Group Component Analysis for Multi-block Data: Common and Individual Feature Extraction

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Abstract—Real world data are often acquired as a collection of matrices rather than as a single matrix. Such multi-block data are naturally linked and typically share some common features and at the same time exhibit their own individual features, reflecting the background in which they are measured and collected. To exploit the linked nature of data, we propose a new framework for common and individual feature extraction (CIFE) which identifies and separates the common and individual features from multi-block data. Two efficient algorithms termed common orthogonal basis extraction (COBE) are proposed to extract the common basis which is shared by all data, independent on whether the number of common components is given or not. Feature extraction is then performed on the common and the individual subspace separately, by incorporating dimensionality reduction and blind source separation techniques. Extensive experimental results on both synthetic and real-world data show significant advantages of the proposed CIFE method in comparison to the state-of-the-art.

Index Terms—Linked blind source separation, common and individual feature extraction, classification, clustering

I. INTRODUCTION AND MOTIVATION

THE emergence of high-dimensional data structures requires new data analysis tools to be able to deal with the many aspects of this multifaceted problem, from data representation and interpretation to information retrieval. In this context, multi-block data analysis techniques are particularly interesting, as they accommodate multiple measurements of the same phenomenon under various experimentation conditions. For example, human electrophysiological signals in response to a certain stimulus, but from different subjects and trials, can be grouped together and naturally linked as multi-block data. Such data blocks share common information, and at the same time they also allow for individual data features to be kept. Intuitively, this common shared information should help to discover connections between members of a data ensemble and can be used to characterize this data ensemble, while the individual features may help to recognize or identify each individual member of the data ensemble. The identification and separation of such common and individual information in order to employ the features highly relevant to the data analysis task at hand promises to significantly improve data analysis 

For example, shared features among tasks have been exploited to improve the performance of supervised and semi-supervised learning. In this paper, we focus on an unsupervised learning framework for the common and individual features across multi-block data.

Common data analysis techniques that relate two data sets include canonical correlation analysis (CCA) which aims to maximize the correlation between two data sets and the class of partial least squares (PLS) methods that maximize the cross-covariance. CCA has also been generalized to multiple data sets in the context of blind source separation (BSS) and feature extraction. For images, the population value decomposition (PVD) jointly analyzes same-size images, and can be considered as a special case of tensor (Tucker) decompositions, an active research topic in high-order data analysis and exploration. Very recently, group independent component analysis (ICA) and independent vector analysis (IVA) were proposed to capture group variables from multi-block data. Common to these methods is that they account for only correlated features within multi-block data, which limits their practical applications. The joint and individual variation explained (JIVE) is a step in the right direction which simultaneously extracts both joint and individual variations across the members of a heterogeneous data ensemble. However, for the potential of JIVE to be fully exploited several issues need to be further addressed: (i) when common components are relatively weak, JIVE often gives principal components rather than true common components, thus compromising the fidelity of the extraction of true common components; (ii) rigorous quantitative analysis regarding the extracted common components is still missing; (iii) for heterogeneous data, JIVE has been mainly used for revealing gene-miRNA associations, but its potential in general machine learning applications remains unclear; iv) ways to improve data analysis performance by incorporating well-established component analysis tools also need further investigation.

To help resolve these issues, we propose a general framework for common and individual feature extraction (CIFE) for multi-block data, and provide:

1) New efficient algorithms for common orthogonal basis extraction (COBE) from multi-block data. These new...
algorithms guarantee the identification of true common components and can be applied to separate common and individual subspaces of multi-block data;

2) A unifying framework for multi-block data analysis, which deals with heterogeneous data structures by respectively applying suitable well-established matrix factorization methods (such as blind source separation (BSS) [23] and nonnegative matrix factorization (NMF) [19]) to the common and individual subspaces separately rather than to the global space of a data ensemble. This allows for more effective information retrieval and feature extraction for multi-block data.

3) Two generic applications of CIFE—classification and clustering, illustrating the extent to which a simultaneous use of common and individual features can improve the performance in practical applications.

We also provide an in-depth analysis of the links between COBE and related methods such as CCA and principal component analysis (PCA). The analysis shows that the introduced common feature extraction can be interpreted as higher-order correlation analysis, where PCA is performed on the common subspace shared by all the available data rather than on a single global data set, as in ordinary PCA.

The rest of the paper is organized as follows. In Section 2 the COBE method is discussed, including the problem statement, model, algorithms, and its links with CCA and other related methods. In Section 3 a general framework for CIFE is presented. The applications of CIFE in classification and clustering are presented in Section 4. Section 5 provides simulations on both synthetic and real-world data, verifying the proposed methods. Finally, concluding remarks and directions for future work are provided in Section 6.

A conference summary of part of our results has been accepted for publication at ICASSP 2015 [24]. Relative to [24], this journal version includes detailed derivations, new algorithms, and potential applications in machine learning, and further experiments.

II. COMMON ORTHOGONAL BASIS EXTRACTION

Consider a set of matrices \( \mathcal{Y} = \{ \mathbf{Y}_n \in \mathbb{R}^{D \times J_n} : n \in \mathcal{N} \} \), \( \mathcal{N} = \{1, 2, \ldots, N\} \), and the following matrix factorization problem, whereby for each matrix \( \mathbf{Y}_n \in \mathcal{Y} \), we seek

\[
\min_{\mathbf{A}_n, \mathbf{B}_n} \| \mathbf{Y}_n - \mathbf{A}_n \mathbf{B}_n^T \|_F^2, \quad n \in \mathcal{N},
\]

where the \( R_n \) columns of \( \mathbf{A}_n \in \mathbb{R}^{D \times R_n} \) represent the latent variables in \( \mathbf{Y}_n \) (sources, bases, loading, etc), and \( \mathbf{B}_n \in \mathbb{R}^{J_n \times R_n} \) denotes the corresponding coefficient matrix (mixing, encoding, etc). We assume that \( R_n < \min(D, J_n) \), which implies that \( \mathbf{A}_n \mathbf{B}_n^T \) provides a compact/compressed or low-rank representation of \( \mathbf{Y}_n \).

A number of matrix factorization techniques exist to solve (1), including PCA, BSS [23], however, these methods consider each matrix \( \mathbf{Y}_n \) separately. This is often both counter-intuitive and physically restrictive, since the members of the data ensemble \( \mathbf{Y}_n \) are likely to be naturally linked, thus sharing some common components. We therefore propose to perform a simultaneous analysis of the ensemble \( \mathcal{Y} \), in order to obtain the latent variables in the form

\[
\mathbf{A}_n = \begin{bmatrix} \bar{\mathbf{A}} & \bar{\mathbf{A}}_n \end{bmatrix}, \quad n \in \mathcal{N},
\]

where \( \bar{\mathbf{A}} \in \mathbb{R}^{D \times C}, \bar{\mathbf{A}}_n \in \mathbb{R}^{D \times (R_n - C)} \), and \( C \leq \min\{R_n : n \in \mathcal{N}\} \). In other words, we assume that the sub-matrix \( \bar{\mathbf{A}} \) contains the common components shared by all the matrices in \( \mathcal{Y} \) while the sub-matrix \( \bar{\mathbf{A}}_n \) contains the individual information in each \( \mathbf{Y}_n \). This allows us to factorize the data matrices in \( \mathcal{Y} \) in a linked way, so that

\[
\mathbf{Y}_n \approx \mathbf{A}_n \mathbf{B}_n^T = \bar{\mathbf{A}} \bar{\mathbf{B}}_n^T + \bar{\mathbf{A}}_n \bar{\mathbf{B}}_n^T = \bar{\mathbf{Y}}_n + \bar{\mathbf{Y}}_n, \quad n \in \mathcal{N},
\]

where \( \bar{\mathbf{B}}_n \) and \( \bar{\mathbf{B}}_n \) are the partitions of the coefficients \( \mathbf{B}_n \) corresponding to \( \bar{\mathbf{A}} \) and \( \bar{\mathbf{A}}_n \). In this way, each data matrix \( \mathbf{Y}_n \) is represented through a combination of components from the common (shared) subspace \( \bar{\mathbf{Y}}_n = \bar{\mathbf{A}} \bar{\mathbf{B}}_n^T \) and the individual (intrinsic) subspace \( \bar{\mathbf{Y}}_n = \bar{\mathbf{A}}_n \bar{\mathbf{B}}_n^T \), as illustrated in Fig.1.

