High Dimensionality Reduction by Matrix Factorization for Systems Pharmacology

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Abstract

The extraction of predictive features from the complex high-dimensional multi-omic data is necessary for decoding and overcoming the therapeutic responses in systems pharmacology. Developing computational methods to reduce high-dimensional space of features in in vitro, in vivo and clinical data is essential to discover the evolution and mechanisms of the drug responses and drug resistance. In this paper, we have utilized the Matrix Factorization (MF) as a modality for high dimensionality reduction in systems pharmacology. In this respect, we have proposed three novel feature selection methods using the mathematical conception of a basis for features. We have applied these techniques as well as three other matrix factorization methods to analyze eight different gene expression datasets to investigate and compare their performance for feature selection. Our results show that these methods are capable of reducing the feature spaces and find predictive features in terms of phenotype determination. The three proposed techniques outperform the other methods used and can extract a 2-gene signature predictive of a Tyrosine Kinase Inhibitor (TKI) treatment response in the Cancer Cell Line Encyclopedia (CCLE).

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**Keywords:** Cancer, Cancer Cell Line Encyclopedia, Dimension Reduction, Feature Selection, Matrix Factorization, Systems Pharmacology

**Key Points:**

- Matrix Factorization (MF) is a useful framework for high dimensionality reduction in systems pharmacology.

- Novel feature selection methods using the incorporation of the mathematical conception of a basis for features into MF increases the performance of feature selection process.

- Feature selection based on the basis-concept in MF can provide predictive gene signatures for therapeutic responses in systems pharmacology.

1 Introduction

One of the challenges in the era of systems medicine is to decode and select personalized features for an individual patient to design personalized therapeutic regimens [1, 2], even if the disease has been modeled analytically [3, 4, 5, 6]. Although systems pharmacology-based mechanistic models which are built based on the known interactions among molecular mechanisms and phenotypes provide a path to solve this problem, but high dimensionality of clinical and molecular characteristics, including multi-omic features, adds more complexity to uncovering the mechanisms causing complex diseases in any model system and patient [7]. Systems-level study of interaction networks constructed from omics data can lead to static models connecting molecular data to phenotypes [8]. Yet, these types of models are still high-dimensional and lack temporal characteristics of disease evolution. A path to design personalized molecular sets assigned to a specific phenotype of a complex disease like cancer is to use phenomenological approaches to reduce the high-dimensional data [9]. The emergence of molecular signatures on this domain requires new methodologies for both model reduction and feature selection in large molecular omic and multi-omic data sets. In systems pharmacology, predicting the therapeutic responses in in vitro and in vivo models and detecting the molecular signatures related to drug resistance help to narrow down the therapeutic regimens for testing in clinical trials [1, 10]. High-dimensional data has noise and redundancy and includes irrelevant information controlled by some unknown parameters required for the model to run. For this reason, analysis of large biological data sets does not always end in robust models and is computationally costly while being prone to over-fitting [11, 12]. In addition, if the modeling process is selected to stand on the distance between molecular features, the designed methods face even more performance degeneration due to adverse impacts of the high-dimensional feature space on the calculation of feature distances. With the emergence of advanced information technology and its increasing applications in electronic communications, a wide variety of modern devices have been designed based on big data whose dimension is exceptionally high [13, 14, 15]. The noise and redundancy, including the irrelevant information, come from a large number of complex parameters in the model, and it results in a limited ability to generalize the learning model and impose inefficient requirements such as more memory usage and longer processing time on the computational devices since the corresponding algorithms suffer enormously from over-fitting [16, 17, 18, 19, 20, 21].

To face the challenge of high dimensionality, two well-known approaches have been proposed. One of these methods is “Feature Extraction” which reduces the dimension by projecting the data given in
a higher-dimensional space, through a proper mapping, into a subspace of this space whose dimension is lower enough than that of the original space [22, 23, 24, 25, 26]. The other technique is “Feature Selection” which reduces the high dimensionality by selecting a smaller number of features compared to the original feature set so that the selected features have the capability to represent the original feature space optimally [27, 28, 29]. It is proved that both techniques, particularly feature selection, substantially contribute to eliminate redundancy and noise, remove the irrelevant parameters and enhance the efficiency of data analysis computations, thus leading to a decrease in the costs associated with time and storage [30, 31]. This way, feature extraction, and feature selection provide considerable improvements in the clustering and classification accuracy, respectively. It should be pointed out that feature selection maintains the original features’ conceptual information by preserving the original features representation more perfectly compared to feature extraction, which constructs new features. For this reason, it is more common to use feature selection than feature extraction to develop novel algorithms for handling dimension reduction tasks in recent years [22, 32].

A broad and significant category of the applications of machine learning techniques discussed can be merged with systems pharmacology methodologies to uncover the quantitative features involved in the evolution and treatment of complex diseases such as cancers [1]. One of the first steps to build systems pharmacology models and platforms is to analyze big datasets such as gene expressions and transcriptomics. Since these datasets, considering phenotypes associated with them, are big and complex datasets, reducing their high dimensionality is a key to be able to use them in systems pharmacology [1]. For instance, to determine the sensitivity and resistance to a specific therapeutic in vitro and in vivo, it is necessary to explore the bio-molecular datasets such as gene expression profiles of hundreds of cancer cell lines [2]. Exploring these datasets to extract features related to the mechanisms causing therapeutic-resistant phenotype requires reducing the high-dimensional space of the molecular features. The complexity of the omic datasets, including gene expressions, originates from two major properties of the biological systems. The first one is intrinsic geometry. It indicates the geometrical properties of these data need to be preserved when the features initially exist in a manifold of high-dimension mapped into a lower-dimension sub-manifold of the base space. The other characteristic of such big data is the noise and enormous amount of outliers.

Over the past decades, subspace learning techniques have demonstrated a considerable ability to deal with feature selection problems and their applications to the analysis of complex biological systems [12, 33, 34, 35, 36]. It is required that the subspace learning techniques and feature selection methods, particularly when they are employed to handle gene expression data, are equipped with some proper and efficient tools. In this regard, some modifications have been incorporated into these methods. One good example is subspace learning for unsupervised feature selection via matrix factorization (MFFS), introduced in [37], in which a subspace learning framework is designed from the perspective of the matrix decomposition. The MFFS framework and the idea of minimum redundancy were merged to adapt unsupervised feature selection via maximum projection and minimum redundancy (MPMR) [38]. Another interesting example is subspace learning-based graph regularized feature selection (SGFS), investigated in [39], which can maintain the features intrinsic information in terms of features geometry. Then utilizing matrix factorization techniques again, an unsupervised method, feature selection based on maximum projection and minimum redundancy (MPMR), was explored. All these algorithms can provide a highly accurate estimate of the feature space by learning a proper subspace and also produce satisfactory outcomes for the corresponding feature selection task.

Although the mentioned methods provide powerful tools to successfully develop new practical models for dimension reduction problems, they have not been vastly applied to systems pharmacology.
The major goal of this paper is to establish a number of novel feature selection frameworks and apply them to cancer gene expression data. In the proposed methods, not only are the notable and outstanding achievements of the MFFS, MPMR and SGFS methods considered, but also the performance of the introduced methods is improved in terms of classification and clustering. For this, the notion of a basis, which is an essential concept in the theory of linear vector spaces, is applied. Normally, a basis possesses two fundamental characteristics, each of which has a positive and constructive impact on the discriminative feature selection process [35]. The first property is that a basis generates the whole feature space making use of the basis elements. This means that the characteristics of original features are included in the basis whose number of elements is far smaller than that of the whole feature space. A major advantage in this case is that rather than considering the original features, a basis can be utilized to represent them. It leads to a considerable reduction in the number of the selected features and the calculating processes. The second property of a basis is its elements that are linearly independent, which indicates a significant decrease in the data redundancy.

Figure 1 illustrates a schematic diagram of the major set of concepts used in the proposed methods. This figure shows that considering an original feature data, a basis of features is primarily built for the original feature space. Accordingly, the feature selection problem is defined based on the constructed basis. Through some regularizers such as the $L_{2,1}$-norm sparsity regularization, the redundancy of the basis space and the manifold of the basis space, three feature selection methods are introduced under these regularizations, and the feature matrix $G$ is learned to enhance the feature learning performance.

Figure 1: A schematic diagram of the fundamental framework of the proposed methods.

2 Motivation and Background

This section discusses the substantial role of a basis of features for the feature selection problems and describes an algorithm to construct a basis for the features space. Throughout the paper, the lowercase letters in Italic are used to denote the scalars, the uppercase letters in boldface represent matrices, the lowercase letters in boldface indicate vectors, and the uppercase letters in italic are used for sets. For a matrix $A$, the set including all linear combinations of its columns is denoted by $\text{span}(A)$. The notation $E_k$ represents the $k \times k$ identity matrix. For a given matrix $A$, $A^T$ and $\text{Tr}(A)$ denote the transpose and the trace operators, respectively. In addition, $\|A\|_F$ is the Frobenius norm of the matrix $A$. For a matrix $A \in \mathbb{R}^{m \times n}$, the $L_{2,1}$-norm of $A$ is given by $\|A\|_{2,1} = \sum_{i=1}^{m} \sqrt{\sum_{j=1}^{n} A_{ij}^2}$. 

