detection has been
noticed in the work in [61] and more recently explored in
[62]. We believe that the full potential of sparsity in robust
object detection and recognition together is yet to be
uncovered. From a practical standpoint, it would also be
useful to extend the algorithm to less constrained condi-
tions, especially variations in object pose. Robustness to
occlusion allows the algorithm to tolerate small pose
variation or misalignment. Furthermore, in the supplemen-
tary appendix, which can be found on the Computer Society
Digital Library at http://doi.ieeecomputersociety.org/
10.1109/TPAMI.2008.79, we discuss our algorithm’s ability
to adapt to nonlinear training distributions. However, the
number of training samples required to directly represent
the distribution of face images under varying pose may be
prohibitively large. Extrapolation in pose, e.g., using only
frontal training images, will require integrating feature
matching techniques or nonlinear deformation models into
the computation of the sparse representation of the test
image. Doing so, in a principled manner, it remains an
important direction for future work.

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![Fig. 14. ROC curves for outlier rejection. Vertical axis: true positive rate. Horizontal axis: false positive rate. The solid red curve is generated by SRC with outliers rejected based on (15). The SCI-based validation and SRC classification together perform almost perfectly for up to 30 percent occlusion. (a) No occlusion. (b) Ten percent occlusion. (c) Thirty percent. (d) Fifty percent.](image)

![Fig. 15. Robust training set design. (a) Varying illumination. Top left: four subsets of Extended Yale B, containing increasingly extreme lighting conditions. Bottom left: estimated neighborliness of the polytope \( \text{conv}(\pm B) \) for each subset. (b) Varying expression. Top right: four facial expressions in the AR database. Bottom right: estimated neighborliness of \( \text{conv}(\pm B) \) when taking the training set from different pairs of expressions.](image)
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