Our objective is to find the common and individual components \( \bar{\mathbf{A}} \) and \( \bar{\mathbf{A}}_n \) which exhibit some desired properties from a given set of matrices \( \mathbf{Y}_n, n \in \mathcal{N} \), without the knowledge of the mixing coefficients \( \mathbf{B}_n \) and possibly even without knowing the number of common components \( C \). Recall that two special cases of (2) have been extensively studied in the past decades:

- The case \( C = 0 \), where no common components exist in \( \mathcal{Y} \) so that the problem boils down to factorizing each matrix \( \mathbf{Y}_n \in \mathcal{Y} \) separately;
- The case \( C = R_n \) for all \( \mathbf{Y}_n \), which is equivalent to matrix factorization of a large global matrix created by stacking together all matrices \( \mathbf{Y}_n \).

It is important to note that the solutions to (2) are not unique. In fact, for any invertible matrix \( \mathbf{Q} \) of appropriate size, we have \( \bar{\mathbf{Y}}_n = (\bar{\mathbf{A}} \mathbf{Q}) (\mathbf{Q}^{-1} \bar{\mathbf{B}}_n^T) \) which is also a solution to (2).

To reduce the solution space and to simplify the computation we shall consider the following three steps:

1. \( \bar{\mathbf{A}} \):
   - Perform any classical method (e.g., PCA) to find the common components and the corresponding coefficients \( \bar{\mathbf{B}}_n \).

2. \( \bar{\mathbf{A}}_n \):
   - Perform a classical method (e.g., PCA) to find the individual components and the corresponding coefficients \( \bar{\mathbf{B}}_n \).

3. \( \bar{\mathbf{A}} \) and \( \bar{\mathbf{A}}_n \):
   - Perform the same classical method (e.g., PCA) to find both the common and individual components and the corresponding coefficients \( \bar{\mathbf{B}}_n \).
Step 1: Consider the QR-decomposition of $$\tilde{A} = UR$$ such that $$U^TU = I$$, where I is the identity matrix (we also use the symbol $$I_C$$ to denote the C-by-C identity matrix). Upon substituting into (3), we obtain

$$A_nB_n^T = [U \tilde{A}] [RB_n^T \tilde{B}_n^T], \quad n \in \mathcal{N},$$

(a comparison between (3) and (4) makes it possible to assume that in (3) $$\tilde{A}^T\tilde{A} = I$$, without loss of generality.

Step 2: Since our aim is to separate “shared” and “individual” latent component subspaces, we can further assume that $$\tilde{A}^T\tilde{A}_n = 0, n \in \mathcal{N},$$ where 0 is a zero matrix, which implies no interaction between the common and individual subspaces, i.e., perfect separability of the common and individual features. This assumption is reasonable and will not introduce any additional factorization error. To see this, consider

$$\tilde{A}_n = A\tilde{A}^T A_n + (I - A\tilde{A}^T)\tilde{A}_n.$$  

Substituting (5) into (3), we have

$$A_n B_n^T = A\tilde{B}_n^T + \tilde{A}_n \tilde{B}_n^T$$

$$= A\tilde{B}_n^T + [A\tilde{A}^T A_n + (I - A\tilde{A}^T)\tilde{A}_n] \tilde{B}_n^T$$

$$= [A\tilde{B}_n^T + A\tilde{A}^T A_n \tilde{B}_n^T] + [(I - A\tilde{A}^T)\tilde{A}_n] \tilde{B}_n^T$$

Upon comparing (6) and (3) and defining $$\tilde{A}_n \doteq (I - A\tilde{A}^T)\tilde{A}_n$$ and $$\tilde{B}_n \doteq \tilde{B}_n + \tilde{A}_n \tilde{B}_n$$, we arrive at $$\tilde{A}^T\tilde{A}_n = 0$$, verifying that this assumption is not only reasonable, but also does not introduce any factorization error.

Step 3: We consider the truncated singular value decomposition (SVD) of $$\tilde{A}_n = U_n A_n V_n^T$$, where $$U_n^T U_n = I$$, $$V_n^T V_n = I$$, and $$A_n \in \mathbb{R}^{R_n \times C}$$ is invertible. Then $$\tilde{A}^T\tilde{A}_n = 0 \iff \tilde{A}^T U_n = 0$$. Setting $$\tilde{A}_n \doteq U_n$$ and $$\tilde{B}_n \doteq V_n^T \tilde{B}_n$$, we have $$\tilde{A}^T\tilde{A}_n = 0$$ and $$\tilde{A}^T\tilde{B}_n = I$$.

Based on the above three steps for a known of number of common components $$C$$, the general factorization problem in (3) can be reformulated as:

$$\begin{align*}
\min_{A_n, B_n} & \sum_{n \in \mathcal{N}} \|Y_n - A\tilde{B}_n^T - \tilde{A}_n \tilde{B}_n^T\|_F^2, \\
\text{s.t.} & \quad \tilde{A}^T \tilde{A} = I_C, \quad \tilde{A}^T\tilde{A}_n = I_{R_n - C}, \quad \tilde{A}^T\tilde{B}_n = I_n, \quad n \in \mathcal{N},
\end{align*}$$

where $$\|\cdot\|_F$$ is the Frobenius norm of matrices. The JIVE method also solves the model (7), although this is not stated explicitly.

Remark 1: Notice that in (7) both $$A_n$$ and $$B_n$$ are to be optimized, this is different from least squares where $$B_n$$ are fixed.

Since from (2)-(6) the separation of individual and common subspaces does not introduce any additional factorization error, by comparing (1) and (7), the matrix $$A_n = [A \quad \tilde{A}_n]$$ essentially gives the optimal rank-$$R_n$$ approximation of $$Y_n$$ with separated common and individual subspaces. This means that (7) actually consists of two major functions: dimensionality reduction (or low-rank approximation) of each single data matrix $$Y_n$$ and separation of common and individual components of the data ensemble. The dimensionality reduction of $$Y_n$$ depends on each $$Y_n$$ only and its purpose is to capture the variation as much as possible; while the latter requires an integrated analysis of all data matrices with the purpose of extracting the common (or highly correlated) components shared by the data ensemble. The JIVE approach performs these two functions simultaneously by applying the alternating least squares (ALS) to (7) and quantifies the amount of joint variation between data types (1). In other words, components extracted by JIVE are largely dominated by variances rather than correlations. As a result, the JIVE may fail to capture the common components with high correlations, especially when the common components in the data ensemble are relatively weak but are consistently present in all data sets. In contrast, we think it could be more natural to realize these two functions.

Fig. 2: Our two-step method for common and individual feature extraction. The separation of the common and individual subspaces via transforming $$A_n$$ will not introduce any additional factorization error.
separately. Hence, we consider a new two-step method to solve (7), which is also illustrated in Fig.2.

Step 1: Dimensionality reduction: update the matrices \( Y_n \) in (7) by their optimal rank-\( R \) approximation \( A_n B_n^T \) by solving (1) separately for each \( Y_n \). We call the original \( Y_n \) raw data and the reduced version \( Y_n \leftrightarrow A_n B_n^T \) cleaned data; the individual subspace \( \bar{A} \bar{B} \) solving (1) separately for each \( Y_n \).