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The data matrix is represented by \( X = [x_1; x_2; \ldots; x_n] = [f_1, f_2, \ldots, f_d] \in \mathbb{R}^{n \times d} \), where \( n \) and \( d \) indicate the number of samples and that of features, respectively. It is also supposed that \( n \leq d \). For a feature subset \( X_I \), the corresponding submatrix of \( X \) is simply represented by \( X_I \) that is a matrix of order \( n \times k \) in which \( I \) and \( |I| \) show the index set of the selected features and the number of indices of \( I \), respectively. Given a vector space \( V \) of dimension \( n \), a basis for \( V \) is a subset \( B = \{v_1, v_2, \ldots, v_n\} \) of \( n \) linearly independent elements of \( V \) such that \( V \) is spanned by the vectors of \( B \). Given the matrix of features \( X = [f_1, f_2, \ldots, f_d] \), it is supposed throughout this article that the set \( B \) is composed of \( n \) elements and is a basis of the vector space spanned by \( X \) as well as assuming that \( \text{rank}(X) = n \). The rank of a matrix \( X \), \( \text{rank}(X) \), is the dimension of the vector space produced by the rows or the columns of \( X \). It can also be proven that \( \text{rank}(X) \) is equal to the maximum number of rows or columns of \( X \) that are linearly independent.

2.1 Motivation

The notion of directional distance between two subspaces has been discussed in learning problems regarding subspaces and several feature selection methods have been established based on it.

**Definition 1 (Directional Distance [37]).** Given two matrices \( S_1 \in \mathbb{R}^{m \times n} \) and \( S_2 \in \mathbb{R}^{m \times p} \), the distance from \( S_1 = \text{span}(S_1) \) to \( S_2 = \text{span}(S_2) \) is calculated by:

\[
\vec{d}(S_1, S_2) = \min_{H \in \mathbb{R}^{p \times n}} \| S_1 - S_2 H \|_F^2.
\] (1)

One most well-known and practical technique for feature selection is presented in Problem 1 that is rested on the directional distance defined above and has been the center of attention as the foundation of a number of powerful state-of-the-art algorithms in recent years [34, 35, 36, 37, 38, 39].

**Problem 1.** Let \( X \in \mathbb{R}^{n \times d} \) and \( k \in \mathbb{Z}^+ \). Considering \( I \) as a feature index set for which \( |I| = k \), a feature selection problem is formulated by Eq. (2).

\[
\min_{|I|=k} \vec{d}(\mathcal{X}, \mathcal{X}_I) = \min_{H \in \mathbb{R}^{k \times d}} \| X - X_I H \|_F^2,
\] (2)

where \( \mathcal{X} = \text{span}(X) \) and \( \mathcal{X}_I = \text{span}(X_I) \).

Saberi-Movahed et al. [35] studied the importance of the representation of features provided by the notion of a basis of \( \text{span}(X) \) in terms of the improvements that this representation makes on the feature selection performance in Problem 1. Specifically, two highly significant properties were investigated as follows.

1. Let \( B \) be a basis of \( \text{span}([f_1, f_2, \ldots, f_d]) \) and \( B = \text{span}(B) \). Then

\[
\vec{d}(\mathcal{X}, B) = 0.
\] (3)

2. Let \( X_I \subseteq X \) be a feature subset. Then

\[
\vec{d}(B, \mathcal{X}_I) \leq \vec{d}(\mathcal{X}, \mathcal{X}_I).
\] (4)
Item (1) states that for a basis $B$ of a feature subspace $\mathcal{X}$, its span, $\mathcal{B}$, can be selected as a representation of $\mathcal{X}$ since $\mathcal{B}$ coincides with $\mathcal{X}$. Furthermore, Item (2) indicates that the representation of $\mathcal{X}$ with $\mathcal{B}$, i.e., assuming that $\mathcal{X} = \mathcal{B}$, encourages us to replace $\overrightarrow{d}(\mathcal{X}, \mathcal{X}_I)$ in Problem 1 by $\overrightarrow{d}(\mathcal{B}, \mathcal{X}_I)$ and it guarantees that this replacement will not make the results of data fitting worse than that of the original problem [35].

Considering the merits of a basis in feature selection problems, Saberi-Movahed et al. [35] proposed a new objective function, given by Eq. (5), in which the original feature space was represented by a basis of this space. Problem 2 summarizes this method which improved the method presented by Problem 1.

**Problem 2.** Let $X$ be the original set of features and $B$ be a basis of the span of $X$. In addition, $B = \text{span}(B)$ and $\mathcal{X}_I = \text{span}(X_I)$. Assuming that $I$ is a set composed of $k$ indices of the features, an objective function can be formulated as:

$$\min_{|I|=k} \overrightarrow{d}(B, \mathcal{X}_I) = \min_{H \in \mathbb{R}^{|I| \times n}} \|B - X_I H\|_{F}^2.$$  \hspace{1cm} (5)

Recently, a supervised feature selection technique, SFS-BMF, rested on Problem 2, was presented in [35] so that a basis of features was applied to the selection process. The advantage of this method over a set of benchmark techniques was demonstrated by conducting diverse experiments on a number of gene expression datasets. Inspired by satisfactory outcomes from supervised feature selection founded on the notion of a basis, we investigate the impacts of this approach on the outcomes of feature selection problems in the unsupervised mode.

The main goal of the feature selection is to choose a feature subset to decrease the noise and to enhance the efficiency and efficacy of a learning task, such as classification, prediction and clustering [40, 41]. To diminish the dimension of feature space, the amount of the lost information needs to be as low as reasonable so that the obtained space can supply sufficient information for the learning task. In fact, it is essential to pick out features that are relevant for a learning task, but at the same time it is important to have a set of features which is not redundant for the purpose of enhancing the robustness [28]. In this regard, one of the most important issues in the process of feature selection is the redundancy. Redundancy evaluates how similar the features are to each other and how much inserting a new feature into a specific feature set plays an effective role in the data analysis [42].

In the context of linear algebra, there is a direct relationship between the redundancy and the notion of linear independence [43]. Specifically, a set of vectors $\{x_1, x_2, \ldots, x_d\}$ in $\mathbb{R}^n$ is called linearly dependent if there exist non-zero scalars $\alpha_1, \alpha_2, \ldots, \alpha_d$ in $\mathbb{R}$ such that $\alpha_1 x_1 + \alpha_2 x_2 + \cdots + \alpha_d x_d = 0$. This means that one of these vectors is a linear combination of some of the others. On the other hand, it is said that the vectors are linearly independent if they are not linearly dependent, that is, the expression $\alpha_1 x_1 + \alpha_2 x_2 + \cdots + \alpha_d x_d = 0$ implies that $\alpha_i = 0$ for $i = 1, \ldots, d$. For an example, let us consider the vectors

$$x_1 = \begin{bmatrix} 1 \\ 1 \end{bmatrix}, \quad x_2 = \begin{bmatrix} 2 \\ 3 \end{bmatrix}, \quad x_3 = \begin{bmatrix} 3 \\ 4 \end{bmatrix}.$$  

It can be easily checked out that $x_1 + x_2 - x_3 = 0$. It is seen that $x_3 = x_1 + x_2$, and the set of vectors $\{x_1, x_2, x_3\}$ in $\mathbb{R}^2$ is linearly dependent. As an immediate result, we can infer from this example that a linearly dependent set of vectors has a redundancy so that one of the vectors can be took away from the original vectors without affecting the span of the other vectors. Considering what is described above, a significant advantage of removing linearly dependent features from the original set of features is that it helps to achieve one of the main goals of feature selection.
that is minimizing the redundancy. On the other hand, every basis for the feature space possesses two major characteristics. Firstly, it spans the feature space, and secondly, all the features included in a basis are linearly independent. Such a basis reduces the feature space so that the redundancy among the features decreases. It is because the reduced features are linearly independent as elements of the basis. Based on this, the role that a basis for the feature space plays in learning tasks is investigated in more depth in this paper and this investigation involves the theoretical foundation as well as some practical aspects.

2.2 An Algorithm to Build a Basis for the Features Space

In this section, we aim to describe how to find a proper basis for the feature space. The technique of Variance Score (VS) is an effective method that determines a discriminative feature as the one whose VS is a large value. This technique is very simple to implement and is commonly used for feature selection problems in unsupervised mode. It is known that the variance of data represents the level of its spread over a given domain. The variance maximization for the data that are reduced and transformed into a lower dimension leads to far more information extraction. Considering the remarkable ability of this method, our proposed techniques apply it to determine which features are the most discriminative to be included in the basis we are going to form. Such a basis is constructed as follows. First, computing every feature’s VS, the features are reordered in terms of their VS so that the feature with the highest VS comes first, and the other features are ordered in a designing manner after it. Next, the first feature, the one whose VS is the highest value, is considered as the first feature of the basis and the set including this single feature is extended to a basis of features by applying the basis extension algorithm to the other features ordered after it. Algorithm 1 provides a detailed description of this process.

Algorithm 1 Building a basis for the feature space.

Input: Data matrix $X = [f_1, f_2, \ldots, f_d] \in \mathbb{R}^{n \times d}$ in which $1 < \text{rank}(X) = p \leq \min\{n, d\}$.

Output: A basis $B$ for the vector space spanned by the set $\{f_1, f_2, \ldots, f_d\}$ of the features.