Step 2: Common and Individual Components Separation: solve (7) using the cleaned data. Therefore, because the procedure (2)-(6) that separates the common and individual subspaces does not introduce any factorization error, in theory, we have

\[
Y_n = A_n B_n^T = \bar{A} \bar{B}_n^T + \bar{A}_n \bar{B}_n^T, \quad \forall n \in N,
\]

where \( Y_n \) is the cleaned data. Obviously, (10) holds if and only if \( \bar{A} \) contain only common components. This property can be further exploited to ensure the extraction of true common components, as detailed in Sections III-A and III-B. Additional important advantages of this two-step method include: (i) the dimensionality reduction is more flexible in the sense that any suitable methods can be applied; (ii) it can be performed in parallel when a large number of data sets are involved; and (iii) it simplifies the subsequent separation of the common and individual components.

In the sequel, we shall assume that \( Y_n \) in (7) are the cleaned data processed by the above Step 1.

A. The COBE Algorithm: The Number of Common Components \( C \) is Unknown

The estimation of the common components \( \bar{A} \) plays a central role in solving (7). In fact, once \( \bar{A} \) has been estimated, from (10), the coefficient matrices \( \bar{B}_n \) can be computed from

\[
\bar{B}_n = \left( Y_n^T - \bar{B}_n \bar{A}_n^T \right) \bar{A} (\bar{A}^T \bar{A})^{-1} Y_n^T \bar{A}, \quad \forall n \in N,
\]

which then allows us to compute the common subspace \( \bar{A} \bar{B}_n^T \) and the individual subspace \( \bar{Y}_n = Y_n - \bar{A} \bar{B}_n^T \), \( \forall n \in N \), respectively.

To estimate \( \bar{A} \) efficiently, from (10) we have

\[
\bar{A} = Y_n B_n^T, \quad A_n^T A_n = I_{R_n}, \quad \forall n \in N,
\]

where \( A_n = \bar{A} \bar{A}_n \), \( B_n = \bar{B}_n \bar{B}_n^T \), and \((\cdot)^T\) denotes the Moore-Penrose matrix pseudo-inverse. In other words, by finding appropriate transformation matrices \( B_n^T \), we can obtain the desired common and individual component subspaces spanned by \( \bar{A} \) and \( \bar{A}_n \), respectively.

Computing the basis vectors of \( \bar{A} \). Let \( Y_n = Q_n H_n \), such that \( Q_n^T Q_n = I \); for each matrix \( Y_n \) this only needs to be computed once by using, e.g., the QR decomposition or a truncated SVD of \( Y_n \). We next define \( Z_n = H_n B_n^T \), so that (12) becomes

\[
\bar{A} \bar{A}_n = Y_n B_n^T, \quad A_n^T A_n = I_{R_n}, \quad \forall n \in N,
\]

and hence for any \( n_1, n_2 \in N \), \( n_1 \neq n_2 \), the following holds

\[
\begin{align*}
Q_{n_1} z_{n_1,k} = & Q_{n_2} z_{n_2,k}, \quad \text{if } k \leq C; \\
Q_{n_1} z_{n_1,k} \neq & Q_{n_2} z_{n_2,k}, \quad \text{if } k > C,
\end{align*}
\]

where \( z_{n,k} \) and \( \bar{a}_k \) are the \( k \)th columns of \( Z_n \) and \( \bar{A} \), respectively. From (14), the first column of \( \bar{A} \), denoted by \( \bar{a}_1 \), can be calculated by solving:

\[
\min_{\bar{a}_1, z_{n,1}} f_1 = \sum_n \| Q_n z_{n,1} - \bar{a}_1 \|_F^2, \quad \text{s.t.} \quad \bar{a}_1^T \bar{a}_1 = 1,
\]

where the objective function \( f_1 \) has to be very small in order to ensure that \( \bar{a}_1 \) is a common basis vector, as governed by (10) and (13). Eq. (15) can be minimized by using alternating least-squares (ALS), whereby by first fixing \( z_{n,1} \) the optimal \( \bar{a}_1 \) is found as

\[
\bar{a}_1 = \sum_n Q_n z_{n,1},
\]

which is then normalized to have a unit norm. Repeating for a fixed \( \bar{a}_1 \) we obtain

\[
z_{n,1} = Q_n^T \bar{a}_1, \quad n \in N,
\]

and so on until convergence. A common column \( \bar{a}_1 \) is considered to be found if \( \min f_1 \leq \epsilon \) for a very small threshold \( \epsilon \geq 0 \); otherwise, no common basis exists in \( \mathcal{Y} \), and we terminate the iterations (16) and (17).

Upon finding a set of common basis vectors \( \bar{a}_1, \bar{a}_2, \ldots, \bar{a}_k \), we need to ensure that repeated common basis vectors are not found when we seek the next common vector \( \bar{a}_{k+1} \). This is achieved by considering the following useful property of \( Z_n \).

Let \( Z_{n,C} = [z_{n,1} \ z_{n,2} \ \cdots \ z_{n,C}] \), then from (13) we have

\[
Z_{n,C}^T Z_{n,C} = Z_{n,C}^T Q_n Z_{n,C} = \bar{A}^T \bar{A} = I.
\]

In other words, \( z_{n,k+1}^T z_{n,k} = 0 \), and \( z_{n,k+1} \) is in the null space of \( Z_{n,k} \), which allows us to update \( Q_n \) as

\[
Q_n^{(k+1)} = Q_n^{(k)} (I - Z_{n,k} Z_{n,k}^T)
\]

\[
= Q_n^{(k)} (I - z_{n,k+1} z_{n,k+1}^T),
\]

where \( Q_n^{(1)} = Q_n \). This then yields \( \bar{a}_{k+1} \) through solving

\[
\min_{\bar{a}_{k+1}, z_{n,k+1}} f_{k+1} = \sum_n \| Q_n^{(k+1)} z_{n,k+1} - \bar{a}_{k+1} \|_F^2, \quad \text{s.t.} \quad \bar{a}_{k+1}^T \bar{a}_{k+1} = 1.
\]

The minimum of \( f_{k+1} \) can be obtained by repeating the procedure in solving (15). We distinguish between the following two cases:

1) For \( \min f_{k+1} \leq \epsilon \), a new common basis vector \( \bar{a}_{k+1} \) is found. We then update \( Q_n^{(k+1)} \) using (19) and then solve (20) to seek the next common basis vector.

2) Otherwise, no common basis vector exists any more and a total of \( C = k \) common orthogonal basis vectors are found as \( \bar{A} = [\bar{a}_1 \ \bar{a}_2 \ \cdots \ \bar{a}_C] \).

We refer to the above procedure for finding sequentially an orthogonal basis of the common space as the common
orthogonal basis extraction (COBE), which is outlined in Algorithm 1.

Remark 2: The parameter \( \epsilon \) controls the degree of similarity (i.e., correlation, in this paper) of the extracted components \( Z_n \), \( Y_n \| \overline{A}_k \| \approx \overline{a}_k \). If \( \epsilon = 0 \), the extracted components are exactly the same and equal to \( \overline{a}_k \), otherwise, approximately identical highly correlated components are extracted (see Section 2.5 for a detailed discussion).

Remark 3: The basis matrix \( \overline{A} \) is not unique as \( \overline{A} \) forms another basis matrix that also spans the common subspace, here \( U \) is an arbitrary orthogonal matrix with proper size.

With the intuition that \( f_k \approx 0 \) if and only if \( k \leq C \) (see [14]), the threshold \( \epsilon \) should be sufficiently small to ensure that the extracted \( \overline{A} \) does contain only common components. In practice, we may specify a very small value of \( \epsilon \) such that the number of common components is overestimated. Then the Second OrDer sTatistic of the Eigenvalues (SORT) method [25], which was proposed to detect the gap between the eigenvalues corresponding to the signal space and those belonging to the noise space, can be applied here to estimate the number of common components. Suppose we have overestimated a total of \( K \) ‘common’ components \( \overline{a}_k, \), \( k = 1, 2, \ldots, K \) with \( R_n \geq K > C \), by using Algorithm 1 and the corresponding errors are \( f_k \) given in (20) or reevaluated as \( f_k \approx \frac{1}{N} \sum_{n=1}^{N} \| Y_n Y_n^T \overline{A}_k - \overline{a}_k \|_{F} \). Then we apply SORTE to detect the gap between the common and individual subspaces, as illustrated in Section V and in the first simulation.

Algorithm 1 The COBE Algorithm

Input: \( Y_n, n \in \mathcal{N}, \epsilon \geq 0 \).
1: Let \( Y_n = Q_n H_n \) such that \( Q_n^T Q_n = I_{R_n} \) for all \( n \in \mathcal{N} \).
2: \( \overline{A} = [\cdot], Q_n^{(1)} = Q_n, \) and \( k = 1 \).
3: while \( f_k \leq \epsilon \) do
4:   while not converged do
5:     \( \overline{a}_k = \sum_n Q_n^{(k)} z_{n,k} / \left\| \sum_n Q_n^{(k)} z_{n,k} \right\|_F \); 
6:     \( z_{n,k} = [Q_n^{(k)}]^T \overline{a}_k, n \in \mathcal{N} \); 
7:   end while
8:   \( f_k = \sum_n \left\| Q_n^{(k)} z_{n,k} - \overline{a}_k \right\|_F^2 \);
9:   \( \overline{A} = [\overline{A} \overline{a}_k] \);
10:  \( k = k + 1 \);
11:  \( Q_n^{(k)} = Q_n^{(k-1)} (I - z_{n,k-1} z_{n,k-1}^T), n \in \mathcal{N} \); 
12: end while
13: return \( \overline{A} = [\overline{a}_1 \overline{a}_2 \cdots \overline{a}_C] \), where \( C = k - 1 \).

Algorithm 2 The COBEC Algorithm

Input: \( C \) and \( Y_n, n \in \mathcal{N} \).
1: Let \( Y_n = Q_n H_n \) such that \( Q_n^T Q_n = I_{R_n} \) for all \( n \).
2: Initialize \( Z_n \) randomly.
3: while not converged do
4:   \( P = \sum_{n \in \mathcal{N}} Q_n Z_n \).
5:   \( \overline{A} = EV^T \), where \([E, \Lambda, V] = tSVD(P, C)\).
6:   \( Z_n \leftarrow Q_n \overline{A} \).
7: end while
8: return \( \overline{A} \).

B. The COBE Algorithm When The Number of Common Components \( C \) is Given

For a given \( C \), following the analysis in Section II-A, the common components can be found by solving:

\[
\min_{Z_n, \overline{A}} \sum_{n=1}^{N} \| Q_n Z_n - \overline{A} \|_F^2, \quad \text{s.t.} \quad \overline{A}^T \overline{A} = I, \tag{22}
\]

through alternating optimization with respect to \( Z_n \) and \( \overline{A} \). In this way, when \( \overline{A} \) is fixed, the optimal \( Z_n \) is computed from

\[
Z_n \leftarrow Q_n^T \overline{A}, n \in \mathcal{N}; \tag{23}
\]

while when \( Z_n, n \in \mathcal{N} \), are fixed, (22) is equivalent to

\[
\max_{\overline{A}} \text{trace}(P^T \overline{A}), \quad \text{s.t.} \quad \overline{A}^T \overline{A} = I, \tag{24}
\]

where \( \text{trace}(\cdot) \) denotes the trace of a matrix and \( P = \sum_{n=1}^{N} Q_n Z_n \).

To solve (24), let \( P = E \Lambda V^T \in \mathbb{R}^{D \times C} \) be the truncated SVD (tSVD) of \( P \), where \( \Lambda = \text{diag}(\lambda_1, \lambda_2, \ldots, \lambda_C) \in \mathbb{R}^{C \times C} \) is a diagonal matrix with \( \lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_C > 0 \). Motivated by the work in [26] (page 601), the optimal solution of (24) becomes

\[
\overline{A} = E \Lambda V^T, \tag{25}
\]

which is easy to see by considering

\[
\text{trace}(P^T \overline{A}) = \text{trace}(V \Lambda E^T \overline{A}) = \text{trace}(\Lambda (E^T \Lambda V)).
\]

In fact, as \( E^T E = I \) and \((\Lambda V)^T (\Lambda V) = I\), we have \( |E^T \Lambda V|_{ii} \leq 1 \), which means that \( \text{trace}(\Lambda (E^T \Lambda V)) \leq \sum_{i=1}^{C} \lambda_i \). Clearly, when \( \overline{A} = E \Lambda V \), then \( \overline{A}^T \overline{A} = I \), \( E^T \Lambda V = I \) and \( \text{trace}(P^T \overline{A}) \) reaches its upper bound \( \sum_{i=1}^{C} \lambda_i \). We refer to this algorithm as the COBEC and its pseudocode is given in Algorithm 2.

C. Pre-processing: Dimensionality Reduction

Like CCA, COBE is meaningless if \( R_n = D \) for all \( n \), since for any \( D \times D \) invertible matrix \( \overline{A} \) there always exist matrices \( \overline{B}_n \) such that \( Y_n = \overline{A} \overline{B}_n^T \), i.e., any \( D \times D \) invertible matrix forms a common basis. For this reason in model [7] the condition \( R_n < D \) is required for all \( Y_n \). This requirement is not too restrictive and has physical justification, as for real-world data of high dimensionality, the latent rank is often significantly lower than the dimensionality.

\footnote{The main difference is that here \( \overline{A} \) is not necessarily square.}
of the observed data. If \( R_n < D \) is not satisfied, we need to perform dimensionality reduction (such as PCA) by, e.g., solving (1) prior to applying COBE, which justifies the necessity of the proposed two-step method. However, this is not the only reason to apply dimensionality reduction—it also helps to reduce the computational complexity, and particularly, to reduce noise, outliers, and artifacts in the original raw data. After dimensionality reduction, we obtain cleaned data \( Y_n \approx A_nB_n^T \) with \( \text{rank}(A_n) = R_n < D \), and can apply COBE to the cleaned data to obtain \( A_nB_n^T = \hat{A}B_n^T + \tilde{A}B_n^T \). Notice that if \( A_nB_n^T \) is interpreted as the PCA of matrix \( Y_n \), COBE simply rotates/transforms the columns of \( A_n \) so that the common subspace and the individual subspace are completely disjoint, as illustrated in Fig.2.

Remark 4: Standard PCA performs dimensionality reduction for i.i.d. Gaussian noise; for sparse distributions we may use robust PCA (RPCA) [27], while the number of latent components \( R_n \) can be estimated using, e.g., SORTIE [25], before applying PCA.

D. Relation to Other Methods

To illustrate the relation between COBE and CCA, recall that for data matrices \( Y_1 \) and \( Y_2 \), CCA finds the vectors \( w_1 \) and \( w_2 \) which maximize the correlation \( \rho = \text{corr}(Y_1w_1, Y_2w_2) \), while COBE extracts only components for which the correlation is higher than a specified threshold. We next shed further light on this relationship by introducing the following bound on the correlations of the variables involved.