1: for $i = 1$ to $d$ do
2: $\text{VS}(i) \leftarrow$ Variance Score (VS) for the $i$th feature.
3: end for
4: Arrange the VSs in a descending order.
5: $X^{\text{new}} \leftarrow$ Rearranging the features in $X$ in terms of the order of their corresponding VS. Set $X^{\text{new}} = [f_1^{\text{new}}, f_2^{\text{new}}, \ldots, f_d^{\text{new}}]$.
6: Set $B = \{f_1^{\text{new}}\}$.
7: for $k = 2 : d$ do // The basis extension technique
8: if $f_k^{\text{new}} \notin \text{span}(B)$ then
9: $B \leftarrow B \cup \{f_k^{\text{new}}\}$.
10: end if
11: if $|B| = p$ then
12: Stop.
13: end if
14: end for

In Algorithm 1, the basis extension technique has been utilized to construct a basis for the feature space. In linear algebra, this technique is a standard commonly used way to produce a basis of a given
vector space when the dimension of the space is finite [44]. For an \( n \times d \) matrix \( V = [v_1, \ldots, v_d] \), where \( 1 < \text{rank}(V) = p \leq \min\{n, d\} \), it is assumed that \( V_1 = \{v_{i_1}\} \) which \( v_{i_1} \in \{v_1, \ldots, v_d\} \) is a non-zero vector of size \( n \). Therefore, the subset \( V_1 \) is linearly independent. Now considering \( 1 < p \), a vector \( v_{i_2} \in \{v_1, \ldots, x_d\} \) - \( \text{span}(V_1) \) can be found. It is evident that the extension of \( V_1 \) defined as \( V_2 = V_1 \cup \{v_{i_2}\} = \{v_{i_1}, v_{i_2}\} \) is a linearly independent subset. Now, continuing this process, a series \( V_3, V_4, \ldots, V_p \) of subsets that are linearly independent extensions of \( V_1 \) is obtained. Considering the fact that \( V_p \) is composed of \( p \) vectors, \( V_p \) is a basis of \( \mathcal{V} = \text{span}\{v_1, \ldots, v_d\} \). Algorithm 2 summarizes the procedure of the basis extension technique to construct a basis for \( \text{span}(V) \).

**Algorithm 2** The basis extension technique [44].

**Input:** Data matrix \( V = [v_1, v_2, \ldots, v_d] \in \mathbb{R}^{n \times d} \) such that \( 1 < \text{rank}(V) = p \leq \min\{n, d\} \).

**Output:** A basis \( B \) for \( \mathcal{V} = \text{span}(V) \).

1. Choose an element of \( \{v_1, \ldots, v_d\} \) such as \( \{v_{i_1}\} \). Set \( B = \{v_{i_1}\} \).
2. for \( k = 2 \) : \( n \) do
3. Find a vector \( v_{i_k} \in V \) in a way that \( v_{i_k} \notin \text{span}(B) \). Set \( B \leftarrow B \cup \{v_{i_k}\} \).
4. end for

3 Proposed Frameworks

This section presents a set of novel and effective feature selection techniques rested on Problem (2) and the notion of a basis. In Section 3.1, an original unsupervised feature selection method is offered based on a basis of the feature space. In addition, Section 3.2 proposes a new unsupervised feature selection technique in which the redundancy among the features is minimized. Finally, Section 3.3 introduces another method for feature selection that preserves the intrinsic geometry of features.

3.1 The B-MFFS Method

A set of methods have been founded on the concept of matrix factorization, including Singular Value Decomposition (SVD) [45], Non-negative Matrix Factorization (NMF) [46] and Principal Components Analysis (PCA) [47]. Since such techniques were originally developed to handle feature extraction tasks, they are not always performed well enough when applied to feature selection problems. Especially, interpretation of the features determined by the mentioned methods in a subspace of lower dimension is not acceptable. One technique to deal with this problem was “Matrix Factorization criterion of Feature selection Subspace learning” (MFFS), proposed by Wang et al. [37]. In this approach, the matrix factorization was used for feature selection. This method was constructed using the problem presented in Eq. (6) and determines the optimal subset of features subject to an orthogonality condition imposed on its optimization problem.

\[
\arg\min_{W \in \mathbb{R}^{d \times k}, H \in \mathbb{R}^{k \times d}} \|X - WH\|_F \quad \text{subject to} \quad W^TW = E_k, W \geq 0, H \geq 0. \tag{6}
\]

The current section aims to present a novel method, B-MFFS, for feature selection problems that is rested on the mathematical notion of a basis. Particularly, the B-MFFS problem is established based on the idea of selecting a basis for the feature space as a representation of this space. This means that first, a basis is determined for the space of the original features. Then, to design an efficient and effective
mechanism for feature selection, the B-MFFS algorithm represents the original feature space by the span of this basis and minimizes the distance between this span and the set of the selected features. In addition, making use of the $L_2,1$--norm, the matrix of feature weights is regulated to guarantee the sparsity of feature selection [48].

Let $X = \{f_1, f_2, \ldots, f_d\}$ be the set of the original features and $B = [f^b_1, f^b_2, \ldots, f^b_n] \in \mathbb{R}^{n \times n}$ be a basis for the span of $X$. Since for a given submatrix $X_I$ of $X = [f_1, f_2, \ldots, f_d]$, there is a unique representation of $X_I$ in the form of a linear combination of elements of $B$, a decomposition of $X_I$ exists as follows: $X_I = BG$, where $G \in \mathbb{R}^{n \times k}$. Considering the aforementioned reduced forms for $X$ and $X_I$, the feature selection problem for B-MFFS is formulated by Eq. (7).

$$\arg\min_{G \in \mathbb{R}^{n \times k}, H \in \mathbb{R}^{k \times n}} \|B - BGH\|^2_F + \alpha \|G\|_{2,1}, \tag{7}$$

where $\alpha$ is a regularization parameter, and the $L_2,1$--norm is imposed on the weight feature matrix $G$ to guarantee that this matrix is sparse. Moreover, the term $\|G\|_{2,1}$ can be rewritten as:

$$\|G\|_{2,1} = \text{Tr}(G^TUG),$$

where $U \in \mathbb{R}^{n \times n}$ is a diagonal matrix with the diagonal elements

$$U_{ii} = \frac{1}{2\|G_i\|_2}, \quad \text{for} \quad i = 1, \ldots, n.$$

In order to find $G$ and $H$, we make use of an alternating least squares iterative method in such a way that we will solve (7) with respect to one variable while the other one is taken to be fixed. For this purpose, we first assume that $H \in \mathbb{R}^{k \times n}$ is fixed. In this respect, it is supposed that

$$\mathcal{L}(G, H) = \|B - BGH\|^2_F + \alpha \text{Tr}(G^TUG).$$

Now, by taking the partial derivative of $\mathcal{L}$ with respect to $G$, we have

$$\frac{\partial \mathcal{L}}{\partial G} = 2B^TBGHH^T - 2B^TBH^T + 2\alpha UG.$$

Set $\frac{\partial \mathcal{L}}{\partial G} = 0$. It follows that the matrix $G$ satisfies Eq. (8).

$$B^TBGHH^T + \alpha UG = B^TBH^T. \tag{8}$$

In the same manner, taking the partial derivative of $\mathcal{L}$ with respect to $H$ when $G \in \mathbb{R}^{n \times k}$ is assumed to be fixed, we get

$$\frac{\partial \mathcal{L}}{\partial H} = 2G^TB^TBG - 2G^TB^TB.$$

Again, putting $\frac{\partial \mathcal{L}}{\partial H} = 0$, it is obtained that $H$ satisfies Eq. (9).

$$G^TB^TBG = G^TB^TB. \tag{9}$$

Algorithm 3 summarizes what has been discussed in this section.
Algorithm 3 The B-MFFS method.

**Input:** Data matrix $X \in \mathbb{R}^{n \times d}$; the regularization parameter $\alpha$; the number of the selected features $k$; maximum number of iterations $\text{maxIter}$.

1: Initialize the basis of features $\{f_1^b, f_2^b, \ldots, f_n^b\}$ according to the technique of VS described in Algorithm 1 and consider the matrix $B = [f_1^b, f_2^b, \ldots, f_n^b]$.

2: Initialize $H \in \mathbb{R}^{k \times n}$.

3: while iteration $\leq \text{maxIter}$ do


5: Fix $G$ and solve $G^T B^T B G = G^T B^T B$ for $H$.

6: end while

**Output:** Sort the norm of rows of the weight feature matrix, $\|G_i\|_2$, in a descending order and put the rows of $G$ in an order induced by the values of $\|G_i\|_2$, where $i = 1, \ldots, n$. Next, from $\{f_1^b, f_2^b, \ldots, f_n^b\}$, select $k$ features whose corresponding rows in $G$ are evaluated as the top score rows according to the norm function.

### 3.2 The B-MPMR Method

There are two major drawbacks when data of high-dimension are applied to machine learning methods. These include the significant increase in the storage costs and level of redundancy which negatively affect the learning process performance. To address these problems, a set of feature selection techniques have been developed. An example of such methods, that was successful in representing the original features effectively and reducing the features redundancy, is “feature selection based on Maximum Projection and Minimum Redundancy” (MPMR) [38], that can be formed as:

$$
\arg\min_{W \in \mathbb{R}^{d \times k}, H \in \mathbb{R}^{k \times d}} \|X - X W H\|_F^2 + \alpha \Tr(W^T X^T X W 1_k), \quad \text{subject to } W \geq 0, W^T W = E_k, \quad (10)
$$

in which $\alpha$ is considered to achieve a proper trade-off between the redundancy rate and the approximation degree.

The current section introduces B-MPMR as an improvement of MPMR. The key idea in B-MPMR is to investigate that to what extent the features in a subset of the basis are independent of each other. Suppose that the basis $B = [f_1^b, f_2^b, \ldots, f_n^b] \in \mathbb{R}^{n \times n}$ is as the one mentioned in Section 3.1. The B-MPMR technique addresses the redundancy problem by computing the rate of redundancy for each feature in the given feature subset and using the results to guide the process of selecting the features.

Given a submatrix of features, $X_I$, there is a representation $X_I = B G$ similar to the one described in Section 3.1. For $X_I$, the rate of redundancy, $\text{red}(I)$, is calculated by Eq. (11).

$$
\text{red}(I) = \frac{1}{k^2} \sum_{p=1}^{k} \sum_{q=1}^{k} x_p^T x_q = \frac{1}{k^2} \Tr(X_I^T X_I 1_k) = \frac{1}{k^2} \Tr(G^T B^T B G 1_k), \quad (11)
$$

in which the matrix $1_k \in \mathbb{R}^{k \times k}$ is defined so that all its entries are one. In this regard, we can see that

$$
\Tr(G^T B^T B G 1_k) = 1_k^T G^T B^T B G 1_k, \quad (12)
$$

in which the vector $1_k \in \mathbb{R}^k$ whose entries are all one. Considering the fact that $G^T B^T B G$ is a symmetric positive semi-definite matrix, it is deduced from Eq. (12) that $\Tr(G^T B^T B G 1_k) \geq 0$.  

10
Regarding the redundancy problem discussed above, the feature selection problem of the B-MPMR
method is formulated as follows:

$$\arg\min_{G \in \mathbb{R}^{n \times k}, H \in \mathbb{R}^{k \times n}} \| B - BHG \|_F^2 + \alpha \text{Tr}(G^T B^T BG_1) + \beta \| G \|_{2,1}, \quad (13)$$

where $\alpha$ and $\beta$ are two regularization parameters, and the $L_{2,1}$-norm is imposed on the weight feature
matrix $G$ to assure the sparsity of this matrix.

To find $G$ and $H$, the alternating least squares iterative method is employed again to solve (13)
with respect to one variable while the other one is fixed. First, assume that $H \in \mathbb{R}^{k \times n}$ is fixed. We
define

$$\mathcal{L}(G, H) = \| B - BGH \|_F^2 + \alpha \text{Tr}(G^T B^T BG_1) + \beta \text{Tr}(G^T U G),$$

where $U \in \mathbb{R}^{n \times n}$ is a diagonal matrix with the diagonal elements $U_{ii} = \frac{1}{2 \| G_i \|_2}$ for $i = 1, \ldots, n$.

Now, by taking the partial derivative of $\mathcal{L}$ with respect to $G$, we have

$$\frac{\partial \mathcal{L}}{\partial G} = 2B^T BGHH^T - 2B^T BH^T + 2\alpha B^T B G_1 + 2\beta U G.$$

Next, assuming that $\frac{\partial \mathcal{L}}{\partial G} = 0$, we get

$$B^T BGHH^T + \alpha B^T B G_1 + \beta U G = B^T BH^T,$$

which can be expressed as

$$B^T BG(HH^T + \alpha I_k) + \beta U G = B^T BH^T. \quad (14)$$

In the same manner, assuming that $G \in \mathbb{R}^{n \times k}$ is fixed and taking the partial derivative of $\mathcal{L}$ with
respect to $H$, we have

$$\frac{\partial \mathcal{L}}{\partial H} = 2G^T B^T BGH - 2G^T B^T B.$$

Setting $\frac{\partial \mathcal{L}}{\partial H} = 0$, it follows that the matrix $H$ satisfies Eq. (15).

$$G^T B^T BGH = G^T B^T B. \quad (15)$$

Algorithm 4 describes the steps of B-MRPR in brief.

### 3.3 The B-SGFS Method

The majority of former approaches consider the original space of features as the domain for the feature
selection process. They also assume that there is a smooth manifold on which the original data are distributed [49, 50, 51]. For instance, the Subspace learning-based Graph regularized Feature Selection
(SGFS) method [39] has been recently developed by including the notion of graph regularization in the
objective function of a former feature selection method. This technique is formulated as:

$$\arg\min_{W \in \mathbb{R}^{d \times k}, H \in \mathbb{R}^{k \times d}} \alpha \| X - XWH \|_F^2 + \text{Tr}(HLH^T) + \beta \| W \|_{2,1}, \quad \text{subject to } W^T W = E_k, W \geq 0, H \geq 0, \quad (16)$$
Algorithm 4 The B-MPMR method.

**Input:** Data matrix $X \in \mathbb{R}^{n \times d}$; the regularization parameters $\alpha$ and $\beta$; the number of the selected features $k$; the maximum number of iterations $maxIter$.

1. Initialize the basis of features $\{f^b_1, f^b_2, \ldots, f^b_n\}$ according to the technique of VS described in Algorithm 1 and consider the matrix $B = [f^b_1, f^b_2, \ldots, f^b_n]$.
2. Initialize $H \in \mathbb{R}^{k \times n}$.
3. **while** iteration $\leq maxIter$ **do**
   4. Fix $H$ and solve the matrix equation $B^T B G (H H^T + \alpha 1_k) + \beta U G = B^T B H^T$ for $G$.
   5. Fix $G$ and solve the matrix equation $G^T B^T B G H = G^T B^T B$ for $H$.
4. **end while**

**Output:** Sort the norm of rows of the weight feature matrix, $\|G_i\|_2$, in a descending order and put the rows of $G$ in an order induced by the values of $\|G_i\|_2$, where $i = 1, \ldots, n$. Next, from $\{f^b_1, f^b_2, \ldots, f^b_n\}$, select $k$ features whose corresponding rows in $G$ are evaluated as the top score rows according to the norm function.

In which $\alpha$ and $\beta$ are nonnegative balance parameters, and $L \in \mathbb{R}^{d \times d}$ is the feature Laplacian matrix. In Eq. (16), the term $Tr(HLH^T)$ handles the geometric structure preservation task for the feature manifold. Moreover, to guarantee that the matrix of feature selection is sparse, this matrix is constrained by the $L_{2,1}$-norm in the objective function of SGFS.

In this section, the B-SGFS method is presented as a technique to select the most discriminative features from a basis of the original feature space. This methodology also preserve the geometric structure of features associated with the basis manifold. This method is different from the SGFS method in terms of two properties including geometric structure preservation and feature manifold reduction. The main idea of our proposed B-SGFS method is to reduce the original feature graph through introducing a basis for the original feature space within which the selected feature subset is chosen. In other words, the reduced graph includes only the elements of the basis determined by the suggested method as its vertices. Instead of the original feature graph, the reduced graph can be applied as a model of the feature manifold. From a geometric point of view, this issue can be interpreted as follows. The manifold of features is represented by one of its sub-manifolds. This sub-manifold is determined by the basis that the proposed method introduces and it is composed of all the features included in that basis. Considering only the sub-manifold associated with a basis of features allows us to access the information about the whole feature manifold as well.

Let $B = [f^b_1, f^b_2, \ldots, f^b_n] \in \mathbb{R}^{n \times n}$ be a basis of features. Assuming that the basis of features exists on a sub-manifold of the original manifold of features, the similarity of the $i$th and the $j$th basis features for $i, j \in \{1, 2, \ldots, n\}$, is denoted by $S^b_{ij}$ and is defined as:

$$S^b_{ij} = \begin{cases} e^{-\frac{||f^b_i - f^b_j||^2_2}{\sigma}} & \text{if } f^b_i \in N(f^b_j) \text{ or } f^b_j \in N(f^b_i), \\ 0 & \text{otherwise} \end{cases},$$

where the set $N(f^b_i)$ is the $k$-nearest neighbor corresponding to $f^b_i$, and $\sigma$ denotes the Gaussian parameter. In addition, the feature Laplacian matrix $L^b$ corresponding to the basis of features is defined as $L^b = D^b - S^b$ in which $D^b$ is an $n \times n$ diagonal matrix with the diagonal entries $D^b_{ii} = \sum_{j=1}^n S^b_{ij}$.

Suppose that the coefficient matrix is $H = [h_1, h_2, \ldots, h_n]$. The problem regarding the preserv-
tion of the geometric structure of features associated with the basis manifold is presented as:

\[
\text{Tr}(H^bL^bH^T) = \frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \|h_i - h_j\|_2^2 S_{ij}^b.
\] (18)

Considering Eq. (18), in case the level of the similarity between the \(i\)th and the \(j\)th features in the basis is sufficiently high, the similarity between the vectors \(h_i\) and \(h_j\) are also considerably strong. It means that the similarity of the feature vectors in the original manifold implies the similarity of the corresponding vectors in the sub-manifold determined by the basis. This representation leads to a substantial reduction of the feature manifold so that the local geometry of features is preserved while the size of the graph is enormously reduced. This issue enhances the performance of our method and reduces the computational costs significantly.

Considering the ultimate goal of an unsupervised feature selection to search a low-dimensional feature subset that best characterizes the original feature space, we define the optimization framework of the proposed method B-SGFS as follows:

\[
\arg\min_{G \in \mathbb{R}^{n \times k}, H \in \mathbb{R}^{k \times n}} \|B - BGH\|_F^2 + \alpha \text{Tr}(H^bL^bH^T) + \beta \|G\|_{2,1},
\] (19)

in which \(\alpha\) and \(\beta\) are two regularization parameters, and the following principle fact has been utilized in (19): every feature subset of \(X\) can be expressed uniquely as a linear combination of the features associated with the basis \(B\), and as an immediate consequence, it is deduced that there is a matrix \(G \in \mathbb{R}^{n \times k}\) such that \(X_I = BG\).