Proposition 1: Let \( Y_n \) be row-centered (i.e., with zero mean) random variables. Suppose that \( \| Y_n w_n - \bar{a} \|_F \leq \epsilon < \sqrt{2} - 1 \) with \( \| \bar{a} \|_F = 1 \), \( \forall n \in \mathcal{N} \), then for \( \forall m, n \in \mathcal{N}, \) we have

\[
\text{corr}(Y_m w_m, Y_n w_n) \geq \frac{1 - (2\epsilon + \epsilon^2)}{1 + (2\epsilon + \epsilon^2)} > 0. \tag{26}
\]

Proof: From \( \| \bar{a} \|_F = 1 \) and \( \| Y_n w_n - \bar{a} \|_F \leq \epsilon < 1 \), we have

\[
0 < 1 - \epsilon \leq \| Y_n w_n \|_F \leq 1 + \epsilon, \quad \forall n \in \mathcal{N}. \tag{27}
\]

Moreover, \( \forall m, n \in \mathcal{N} \), the following holds

\[
\| Y_m w_m - Y_n w_n \|_F \leq \| Y_m w_m - \bar{a} \|_F + \| Y_n w_n - \bar{a} \|_F \leq 2\epsilon. \tag{28}
\]

Hence,

\[
2w_m^T Y_m^T Y_n w_n = \| Y_m w_m \|_F^2 + \| Y_n w_n \|_F^2 - \| Y_m w_m - Y_n w_n \|_F^2 \geq 2(1 - \epsilon)^2 - 4\epsilon^2, \tag{29}
\]

and from (27) and (29), we have

\[
\text{corr}(Y_m w_m, Y_n w_n) = \frac{w_m^T Y_m^T Y_n w_n}{\| Y_m w_m \|_F \| Y_n w_n \|_F} \geq \frac{1 - 2\epsilon - \epsilon^2}{1 - 2\epsilon - \epsilon^2} \geq \frac{1 - 2\epsilon - \epsilon^2}{(1 + \epsilon)^2} = 1 - 2\epsilon + \epsilon^2.
\]

In other words, if \( f_i \) in (15) and (20) are upper bounded, this, in turn, leads to lower bounded correlations between the projected variables \( Y_n w_n, n \in \mathcal{N} \). Since \( \text{corr}(Y_1 w_1, Y_2 w_2) \rightarrow 1 \) as \( \epsilon \rightarrow 0 \), COBE can be interpreted as a higher order correlation analysis (HCA) method. Fig.3 illustrates the difference in operation between COBE and CCA for multiple data sets. In this simulation we first generated two matrices \( S_n \in \mathbb{R}^{1000 \times 10}, n = 1, 2 \). The first column of \( S_1 \) was \( s_{1,1}(t) = \sin(0.01t) \), and that of \( S_2 \) was \( s_{2,1}(t) = \text{sign}(s_{1,1}(t)), t = 1, 2, \ldots, 1000 \). Other entries were drawn from independent standard normal distributions. The matrices \( S_n \) were mixed via different coefficient matrices \( M_n \in \mathbb{R}^{10 \times 10}, n = 1, 2 \), whose entries were drawn from independent standard normal distributions such that \( Y_n = S_n M_n^T \). Apparently, \( s_{1,1}(t) \) and \( s_{2,1}(t) \) are highly correlated. Hence, by applying CCA we obtained the first pair of canonical variables \( \hat{s}_{n,1}(t) = Y_n w_n \) with the correlation \( \text{corr}(\hat{s}_{1,1}(t), \hat{s}_{2,1}(t)) = 0.8867 \), where \( w_n \) were the corresponding canonical coefficients, \( n = 1, 2 \). The red line in Fig.3(a) shows the common components \( \bar{a} \) extracted by COBE, together with the blue and green lines corresponding highly correlated components extracted from \( Y_n \), i.e., \( Y_n(Y_n^T \bar{a}) \), \( n = 1, 2 \), which matched very well the canonical variables obtained by CCA (For this reason, we also call \( Y_n(Y_n^T \bar{a}) \) normalized common components). However, COBE will extract only the components with very high correlations, as stated in Proposition 1. From the figure, \( \bar{a} \) can be interpreted as the principal component of the normalized common components, the information that is not provided in CCA.

Further to showing in (7) and (8) that COBE has a close relation with PCA, Fig.4 illustrates the difference between COBE, JIVE and PCA using the same data as in Fig.3. The principal component was computed from a concatenated version of \( Y \), defined in (7). Basically, COBE identifies the principal components \( \hat{A} \) of normalized common components (corresponding to the canonical variables in CCA), whereas PCA seeks the principal components of the global data set \( Y \), and JIVE captures the joint variation. In this example, JIVE gave principal components of sorts, rather than common components, illustrating that the power of components dominates the results obtained by JIVE, see Fig.4. In this sense, COBE is closer to CCA while JIVE is closer to PCA (and PLS); COBE can also be viewed as a regularized version of JIVE. In terms of computational demands, compared with JIVE which in the computation of the common subspace involves frequent SVDs of huge matrices containing all data, COBE (COBE(\( C \)) is more...
as these fake common components can be easily detected by
P. Let requirements of COBE can be reduced in the following way.
D. On the other hand, for a very large
Y mon (or highly correlated) components
interpreted as the principal component of the normalized com-
¯ Y a
common information between two data sets. Here
Fig. 3: (a) Illustration of how COBE was able to capture the
common information between two data sets. Here \( \bar{a} \) can be
interpreted as the principal component of the normalized common
(or highly correlated) components \( Y_n(\bar{Y}^T_n \bar{a}_n), n = 1, 2 \).
(b) The canonical variables (i.e., \( Y_n w_n, \ n = 1, 2 \)) obtained
by using CCA.

efficient in the optimization and more physically intuitive and
flexible in the estimation of number of common components.

E. Scalability For Large-Scale Problems
For large-scale data the indices \( D \) and \( J_n (n \in \mathcal{N}) \) in \ref{a}
are quite large. First, recall that in \ref{15}, \ref{20}, and \ref{22} we
use the dimensionality reduced matrices \( Q \in \mathbb{R}^{D \times D_n} \) with
\( R_n < D \), and hence the value of \( J_n \) is generally not an issue.
On the other hand, for a very large \( D \), the time and memory
requirements of COBE can be reduced in the following way.
Let \( P \in \mathbb{R}^{D_p \times D} \) be a random matrix with \( n_{\max} n \in \mathcal{N} \) \( R_n \) \( D_p \ll D \). Then, from \ref{12} we can first solve the model:
\[
\min_{n \in \mathcal{N}} \sum_{n \in \mathcal{N}} \| Y_n^P W_n - \bar{A}^p \|^2_F \tag{30}
\]
where \( Y_n^P = P Y_n \in \mathbb{R}^{D_p \times J_n} \) is much smaller than \( Y_n \), and
\( \bar{A}^p = P \bar{A} \). After the matrices \( W_n \) have been estimated by
using COBE or COBEC, the corresponding common basis can be
computed from \( \bar{A} = Y_n W_n \). Obviously, \( P Y_n W_n = P \bar{A} \)
as long as \( Y_n W_n = \bar{A} \), in other words, no common basis
vectors are lost. In the worst case, this approach may give
fake common components \( \bar{a}_k \) when \( Y_n w_{n,k} \bar{a}_k \) occasionally
lies in the null space of \( P \). In practice, this is not an issue,
as these fake common components can be easily detected by
examining the value of \( \| Y_n w_{n,k} - \bar{a}_k \|^2_F \).

III. Common And Individual Feature Extraction
(CIFE)
We shall now illuminate the versatility of the COBE ap-
proach over several established feature extraction paradigms.

Fig. 4: Relation between COBE, JIVE, and PCA: COBE
extracted the principal component of normalized common
(or highly correlated) columns whereas PCA gave principal
components of all columns, and JIVE captured the joint
variation and hence failed to extract the highly correlated
components with relatively weak energy.