In order to find \(G\) and \(H\), we make use of an alternating least squares iterative method in such a way that we will solve (19) with respect to one variable while the other variable is assumed to be fixed. For this purpose, first, let \(H \in \mathbb{R}^{k \times n}\) is taken to be fixed and suppose that

\[
\mathcal{L}(G, H) = \|B - BGH\|_F^2 + \alpha \text{Tr}(H^bL^bH^T) + \beta \text{Tr}(G^TUG),
\]

where \(U \in \mathbb{R}^{n \times n}\) is a diagonal matrix with the diagonal elements \(U_{ii} = \frac{1}{2\|G_i\|_2}\) for \(i = 1, \ldots, n\).

Now, by taking the partial derivative of \(\mathcal{L}\) with respect to \(G\), we get

\[
\frac{\partial \mathcal{L}}{\partial G} = 2(B^TBGHH^T - B^T BH^T) + 2\beta U G.
\]

Next, we assume that \(\frac{\partial \mathcal{L}}{\partial G} = 0\). It follows that the matrix \(G\) satisfies Eq. (20).

\[
B^TBGHH^T + \beta U G = B^T BH^T.
\] (20)

In the same manner, we assume that \(G \in \mathbb{R}^{n \times k}\) is fixed and taking the partial derivative of \(\mathcal{L}\) with respect to \(H\), we have

\[
\frac{\partial \mathcal{L}}{\partial H} = 2(G^TB^T B G H - G^T B^T B) + 2\alpha H L^b.
\]

Now, suppose that \(\frac{\partial \mathcal{L}}{\partial H} = 0\). It results that the matrix \(H\) satisfies Eq. (21).

\[
G^T B^T B G H + \alpha H L^b = G^T B^T B.
\] (21)

Algorithm 5 presents a summary of what discussed in this section.
Algorithm 5 The B-SGFS method.

**Input:** Data matrix $X \in \mathbb{R}^{n \times d}$; the regularization parameters $\alpha$ and $\beta$; the number of the selected features $k$; the maximum number of iterations $\text{maxIter}$.

1. Initialize the basis of features $\{f^b_1, f^b_2, \ldots, f^b_n\}$ according to the technique of VS described in Algorithm 1 and consider the matrix $B = [f^b_1, f^b_2, \ldots, f^b_n]$.
2. Compute the feature Laplacian matrix $L^b$ corresponding to the basis of features.
3. Initialize $H \in \mathbb{R}^{k \times n}$.
4. Initialize the diagonal matrix $U \in \mathbb{R}^{n \times n}$.

5. while iteration $\leq \text{maxIter}$ do
6. Fix $H$ and solve the matrix equation $B^T B G H H^T + \beta U G = B^T B H^T$ for $G$.
7. Fix $G$ and solve the matrix equation $G^T B^T B G H + \alpha H L = G^T B^T B$ for $H$.
8. Update the diagonal matrix $U$ according to the rule $U_{ii} = \frac{1}{2\|G_i\|_2^2}$, where $i = 1, \ldots, n$.
9. end while

**Output:** Sort the norm of rows of the weight feature matrix, $\|G_i\|_2$, in a descending order and put the rows of $G$ in an order induced by the values of $\|G_i\|_2$, where $i = 1, \ldots, n$. Next, from $\{f^b_1, f^b_2, \ldots, f^b_n\}$, select $k$ features whose corresponding rows in $G$ are evaluated as the top score rows according to the norm function.

### 3.4 Solving the Matrix Equations in B-MFFS, B-MPMR and B-SGFS

As it was discussed in the previous sections, the following matrix equations have been derived for the three proposed methods B-MFFS, B-MPMR and B-SGFS.

**B-MFFS:**

\[
\begin{align*}
\text{To obtain } G: & \quad B^T B G H H^T + \alpha U G = B^T B H^T. \\
\text{To obtain } H: & \quad G^T B^T B G H = G^T B^T B.
\end{align*}
\]

**B-MPMR:**

\[
\begin{align*}
\text{To obtain } G: & \quad B^T B G (H H^T + \alpha 1_k) + \beta U G = B^T B H^T. \\
\text{To obtain } H: & \quad G^T B^T B G H = G^T B^T B.
\end{align*}
\]

**B-SGFS:**

\[
\begin{align*}
\text{To obtain } G: & \quad B^T B G H H^T + \beta U G = B^T B H^T. \\
\text{To obtain } H: & \quad G^T B^T B G H + \alpha H L = G^T B^T B.
\end{align*}
\]

Since the coefficient matrices described in the above matrix equations are symmetric positive semi-definite, according to the results mentioned in [52], we can employ some reliable techniques such as the global conjugate gradient method to solve such matrix equations. In Appendix A, the details of the global conjugate gradient method will be explained. For more information about this method as well as the global version of this method, see [52, 53, 54] and references therein.

### 3.5 Comparison Between Our Proposed Methods and Corresponding Conventional Methods

One of the main advantages of using the basis of features in the frameworks presented in this study is that it significantly reduces the volume of calculations, and consequently, the time required in the process of reducing the high-dimensional space. Basically, the original formula used in the feature selection problem, presented in some state-of-the-art methods MFFS, MPMR and SGFS, is defined as:

\[
\min_{W \in \mathbb{R}^{d \times k}, H \in \mathbb{R}^{k \times d}} \|X - X W H\|_F, \tag{22}
\]

14
where $X = [f_1, f_2, \ldots, f_d] \in \mathbb{R}^{n \times d}$ is the data matrix, in which $n$ and $d$ indicates the number of samples and that of features, respectively. It should be here noted that in the high-dimensional datasets such as those examined in our experiments, the value of $d$ can be very high. Furthermore, as it is discussed in the literature of the methods MFFS, MPMR and SGFS, the computational cost of these methods is strongly dependent on the number of features (i.e., $d$), and the greater the number of features, the longer the execution time of these methods [37, 38, 39].

On the other hand, the feature selection problem used in the frameworks presented in this study, i.e., B-MFFS, B-MPMR and B-SGFS, is defined as:

$$\min_{G \in \mathbb{R}^{n \times k}, H \in \mathbb{R}^{k \times n}} \|B - BGH\|_F,$$

(23)

where $B = [f_1^b, f_2^b, \ldots, f_n^b] \in \mathbb{R}^{n \times n}$ is a basis for the feature space generated by $\{f_1, f_2, \ldots, f_d\}$. Contrary to what is seen in Problem (22), the computational cost of our methods depends on the value of $n$, which is much smaller than $d$. In addition to this, a comparison between the size of the matrices $X$ and $B$ shows that the size of $B$ can be much smaller than that of $X$. For this reason, it would be perfectly reasonable to expect that the time required for our proposed methods B-MFFS, B-MPMR and B-SGFS in the process of reducing the high-dimensional space to be much lower than the time required by the original methods MFFS, MPMR and SGFS.

In order to verify the theoretical results given here, the running time of all the proposed methods and their corresponding conventional versions are illustrated in Section 4.10. The numerical results indicate that the running time corresponding to all the proposed methods B-MFFS, B-MPMR and B-SGFS is much shorter compared to all the conventional methods MFFS, MPMR and SGFS. This means that applying the notion of basis to modify MFFS, MPMR and SGFS succeeds to enhance these methods in terms of their running time. As discussed above, the reason for such an improvement comes from the fact that representing the original data by a basis of the feature space reduces the consequent level of computations. Therefor, he proposed methods present modifications of the corresponding conventional methods in terms of the amount of calculations and running time.

4 Computational Experimental Results

To validate the three proposed methods, B-MFFS, B-MPMR and B-SGFS, in terms of their performance, some experiments are conducted in this section on a number of well-known biological datasets. To analyze the results, two essential metrics are used. First, these metrics are described in brief. Following this, the results of the experiments regarding the efficiency and efficacy of the proposed techniques are discussed.

4.1 Datasets

In this paper, to evaluate the proposed methods, eight cancer gene expression datasets are used. A description of these datasets is presented in Table 1. Embryonal Tumors of Central Nervous System (CNS) dataset comprises tumor samples gene expression data from patients with different types of brain tumors. This dataset provides the gene expression profiles and clinical data on the progression of the disease and therapeutic responses. This dataset was used to predict the disease outcomes based on molecular signatures [55]. The Colon cancer dataset that we have used has gene expression profiles of the tumor and normal colon samples (40 tumor tissue samples and 22 normal colon tissue). This
dataset has been used to detect gene expression signatures and patterns differentiating colon tissue types in normal conditions and neoplasia[56]. The dataset of Diffuse Large B-cell Lymphomas (DLBCL) has the gene expression profiles of tumor samples from patients with DLBCL, which is the most common form of non-Hodgkin’s lymphoma [57]. Glioblastoma Multiforme (GBM) dataset includes gene expression profiles of samples from GBM tumors with the clinical characteristics of the patients in terms of outcome [58]. TOX-171 dataset is a toxicology-based gene expression set with integrated biological phenotypes and characteristics [28]. The Lung Cancer and Prostate Cancer datasets are integrated gene expression profiles from lung and prostate tumors [59]. The Cancer Cell Line Encyclopedia (CCLE) has the gene expression data of common cancer cell lines used in cancer research and contains metrics of the therapeutic responses and mutational properties of their genomes [60].

Table 1: The information of eight datasets utilized in the computational experiments.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Abbreviation</th>
<th># Samples</th>
<th># Features</th>
<th># Classes</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Embryonal Tumors of CNS</td>
<td>CNS</td>
<td>60</td>
<td>7129</td>
<td>2</td>
<td>[55]</td>
</tr>
<tr>
<td>Colon Cancer</td>
<td>Colon</td>
<td>62</td>
<td>2000</td>
<td>2</td>
<td>[56]</td>
</tr>
<tr>
<td>Diffuse Large B-cell Lymphomas</td>
<td>DLBCL</td>
<td>47</td>
<td>4026</td>
<td>2</td>
<td>[57]</td>
</tr>
<tr>
<td>Glioblastoma Multiforme</td>
<td>GBM</td>
<td>85</td>
<td>22283</td>
<td>2</td>
<td>[58]</td>
</tr>
<tr>
<td>TOX-171</td>
<td>TOX-171</td>
<td>171</td>
<td>5748</td>
<td>4</td>
<td>[28]</td>
</tr>
<tr>
<td>Lung Cancer</td>
<td>Lung</td>
<td>73</td>
<td>325</td>
<td>7</td>
<td>[59]</td>
</tr>
<tr>
<td>Prostate Cancer</td>
<td>Prostate</td>
<td>102</td>
<td>5966</td>
<td>2</td>
<td>[59]</td>
</tr>
<tr>
<td>Cancer Cell Line Encyclopedia</td>
<td>CCLE</td>
<td>343</td>
<td>1900</td>
<td>2</td>
<td>[60]</td>
</tr>
</tbody>
</table>

4.2 Computational Experimental Setting

Before performing the experiments, the parameters included in the methods are set as follows. For all methods, the tuning of the regular parameters is carried out making use of the grid-search algorithm. This issue is a leading factor in drawing reasonable and fair comparisons. This way, we set $\alpha = 1$ in MPMR and $\rho = 10^8$ in MPMR and MFFS [37, 38]. In addition, the search domain is selected as $\{10^i\mid i = -8, -7, \ldots, 7, 8\}$ for the rest of the balancing parameters in all methods. Moreover, for all methods, the maximum number of iterations and the number of the selected features from each dataset are set to 30 and $\{10, 20, 30, \ldots, 100\}$, respectively. For methods that preserve the geometric structure of features, $\{10^i\mid i = 0, 1, \ldots, 5\}$ is considered as the search domain for $\sigma$, and each neighbor size is selected to be 5 on each dataset. Employing the $k$-means technique, the performance of the process of clustering is examined subject to the selected features, where for a given dataset, the true number of the classes is denoted by $c$. When the feature selection task is performed and is finished, the selected features are used to guide the $k$-means algorithm to carry out the samples clustering task. Since the clustering performance of the $k$-means algorithm relies upon initialization, in the experiments, the clustering process is repeated 20 times where diverse initializations take place at random. Then, the average clustering results are reported.

4.3 Evaluation Metrics for Comparison

In order to carry out proper evaluation of the proposed and benchmark algorithms, two commonly used performance evaluation metrics for unsupervised methods, including Clustering Accuracy (ACC) and Normalized Mutual Information (NMI), are employed. Notice that a greater value of ACC/NMI corresponding to a method indicates that the method under consideration achieves a better performance.
The metric ACC is formulated by Eq. (24).

\[
\text{ACC} = \frac{1}{k} \sum_{j=1}^{k} \delta(d_j, \text{map}(e_j)),
\]

(24)

in which \(k\) denotes the total number of samples, \(d_j\) and \(e_j\) symbolize respectively the ground truth label for the sample \(x_i\) and the clustering label, \(\text{map}(\cdot)\) represents an optimal mapping function that handles the task of matching the labels predicted by the given feature selection method to the real ones, and the indicator function \(\delta(a, b)\) is defined by Eq. (25).

\[
\delta(a, b) = \begin{cases} 
1 & \text{if } a = b \\
0 & \text{otherwise} 
\end{cases}
\]

(25)

Let \(W\) and \(Z\) be random variables. The metric NMI is given by Eq. (26).

\[
\text{NMI}(W, Z) = \frac{I(W, Z)}{\sqrt{H(W)H(Z)}},
\]

(26)

where \(I(W, Z)\) indicates the mutual information between \(W\) and \(Z\), and \(H(\cdot)\) is the entropy function.

### 4.4 Methods and Modalities

To conduct the experiments, we make comparisons between the proposed techniques and a number of benchmark methods including the Baseline, Variance Score (VS) [61], Laplacian Score (LS) [62], MFFS [37], MPMR [38] and SGFS [39]. In Baseline, all original features are used. Variance Score computes every feature’s variance, and determines the optimal features as the ones whose variance is the largest compared to the other features. In Laplacian Score, the intrinsic geometry of data is taken into consideration. The last three methods, MFFS, MPMR and SGFS, were also described in Section 3.

### 4.5 Results and Discussion

For comparisons of the proposed methods with other techniques, six various state-of-the-art methods were examined in terms of their performance on seven benchmark datasets. To this aim, ACC and NMI were computed for the selected methods. The results of these experiments are presented in Tables 2 and 3 in which the first and the second best results are marked in boldface and are underlined, respectively.

According to Table 2, on all the datasets, the first and the second best values of ACC were achieved by the three proposed methods, B-SGFS, B-MPMR and B-MFFS. In addition, B-MFFS outperformed all the conventional methods in terms of ACC. With the exception of the ACC value for MPMR which equals that value for B-MPMR on DLBCL, B-MPMR gave a better performance compared to the conventional methods. It is also clear that B-SGFS showed a better performance compared to the state-of-the-art methods except for SGFS on CNS. In this case the difference between the ACC value of the two methods is very small. A key aspect of the proposed methods is the lower number of the selected features compared to that number corresponding to the conventional methods. It is clear from Table 2 that this number, presented in parentheses, is far smaller for our three methods than that number for the other methods in most cases which is a big advantage. Figure 2 illustrates the bar chart of ACC values for all the methods indicated in Table 2 on the seven benchmark datasets presented in Table 1.
Table 2: Clustering accuracy (ACC±STD%) results for different feature selection methods on seven datasets. The numbers in parentheses are the numbers of selected features corresponding to the best clustering results. The best results in each row are highlighted in bold and the second best results are underlined. Note that the higher ACC values are, the better the clustering performance is.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Baseline</th>
<th>VS</th>
<th>LS</th>
<th>MFFS</th>
<th>MPMR</th>
<th>SGFS</th>
<th>B-MFFS</th>
<th>B-MPMR</th>
<th>B-SGFS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNS</td>
<td>53.33 ± 1.45</td>
<td>56.66 ± 0.04</td>
<td>58.66 ± 1.02</td>
<td>61.75 ± 2.78</td>
<td>61.66 ± 0.00</td>
<td>61.12 ± 0.00</td>
<td>62.33 ± 3.05</td>
<td>62.25 ± 3.37</td>
<td>64.35 ± 3.35</td>
</tr>
<tr>
<td>Colon</td>
<td>78.54 ± 6.95</td>
<td>58.06 ± 1.14</td>
<td>56.66 ± 9.38</td>
<td>74.19 ± 0.29</td>
<td>74.21 ± 2.91</td>
<td>77.41 ± 1.45</td>
<td>80.64 ± 0.03</td>
<td>84.91 ± 2.52</td>
<td>85.48 ± 0.00</td>
</tr>
<tr>
<td>DLBCL</td>
<td>59.57 ± 1.23</td>
<td>65.95 ± 0.01</td>
<td>57.44 ± 0.72</td>
<td>85.10 ± 0.14</td>
<td>87.23 ± 0.14</td>
<td>80.85 ± 0.43</td>
<td>95.74 ± 0.29</td>
<td>87.23 ± 0.29</td>
<td>84.91 ± 2.85</td>
</tr>
<tr>
<td>GBM</td>
<td>67.05 ± 0.01</td>
<td>68.23 ± 0.00</td>
<td>69.41 ± 0.00</td>
<td>70.58 ± 0.00</td>
<td>72.23 ± 0.00</td>
<td>75.29 ± 0.00</td>
<td>75.29 ± 1.45</td>
<td>75.29 ± 1.45</td>
<td>76.47 ± 0.26</td>
</tr>
<tr>
<td>TOX-171</td>
<td>41.25 ± 0.72</td>
<td>42.13 ± 0.99</td>
<td>43.35 ± 0.72</td>
<td>41.73 ± 0.13</td>
<td>41.49 ± 1.65</td>
<td>38.26 ± 0.87</td>
<td>45.08 ± 0.54</td>
<td>44.82 ± 0.91</td>
<td>44.44 ± 0.72</td>
</tr>
<tr>
<td>Lung</td>
<td>76.78 ± 5.03</td>
<td>59.38 ± 5.68</td>
<td>65.82 ± 3.72</td>
<td>77.80 ± 4.6130</td>
<td>79.38 ± 6.57</td>
<td>80.75 ± 5.45</td>
<td>81.16 ± 4.96</td>
<td>82.46 ± 5.83</td>
<td>82.09 ± 6.91</td>
</tr>
<tr>
<td>Prostate</td>
<td>57.84 ± 0.00</td>
<td>65.68 ± 3.86</td>
<td>60.78 ± 0.11</td>
<td>61.76 ± 0.00</td>
<td>62.84 ± 1.38</td>
<td>63.72 ± 1.45</td>
<td>69.60 ± 0.02</td>
<td>69.72 ± 0.01</td>
<td>67.64 ± 0.00</td>
</tr>
</tbody>
</table>

Figure 2: The average results of the ACC metric on seven datasets. The x-axis indicates the different datasets, and the y-axis represents the values of the ACC results obtained for each feature selection method. Note that the higher ACC values are, the better the clustering performance is.