A. Linked BSS with Pre-whitening
We have so far considered the orthogonal components \( \bar{A} \),
this however does not guarantee unique common components
as the columns of \( \bar{A} U \) also form a common orthogonal basis
for any orthogonal matrix \( U \). If our aim is to project the
common components onto a feature space with some desired
properties (uniqueness), this can be achieved by BSS \cite{[23]},
which finds latent variables \( S \) from their linear mixtures
\( Y = S M^T \) such that
\[
\hat{S} = \Psi(Y) = S P A \tag{31}
\]
where \( \Psi \) denotes a suitable BSS algorithm, \( M \) is the unknown
mixing matrix, while \( P \) and \( A \) are a permutation matrix
and a scaling matrix, respectively, that model the unavoidable
ambiguities of BSS. Assuming that the latent sources \( \bar{F} \) satisfy
\[
\bar{Y}_n = \bar{F} M_n^T \tag{32}
\]
where \( \bar{Y}_n = \bar{A} B_n^T \), we have
\[
\bar{A} = \bar{F} (\bar{B}_n^T \bar{M}_n)^T \tag{33}
\]
illustrating that the columns of \( \bar{A} \) are simply linear mixtures
of the sources \( \bar{F} \), so that \( \bar{F} \) can be estimated via BSS as
\[
\bar{F} = \Psi(\bar{A}) \tag{34}
\]
In this case $\bar{A}$ is a pre-whitened version of $\bar{X}$, which stems from (33) and the fact that $\bar{A}^T \bar{A} = I$. The major advantage of using BSS in this context is that we may obtain common features which exhibit some desired properties such as sparsity, independence, nonnegativity. This is achieved by imposing appropriate constraints and penalties on $\bar{F}$. Moreover, we may extract even more than $C$ common signals from $\bar{A}$ or $\bar{X}_n$, which is a challenging problem referred to as underdetermined BSS. To this end, for example, we can apply the novel tensor based approach proposed in [28]. This method advanced the study of this topic in the sense that it can exactly recover as on a global data space; (ii) the linked BSS is capable of capturing components with various diversities or properties. In other words, within the linked BSS we may apply not only ICA, but also NMF (see Section 3.2) or any other suitable component analysis methods. These two distinguishing properties make our linked BSS more flexible and versatile and allow us to extract more physically meaningful components.

B. Common Nonnegative Features Extraction (CNFE)

For nonnegative latent sources $\bar{F}$, we cannot apply NMF methods on $\bar{A}$ directly. Instead, we need to first extract the common subspace using (11), i.e., $\bar{Y}_n = \bar{A} \bar{A}^T \bar{Y}_n$, and subsequently use the following low-rank approximation based (semi-) nonnegative matrix factorization (NMF) model [30]:

$$\min \sum_n \| \bar{F} \bar{M}_n - \bar{A} \bar{B}_n^T \|_F^2, \quad s.t. \quad \bar{F} \succeq 0. \tag{35}$$

The subsequent use of low-rank NMF (if $\bar{M}_n$ is also nonnegative) or low-rank semiNMF (where $\bar{M}_n$ is real-valued) allows us to extract the common nonnegative components $\bar{F}$. For example, the following iterative multiplicative update rules produce nonnegative components $\bar{F}$ and $\bar{M}$:

$$\bar{F} \leftarrow \bar{F} \odot [\bar{A} \sum_n (\bar{B}_n^T \bar{M}_n)]_+ \odot \bar{F} (\sum_n \bar{M}_n^T \bar{M}_n),$$

$$\bar{M}_n \leftarrow \bar{M}_n \odot [\bar{B}_n (\bar{A}^T \bar{F})]_+ \odot \bar{M}_n (\bar{F}^T \bar{F}), \quad n \in N', \tag{36}$$

where $\odot$ and $\odot$ are element-wise product and division of matrices, see [30] for detailed convergence analysis. Similar to BSS, we may impose additional constraints to extract unique nonnegative components, see [31], [32], [33], [34].

C. Individual Feature Extraction (IFE)

Besides the above common features $\bar{F}$ or $\bar{A}$ extracted using common feature extraction (CFE) methods, each data matrix also has its own individual features contained in the matrix $\bar{Y}_n = \bar{Y}_n - \bar{Y}$; these are often helpful in classification and recognition. Notice that although $\bar{Y}_n$ is of the same size as $\bar{Y}_n$, it is rank deficient since $\text{rank}(\bar{Y}_n) + \text{rank}(\bar{Y}) = R_n$. Hence, dimensionality reduction of $\bar{Y}_n$ should be carefully

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**Fig. 5: General framework of common and individual feature extraction (CIFE), i.e., group component analysis from multi-block data.**
addressed prior to further analysis. To estimate \( \hat{A}_n \) and \( \hat{B}_n \),
we can apply any standard dimensionality reduction method discussed in Section 2.4
(and the associated rank estimation techniques) to each \( Y_n \) separately, followed by BSS or related
methods to extract the features in \( \hat{A}_n \) and \( \hat{B}_n \). However, there is
a major difference between the dimensionality reduction
methods considered here and those in the pre-processing
stage. In the pre-processing stage dimensionality reduction
is rather general-purpose and relatively simpler, whereas here
the dimensionality reduction is related to a specific task at
hand. For example, we may wish to visualize data in a low-
dimensional space [35], or to extract discriminative information,
or to establish neighbor relationship. The above procedure
is referred to as individual feature extraction (IFE), as the
extracted features are only presented in individual datasets.

Fig. 5 shows the concept of the proposed common and
individual feature extraction (CIFE).

IV. EXAMPLE APPLICATIONS

A. Classification Using Common Features

In classification and pattern recognition, training data con-
tain training samples and their labels, while the objects be-
longing to the same category naturally share some common
features. More specifically, for a set of common features
extracted from the \( k \)th category, \( k \in \mathcal{K} = \{1, 2, \ldots, K\} \),
denoted by \( \bar{F}_k \), upon arrival of a new test sample \( y_t \in \mathbb{R}^D \),
we can compute its matching score \( r_t(k) \) with each \( \bar{F}_k \) as:

\[
 r_t(k) = \text{Matching}(y_t, \bar{F}_k), \quad k \in \mathcal{K}.
\]

Since the samples within a certain class should share some
features, the label of \( y_t \) is estimated as

\[
 l_t = \arg \max_{k \in \mathcal{K}} r_t(k).
\]

The matching score \( r_t(k) \) can be defined in many ways,
such as the Euclidean distance or correlation (angle) between
\( y_t \) and the space spanned by \( \bar{F}_k \), which can be solved via least-
squares and CCA, respectively. See Fig. 6 for the diagram.

B. Clustering Using Individual Features

Cluster analysis assigns a set of objects to clusters in
such a way that the objects belonging to the same cluster
are most similar. Unlike classification, clustering employs an
unsupervised learning approach. In the clustering analysis, it
is usual that all the samples have some common features
although they may be from different clusters. For example,
in human face image analysis, every face has common facial
features such as cheek, nose, eyes, and mouth, whose shapes
and locations are similar. Their common features are not
meaningful for clustering as they do not provide any discrimi-
native information. It is therefore logical to first remove these
common/similar features across all the samples and then to
use their individual features to cluster the objects.

Fig. 7 shows that COBE incorporating CNFE is capable
of extracting common faces (features) in the PIE database
details are given in the next section). We empirically set
\( C = 2 \) and used CNFE to extract common nonnegative
components; the common faces in Fig. 7(a) contain some basic
features of human faces. On the other hand, Fig. 7(b) shows the
accentuated individual local features. These individual features
are quite helpful to improve the accuracy of clustering and
recognition. In our individual feature based clustering method
we used the following steps:

1) Randomly split the samples \( y_t \) into \( N \) groups to con-
struct \( Y_n \), where \( t \in \mathcal{T} = \{1, 2, \ldots, T\} \) and \( n \in \mathcal{N} \);
2) Apply COBE to extract the common features \( \tilde{A} \) of
\( \{Y_n, n \in \mathcal{N}\} \);
3) Remove the common features from \( Y_n \) by letting \( \tilde{Y}_n =
Y_n - \tilde{Y}_n = Y_n - \tilde{A}\tilde{A}^T Y_n \);
4) Perform dimensionality reduction and feature extraction
on \( \{\tilde{Y}_1, \tilde{Y}_2, \ldots, \tilde{Y}_N\} \) to obtain the features \( \tilde{F} =
[\tilde{f}_1, \tilde{f}_2, \ldots, \tilde{f}_T] \);
5) Apply clustering algorithms on \( \{\tilde{f}_t, t \in \mathcal{T}\} \), where \( \tilde{f}_t \)
corresponds to the original objects \( y_t \).
V. Simulations and Validation

Linked BSS. In this simulation, we generated a total of ten matrices $S_n \in \mathbb{R}^{5000 \times 10}$, $n = 1, 2, \ldots, 10$, for which the first four columns were speech signals included in the ICALEB benchmark (named Speech4.mat) [36], and the other six components were drawn from independent standard normal distributions. The entries of the mixing matrices $M_n \in \mathbb{R}^{50 \times 10}$ were also drawn from independent standard normal distributions. We used the model $Y_n = S_n M_n^T + E_n$, where $E_n$ contains white Gaussian noise (SNR=20dB), and first employed the COBE, JIVE [1], JBSS [14], and PCA methods to extract the common bases. The distinguishing properties of COBE verified by this simulation example include:

- Finding the number of common components in COBE is physically intuitive and simple;
- The threshold $\epsilon$ within COBE also has physical interpretation—the degree of similarity between the components.