The blue, black and red bars that correspond to B-SGFS, B-MPMR and B-MFFS, respectively, show the superiority of our proposed methods over all the other ones except for two mentioned cases.

Moving to Table 3, it is clear that B-MFFS, B-MPMR and B-SGFS are infinitely preferable in terms of NMI than the conventional methods on all the datasets used in this paper with the exception of two cases. The first exception occurred for SGFS on CNS, where this algorithm performed better compared with all our three methods, and the second one was raised on DLBCL in which the values of NMI for B-PPMR and MPMR were equal. In all the other cases, the first and second best results were obtained by the proposed methods. Again, the number of the selected features can be considered as an outstanding achievement of our methods since in almost all cases, these algorithms selected lower number of the features in compared to the conventional methods. In Figure 3, the bar chart of the values of NMI corresponding to the proposed and conventional methods on the seven benchmark datasets are shown. It can be seen from Figure 3 that B-MFFS, B-MPMR and B-SGFS, illustrated by blue, black and red bars, respectively, outperformed the conventional methods with two aforementioned exceptions.
Table 3: Normalized mutual information (NMI±STD%) results for different feature selection methods on seven datasets. The numbers in parentheses are the number of selected features corresponding to the best clustering results. The best results in each row are highlighted in bold and the second best results are underlined. Note that the higher NMI values are, the better the clustering performance is.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Baseline</th>
<th>VS</th>
<th>LS</th>
<th>MFFS</th>
<th>MPMR</th>
<th>SGFS</th>
<th>B-MFFS</th>
<th>B-MPMR</th>
<th>B-SGFS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNS</td>
<td>1.17 ± 0.00</td>
<td>2.25 ± 0.45 (70)</td>
<td>2.44 ± 0.45 (90)</td>
<td>2.93 ± 0.45 (100)</td>
<td>1.60 ± 0.80 (30)</td>
<td>8.79 ± 0.00 (30)</td>
<td>3.91 ± 0.59 (10)</td>
<td>3.61 ± 0.31 (30)</td>
<td>3.26 ± 0.33 (20)</td>
</tr>
<tr>
<td>Colon</td>
<td>2.93 ± 0.19</td>
<td>2.96 ± 1.63 (10)</td>
<td>7.27 ± 1.23 (100)</td>
<td>24.30 ± 0.14 (100)</td>
<td>21.08 ± 0.54 (40)</td>
<td>30.03 ± 0.00 (10)</td>
<td>33.03 ± 0.01 (10)</td>
<td>36.67 ± 5.43 (10)</td>
<td>37.89 ± 1.82 (10)</td>
</tr>
<tr>
<td>DLBCL</td>
<td>2.93 ± 0.19</td>
<td>2.04 ± 0.45 (40)</td>
<td>2.45 ± 0.00 (40)</td>
<td>48.46 ± 0.72 (70)</td>
<td>30.54 ± 0.72 (40)</td>
<td>78.80 ± 0.00 (10)</td>
<td>58.06 ± 0.07 (10)</td>
<td>31.64 ± 0.27 (10)</td>
<td></td>
</tr>
<tr>
<td>GBM</td>
<td>21.74 ± 0.03</td>
<td>6.61 ± 1.82 (40)</td>
<td>5.52 ± 2.75 (60)</td>
<td>7.10 ± 0.18 (30)</td>
<td>7.10 ± 0.82 (10)</td>
<td>8.75 ± 0.00 (60)</td>
<td>23.91 ± 0.36 (10)</td>
<td>25.28 ± 1.09 (10)</td>
<td>18.51 ± 0.01 (10)</td>
</tr>
<tr>
<td>TOX-171</td>
<td>13.54 ± 0.23</td>
<td>12.27 ± 0.65 (20)</td>
<td>11.68 ± 1.97 (90)</td>
<td>12.82 ± 1.52 (40)</td>
<td>16.22 ± 1.89 (60)</td>
<td>10.41 ± 1.02 (10)</td>
<td>17.39 ± 0.89 (10)</td>
<td>16.83 ± 1.03 (10)</td>
<td>18.51 ± 0.01 (10)</td>
</tr>
<tr>
<td>Lung</td>
<td>13.13 ± 2.52</td>
<td>56.36 ± 3.32 (100)</td>
<td>62.85 ± 2.16 (100)</td>
<td>73.22 ± 4.43 (100)</td>
<td>73.67 ± 4.49 (100)</td>
<td>75.61 ± 4.38 (50)</td>
<td>76.41 ± 3.61 (50)</td>
<td>77.29 ± 4.50 (50)</td>
<td>79.01 ± 5.33 (50)</td>
</tr>
<tr>
<td>Prostate</td>
<td>1.85 ± 0.03</td>
<td>5.25 ± 0.00 (10)</td>
<td>5.28 ± 0.06 (10)</td>
<td>5.29 ± 0.03 (50)</td>
<td>7.74 ± 1.28 (60)</td>
<td>7.45 ± 0.91 (10)</td>
<td>13.43 ± 0.79 (10)</td>
<td>16.03 ± 0.27 (10)</td>
<td>10.21 ± 0.22 (10)</td>
</tr>
</tbody>
</table>

Figure 3: The average results of the NMI metric on seven datasets. The x-axis indicates the different datasets, and the y-axis represents the values of the NMI results obtained for each feature selection method. Note that the higher NMI values are, the better the clustering performance is.

Furthermore, Figure 4 presents the bar charts of both the ACC and NMI values of all the conventional and proposed algorithms averaged over all the benchmark datasets. This figure shows that for all the three proposed methods, the average values of ACC and NMI, illustrated by the gray and red bars, respectively, are higher than that values corresponding to the conventional methods. This means that overall, the B-MFFS, B-MPMR and B-SGFS algorithms show a stronger performance in terms of ACC and NMI compared to the benchmark methods.

4.6 Comparison Between MFFS and B-MFFS

The data in Table 2 show that in terms of ACC, B-MFFS outperforms the MFFS on all the datasets. Those data also indicate that B-MFFS stands at the first and second ranks on two datasets, DLBCL and TOX-171, and on the dataset GBM, respectively. In addition, it is apparent from Table 2 that compared
Figure 4: The average values of the metrics ACC and NMI. The $x$-axis indicates the different feature selection methods, and the $y$-axis represents the obtained values of the metrics. Note that the higher ACC and NMI values are, the better the clustering performance is.

to MFFS, B-MFFS selects a lower number of features on all datasets except for Lung, where both numbers are equal. Based on the data given in Table 3, it is clear that B-MFFS is superior in terms of NMI to MFFS on all the datasets and comparing all the considered methods, its results achieves the first and the second ranks respectively on the dataset DLBCL and on the datasets TOX-171 and Lung. Moreover, in terms of NMI, the number of the features selected by B-MFFS is lower than that number corresponding to MFFS. In Figure 5, the bar charts of the values of ACC and NMI, relevant to MFFS and B-MFFS and averaged over all the benchmark datasets, are presented. As Figure 5 shows, the average values of both indices for B-MFFS, represented by blue bars, are greater than that values for B-MFFS, represented by red bars. Thus, the average performance of B-MFFS is better than that of its corresponding conventional method.

Figure 5: Comparison results of the MFFS and B-MFFS methods in respect of the average values of the metrics ACC and NMI.
4.7 Comparison Between MPMR and B-MPMR

To compare B-MPMR with its corresponding conventional method, the data in Tables 2 and 3 should be considered. It is obvious from these tables that in terms of both ACC and NMI, B-MPMR performs better than the MPMR over all the datasets except for DLBCL over which both the methods have the same performance. Furthermore, compared to all the methods considered in the experiments, B-MPMR stands at the first and the second ranks in terms of ACC respectively on two datasets Lung and Prostate and on the datasets CNS, Colon, GBM and TOX-171. In terms of NMI, B-MPMR also achieves the first and second best results on the dataset Prostate and on the datasets Colon, GBM and Lung, respectively. Moreover, it can be seen from Tables 2 and 3 that the number of the features selected by B-MPMR is lower compared to that number for MPMR over all the datasets with three exceptions including GBM in Table 2 and CNS and GBM in Table 3, where both methods selects the same number of features. The bar charts of the ACC and NMI values for MPMR and B-MPMR, averaged over all the benchmark datasets, are depicted in Figure 6. It is apparent from Figure 6 that the overall performance of B-MPMR is stronger in terms of ACC and NMI compared to that of MPMR since for B-MPMR, the ACC and NMI average values, that are illustrated by black bars, are greater compared with that values for MPMR, which are represented by purple bars.

![MPMR vs. B-MPMR](image)

Figure 6: Comparison results of the MPMR and B-MPMR methods in respect of the average values of the metrics ACC and NMI.