Fig. 8 illustrates the underlying principle of the estimation of the number of components by tracking the parameter $f_i$, while Fig. 9 illustrates the average (over 50 runs) performance in terms of the running time and separation accuracy of COBE with the observations projected onto a lower $D_p$-dimensional space over 50 Monte-Carlo runs.

\[
SIR(s, \hat{s}) = 10 \log_{10} \frac{\sum_i s_i^2}{\sum_i (s_i - \hat{s}_i)^2},
\]

where $s$ and $\hat{s}$ are normalized random variables with zero mean and unit variance, and $\hat{s}$ is an estimate of $s$. Observe that JIVE and COBE achieved higher SIRs than JBSS and PCA, although the performance of JBSS improved after incorporating SOBI. Moreover, although PCA has a close relation with COBE, the common features extracted by PCA are often contaminated by individual features; also while COBE and JIVE achieved almost the same separation accuracy, COBE was much faster (The time for dimensionality reduction has been included for all algorithms). Moreover, after we reduced the power of common components such that $\mathbf{1}^T \hat{A}^T \hat{S} \mathbf{1} = 100$, $n = 1, 2, 3, 4$, COBE obtained similar accuracy whereas JIVE failed, which conforms with the analysis in Section 2.4. Also, the performance of JIVE was sensitive to the correct estimate of the rank of joint/common and individual components. If the ranks of individual components were mis-specified, for instance as 7 (denoted as JIVE* in TABLE I), instead of the actual 6, JIVE took more than 77 seconds to converge. Since the estimation of the number of components in JIVE is quite time consuming and the performance is sensitive to selection of its parameters (for this instance, JIVE took more than two hours to estimate the rank); this limits the practical applications of JIVE for large-scale problems.

The distinguishing properties of COBE verified by this simulation example include:

- COBE is able to identify true common subspace even if the common components are relatively weak;
- Computational complexity of COBE depends only on the natural parameters: the size of the data and the number of common components $C$, making it much more efficient than JIVE;
- Finding the number of common components in COBE is physically intuitive and simple;
- The threshold $\epsilon$ within COBE also has physical interpretation—the degree of similarity between the components.

Finding the number of common components in COBE is physically intuitive and simple;
TABLE I: Performance comparison in linked BSS. The latent signals were estimated by applying the SOBI method to the common components extracted by each algorithm.

| Algorithm | SIR₁ | SIR₂ | SIR₃ | SIR₄ | Runtime (s) |
|-----------|------|------|------|------|-------------|
| COBE      | 21.1 | 24.3 | 27.2 | 24.6 | 0.1         |
| COBEC     | 21.1 | 23.3 | 24.2 | 25.2 | 0.1         |
| JIVE      | 21.2 | 23.8 | 24.2 | 25.0 | 7.4         |
| JIVE⁺     | 21.2 | 23.8 | 24.1 | 24.9 | 77.5        |
| JBSS      | 15.1 | 15.4 | 15.9 | 16.3 | 1.7         |
| PCA       | 15.8 | 17.1 | 17.8 | 19.4 | 0.5         |

---

Fig. 10: Illustration of common nonnegative feature extraction.

The soft and bone tissues (both nonnegative) and the remaining eight components were drawn from independent uniform distributions between 0 and 1 to model the interference. The elements of nonnegative mixing matrices \( M_n \in \mathbb{R}^{20 \times 10}, \) \( n = 1, 2, 3, 4, \) were also drawn from independent uniform distributions between 0 and 1. Then four set of observations were generated by using \( Y_n = SM_n^T. \) The sources in this example were highly correlated and could not be separated by ICA methods, while the presence of random dense noise makes the separation by ordinary NMF on each single set of mixtures difficult. We applied COBE to extract a basis of common components extracted by nLCA-IVM [38].

Owing to dense noise, the identifiability conditions of nLCA-IVM were not satisfied, and nLCA-IVM could not extract the desired source images, while COBE performed nonnegative high correlation analysis well.

Clustering Analysis Using Individual Features. We considered two data sets:

Extended Yale Database B [6] The database contains 16,128 images of 28 human subjects under 9 poses and 64 illumination conditions [39]. A total of 5,820 approximately frontal faces were automatically detected by using the classification and regression tree analysis (CART) and cropped for our clustering analysis.

PIE Database [4] This database is a collection of face images of 68 persons taken under different poses, illumination conditions, and expressions. We used a pre-processed version from [40] which consists of 2,856 full frontal face gray scale images taken at the pose c27.

All images were re-scaled to the size of 32 × 32. We randomly selected \( K \) clusters each time, and repeated the experiment 50 times for each selected \( K. \) In each run, the images were first permuted randomly and then split into \( N = \left\lfloor \frac{T}{50} \right\rfloor \) groups to form multi-block data \( Y_n \) with \( J_n \approx 50, \) \( n = 1, 2, \ldots, N, \) and \( T \) is the number of faces, (each group consisted of face images from unknown different clusters). COBE was used to extract the common features followed by CNFE to obtain nonnegative common features. The number of common components was specified as 2 in all experiments, and the two t-SNE components of their individual parts \( \bar{Y}_n \) were used to cluster the data by using K-means (see [41] for the t-SNE method). As K-means is influenced by initial centers of clusters, we replicated K-means 20 times in each run. Two widely used performance indices: Accuracy (%) and Normalized Mutual Information (NMI) were adopted to evaluate the clustering results, see [40] for detailed definitions of these two metrics. The proposed method was compared with PCA with \( K \) principal components, GNMF [40] (using their recommended settings), and the improved MinMax Cut (MMCut) method [42].

Except for MMCut that returns the cluster indicators directly, all the other methods used the K-means to cluster the features extracted by them with the same configuration. To illustrate that the performance of the proposed method was not completely due to t-SNE, two t-SNE components of their individual parts of common components was specified as 2 in all experiments, and nLCA-IVM could not extract the desired source images, while COBE performed nonnegative high correlation analysis well.
TABLE II: Clustering Performance on Extended Yale B

| k   | PCA  | tSNE | GNMF | MMCut | CIFE | PCA  | tSNE | GNMF | MMCut | CIFE |
|-----|------|------|------|-------|------|------|------|------|-------|------|
| 10  | 44.9 ± 4.5 | 52.8 ± 6.3 | 60.3 ± 6.4 | 53.5 ± 3.4 | **71.2 ± 7.1** | 48.5 ± 4.8 | 51.2 ± 5.8 | 61.0 ± 3.9 | 51.4 ± 2.3 | **67.3 ± 6.1** |
| 15  | 45.6 ± 6.2 | 48.7 ± 4.4 | 60.0 ± 6.4 | 56.0 ± 5.6 | **65.2 ± 6.1** | 44.0 ± 5.8 | 51.1 ± 4.7 | 63.9 ± 5.5 | 55.9 ± 4.8 | **66.9 ± 5.2** |
| 20  | 41.3 ± 2.4 | 44.2 ± 2.5 | 52.4 ± 3.8 | 49.5 ± 4.0 | **62.2 ± 4.6** | 43.5 ± 2.3 | 49.2 ± 2.8 | 59.6 ± 3.3 | 53.1 ± 3.3 | **65.7 ± 3.7** |
| 25  | 40.5 ± 2.3 | 40.2 ± 2.5 | 48.7 ± 2.7 | 46.5 ± 3.7 | **57.8 ± 2.9** | 45.4 ± 2.1 | 46.2 ± 2.4 | 57.4 ± 2.5 | 51.9 ± 3.2 | **63.3 ± 2.3** |
| 28  | 37.3 ± 1.4 | 39.2 ± 1.5 | 47.0 ± 1.7 | 47.7 ± 1.4 | **57.1 ± 1.1** | 8.2 ± 5.8 | 8.8 ± 6.3 | 10.8 ± 7.6 | 10.4 ± 7.4 | **12.0 ± 8.5** |
| Avg. | 42.2 | 45.5 | 34.4 | 51.0 | **63.3** | 42.9 | 49.1 | 60.2 | 53.3 | **65.3** |

TABLE III: Clustering Performance on PIE

| k   | PCA  | tSNE | GNMF | MMCut | CIFE | PCA  | tSNE | GNMF | MMCut | CIFE |
|-----|------|------|------|-------|------|------|------|------|-------|------|
| 10  | 79.9 ± 0.5 | 34.0 ± 0.3 | 94.2 ± 0.8 | 63.7 ± 2.7 | **99.3 ± 0.2** | 82.0 ± 0.3 | 39.8 ± 0.5 | 92.7 ± 0.7 | 71.7 ± 1.4 | **98.9 ± 0.3** |
| 20  | 86.8 ± 1.2 | 29.4 ± 0.7 | 90.5 ± 1.1 | 61.4 ± 0.5 | **100.0 ± 0.0** | 88.3 ± 1.0 | 47.7 ± 0.2 | 92.8 ± 0.4 | 76.5 ± 0.9 | **100.0 ± 0.0** |
| 30  | 74.8 ± 0.0 | 40.8 ± 0.0 | 83.9 ± 0.0 | 60.2 ± 0.0 | **90.6 ± 0.0** | 85.3 ± 0.0 | 60.2 ± 0.0 | 92.4 ± 0.0 | 77.0 ± 0.0 | **96.6 ± 0.0** |
| 40  | 67.8 ± 2.4 | 33.5 ± 0.9 | 74.7 ± 2.3 | 62.7 ± 0.2 | **86.3 ± 0.2** | 83.6 ± 1.2 | 57.0 ± 1.0 | 88.5 ± 0.7 | 78.8 ± 0.2 | **95.5 ± 0.1** |
| 50  | 72.3 ± 0.7 | 36.4 ± 0.8 | 73.7 ± 0.6 | 60.4 ± 0.0 | **83.1 ± 0.2** | 86.6 ± 0.2 | 61.1 ± 0.4 | 87.7 ± 0.3 | 79.1 ± 0.2 | **94.5 ± 0.1** |
| 60  | 76.3 ± 0.4 | 35.0 ± 0.6 | 75.4 ± 0.5 | 58.4 ± 0.3 | **84.3 ± 0.6** | 87.2 ± 0.2 | 62.1 ± 0.3 | 88.4 ± 0.3 | 79.2 ± 0.0 | **94.6 ± 0.1** |
| 68  | 77.8 ± 0.5 | 30.2 ± 0.6 | 71.8 ± 1.1 | 60.3 ± 0.1 | **85.3 ± 0.1** | 86.6 ± 0.2 | 60.6 ± 0.1 | 87.2 ± 0.3 | 80.2 ± 0.1 | **95.0 ± 0.0** |
| Avg. | 76.5 | 34.3 | 80.6 | 61.0 | 89.8 | 85.6 | 55.5 | 90.0 | 77.5 | **96.4** |

Fig. 11: Influence of the number of common components \( C \) on the clustering performance.

![Extended Yale B (K=20), PIE (K=40)](image)

(a) The eight categories of the ETH-80 database

(b) The 10 objects in the forth category

Fig. 12: The ETH-80 database.

the components in \( \mathbf{Y}_n \) are quite small (close to 0, ideally), and \( \mathbf{b}_{n,C} = \mathbf{a}_C \mathbf{Y}_n \), i.e., the projection of \( \mathbf{Y}_n \) on \( \mathbf{a}_C \) becomes very small. In other words, the loss of individual features tends to be very small if \( C \) has been slightly overestimated.

Applications in classification. The classification performance of the proposed method was evaluated by using the ETH-80 Dataset. The ETH-80 dataset consists of a total of 3,280 images grouped in 8 categories containing 10 objects with 41 views per object, spaced equally over the viewing hemisphere [33]. Each category contains 10 different objects. Although the objects belonging to the same category share some common features, they also have their individual features different from the other objects in the same category, which makes this database widely adopted to evaluate classifying methods. See Fig[12] for the 8 categories in ETH-80 and the 10 objects in the forth category. We compared our CIFE based classifier (see Fig[6]) with the KNN classifier included in MATLAB 2010b, the SVM classifier [44], the Eigenfeature Regularization and Extraction method [45] (ERE), and the shrinkage LDA [46] (SLDA). As the ERE method needs to perform eigenvalue decomposition of full covariance matrices, we used the grayscale images with the size of 32 × 32 to avoid ERE running out of memory. For the KNN classifier, nearest (measured by correlation) neighbor was used for classification. For SVM, we used 5-fold cross validation and
the best parameters of $c$ and $\gamma$ were found using grid search following the guidelines provided by the authors. While the SLDA can determine the optimal parameters automatically, the ERE has two tunable parameters: $\mu$ that is used to determine the noise region and the number of features $d$. For fair comparison we set $\mu = 1$, as suggested by the authors. To achieve the best performance $d$ was selected as the number of nonzero eigenvalues. In the CIFE based classifier, we split the training samples belonging to each class into $\max(2, \lfloor T/50 \rfloor)$ subgroups and then used COBEC to extract their common features, where $T$ was the number of training samples in this class. For simplicity, we empirically set $C = \min_n J_n \times 80\%$, although the optimal $C$ and number of subgroups could be estimated by using cross validations. Correlation was adopted as the matching score to classify a new test sample (see (37) and Fig. 6 for details). In each run, we randomly selected a certain percentage of samples as the training data and the remainder as the test data. The mean values and standard derivations of classification accuracy over 20 random runs are plotted in Fig. 13, showing that the CIFE based classifier yielded the best classification accuracy among the classifiers considered.

VI. CONCLUSIONS AND FUTURE WORK

A new scheme for common and individual feature extraction (CIFE) for naturally linked multi-block data has been proposed, together with two new efficient algorithms for the extraction of common orthogonal bases (COBE) according to whether the number of common components is known or not. We have also introduced the concept of linked Blind Source Separation (BSS) of multi-block data in order to perform effective task-dependent feature extraction in common and individual subspaces rather than in a global high dimensional space. The proposed CIFE scheme has been validated on classification and clustering tasks, by exploiting the separated common and individual features. Comprehensive simulations have illustrated the ability of the proposed methods to extract common features existing in multi-block data efficiently and accurately.

In this study we have concentrated on developing a unifying and versatile scheme of common and individual feature extraction. Questions that remain to be investigated in the future include:

1) The number of common features, i.e., $C$, which is controlled by the parameter $\epsilon$, often plays a quite important role in practical applications: it controls the degrees of similarity (correlation, in this paper) of common components extracted from different data sets. However, determining its optimal theoretical value is challenging and needs further investigation;

2) For some practical applications, we need to split the data into subgroups manually in order to discover their common features. Optimal grouping of data is a important factor to achieve better performance;

3) In this paper we considered common features in one dimension. Further extensions of the proposed method to higher-order data will yield general and flexible CIFE tools for tensor data.

4) We provided two example applications of CIFE and corresponding experimental evidences to justify their validity and high performance. However, how to unlock the full potential of CIFE in machine learning deserves further investigation.

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