4.8 Comparison Between SGFS and B-SGFS

As it can be seen from Tables 2 and 3, the values of ACC and NMI for B-SGFS is better than that of SGFS over all the datasets with the exception of CNS over which the NMI value for SGFS is greater than that of B-SGFS. In addition, compared to all the conventional methods and the two other proposed methods, B-SGFS holds the first and the second ranks in terms of ACC over the datasets CNS, Colon and GBM and over the datasets DLBCL and Lung, respectively. When considering NMI, the first and the second best results of B-SGFS is obtained over the datasets Colon, GBM, TOX-171 and Lung and on the datasets CNS and DLBCL, respectively. Furthermore, B-SGFS selects a smaller number of features in compared to SGFS over all the benchmark datasets except for Prostate in Table 2 and Colon and Prostate in Table 3, where the number of features selected by both methods are the same. Figure 7 illustrates the bar charts of the ACC and NMI values averaged over all the benchmark datasets corresponding to SGFS and B-SGFS. In this figure, the green bars represent the average value
of SGFS and the red bars illustrate that of B-SGFS. From Figure 7, it is clear that B-SGFS achieves a better average performance in terms of ACC and NMI than does SGFS.

![Comparison results of the SGFS and B-SGFS methods in respect of the average values of the metrics ACC and NMI.](image)

Figure 7: Comparison results of the SGFS and B-SGFS methods in respect of the average values of the metrics ACC and NMI.

### 4.9 Comparison Among the Proposed Methods

To compare the proposed methods with each other, first, we consider their values of ACC in Table 2. Based on these data, B-SGFS holds the first rank on datasets CNS, Colon and GBM. Over CNS and Colon, B-MPMR stands at the second rank and on GBM, both B-MFFS and B-MPMR achieves the second rank simultaneously. The B-MFFS algorithm obtains the best results on the datasets DLBCL and TOX-171. The B-SGFS algorithm stands after it on DLBCL, while the B-MPMR algorithm produces the second best results on TOX-171. Finally, on the datasets Lung and Prostate, B-MPMR stands at the first rank. On Lung, B-SGFS holds the second rand while over Prostate, B-MFFS achieves the second best results. In terms of the number of the selected features, B-MFFS obtains the best results since it selects 10 features over all the benchmark datasets except for Lung over which it selects 50 features that is the same as the number of features the two other proposed methods select on this dataset. The B-MPMR algorithm selects 10 features over the rest of the datasets with the exception of TOX-171 over which it selects 40 features, while B-SGFS selects 20 features on CNS and TOX-171 and 10 feature on the rest of the datasets except for Lung.

Moving to Table 3, the NMI values are considered to make comparisons of our proposed methods. It is clear form this Table that with the exception of CNS on which SGFS produces the best results, for rest of the datasets, our methods obtains the best results. Even on CNS, B-SGFS holds the second best rank. This method also stands at the first rank over four datasets including Colon, GBM, TOX-171 and Lung. The B-MPMR method stands after it over Colon, GBM and Lung, while B-MFFS holds the second rank on TOX-171. The data in Table 3 also indicates that on the dataset Prostate, B-MPMR and B-MFFS achieve the first and the second best results, respectively, while over the dataset DLBCL, B-MFFS and B-SGFS produce respectively the first and the second best outcomes. Moreover, B-MPMR selects 30 and 50 features on CNS and Lung, respectively, and it selects 10 features over the rest of the datasets. In terms of the number of the selected features, B-MPMR perform better than the two other proposed methods. The B-MFFS algorithm selects 50 features on each of TOX-171 and Lung, while it selects 10 features over the rest of the datasets. The number of the features selected by B-SGFS is 20 on each of CNS and TOX-171, 50 on Lung and 10 over the rest of benchmark datasets.
4.10 Methods and Their Running Times

In Table 4, the running time of all the proposed and the conventional methods are illustrated on all the benchmark datasets and for the case of $k = 40$.

Table 4: The CPU time in seconds for B-MFFS, B-MPMR, and B-SGFS as well as for their corresponding conventional methods. Here, the number of the selected features is set to $k = 40$.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MFFS</th>
<th>MPMR</th>
<th>SGFS</th>
<th>B-MFFS</th>
<th>B-MPMR</th>
<th>B-SGFS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNS</td>
<td>20.11</td>
<td>25.28</td>
<td>55.61</td>
<td>3.03</td>
<td>4.22</td>
<td>5.16</td>
</tr>
<tr>
<td>Colon</td>
<td>1.58</td>
<td>2.03</td>
<td>4.32</td>
<td>3.33</td>
<td>3.55</td>
<td>4.28</td>
</tr>
<tr>
<td>DLBCL</td>
<td>6.48</td>
<td>8.13</td>
<td>17.52</td>
<td>2.01</td>
<td>2.15</td>
<td>2.46</td>
</tr>
<tr>
<td>GBM</td>
<td>190.15</td>
<td>239.18</td>
<td>&gt;1000</td>
<td>2.05</td>
<td>2.34</td>
<td>3.18</td>
</tr>
<tr>
<td>TOX-171</td>
<td>12.82</td>
<td>16.52</td>
<td>35.82</td>
<td>2.75</td>
<td>2.80</td>
<td>3.43</td>
</tr>
<tr>
<td>Lung</td>
<td>1.06</td>
<td>1.07</td>
<td>1.1</td>
<td>1.46</td>
<td>1.43</td>
<td>1.78</td>
</tr>
<tr>
<td>Prostate</td>
<td>14.07</td>
<td>18.34</td>
<td>39.68</td>
<td>2.35</td>
<td>2.41</td>
<td>3.32</td>
</tr>
</tbody>
</table>

Table 4 shows that the running time corresponding to all the proposed algorithms is much shorter compared to all the conventional methods with one exception including MFFS, MPMR, and SGFS over the Lung dataset. In specific, it is clear that for the case of GBM, the running time of all the three proposed methods is far less than that of MFFS, MPMR, and SGFS.

In addition, the CPU time in seconds, required to build the basis matrix corresponding to each dataset, is reported in Table 5. It can be seen from this table that building the basis matrix needs a very short time to be completed. In point of fact, since the size of the basis matrix is directly related to its rank, the higher the rank of the dataset is, the longer the CPU time will be.

Table 5: The CPU time in seconds required for building the basis matrix corresponding to each dataset.

<table>
<thead>
<tr>
<th>Rank of dataset</th>
<th>CNS</th>
<th>Colon</th>
<th>DLBCL</th>
<th>GBM</th>
<th>TOX-171</th>
<th>Lung</th>
<th>Prostate</th>
<th>CCLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU time</td>
<td>0.0222</td>
<td>0.0261</td>
<td>0.0174</td>
<td>0.0478</td>
<td>0.2133</td>
<td>0.0413</td>
<td>0.0598</td>
<td>1.8154</td>
</tr>
</tbody>
</table>

The most significant finding that Tables 4 and 5 provide is that all the proposed methods are faster than their corresponding conventional methods, MFFS, MPMR and SGFS, over all datasets (Lung is excepted). This means that applying the notion of basis to modify MFFS, MPMR and SGFS succeeds to enhance these methods in terms of their running time. The reason for such an improvement lies in the fact that representing the original data by a basis of the feature space reduces the consequent level of computations. Hence, the proposed methods present modifications of the corresponding conventional methods in terms of the amount of calculations and running time as well as improving their performance as mentioned earlier.

5 Application to the Extraction of Gene Signatures for Drug Responses in Cancer

Over the last recent years, dimensionality reduction techniques and feature selection methods have been applied to a variety of real-world problems in different fields, particularly to large molecular and clinical data including gene expression datasets [63, 64]. For instance, to study complex diseases such...
as cancer from a molecular perspective, it is required to reduce the dimension of the molecular data and to deal with the noise and redundancy included in such types of data. In this regard, a combination of molecular signature approaches with feature selection and reduction techniques can substantially help to provide new solutions.

In this section, the newly developed methods, B-MFFS, B-MPMR and B-SGFS, are incorporated into gene signature extraction to develop an analytical framework for cancer cell lines’ responses to Nilotinib treatment. In addition, the efficiency and productivity of these methods are evaluated using the CCLE dataset [60]. Furthermore, the proposed methods are compared to MFFS, MPMR and SGFS in terms of their performance.

5.1 Computational Experimental Metrics

To assess the proposed methods in terms of their classification performance and to corroborate their efficiency and efficacy, four types of evaluation metric including the classification accuracy (ACC), the true positive rate (TPR), the true negative rate (TNR), and the F1-score are employed. For a given dataset, these metrics are defined as:

\[
\text{ACC} = \frac{n_{\text{correct}}}{n}, \quad \text{TPR} = \frac{TP}{P}, \quad \text{TNR} = \frac{TN}{N}, \quad \text{and} \quad \text{F1-score} = \frac{2TP}{2TP + FP + FN},
\]

where \(n\) and \(n_{\text{correct}}\) indicate the number of the whole samples and the correct classified samples, respectively. Moreover, TP denotes the number of true positive samples, TN is the number of true negative samples, FP is the number of false positive samples, FN is the number of false negative samples, P indicates the number of positive samples and N is the number of negative samples.

The selected gene sets discovered by our proposed methods in each run have different sizes, specifically, the number of the selected genes is set to \(k = 2, 5, 10\). It should be pointed out that when the feature selection process is completed, the methods used for feature selection are appraised in terms of their performance making use of the Support Vector Machine (SVM) classifier [65]. To draw a comparison between the proposed and the benchmark feature selection techniques, a 10-fold cross-validation (CV) process is applied such that the whole dataset is randomly split into 10 groups in each CV process iteration. Then, nine of these groups are used to handle the classifier training phase and the remaining one is utilized to perform the testing phase. This process guarantees the generalization ability of the trained classifier to make accurate predictions of unseen samples. Moreover, the tuning process for setting the hyperparameters of the methods in the training phase is guided in each iteration by an internal five-fold CV grid search on the values presented in Subsection 4.2.

5.2 Results and Discussion

The experimental results regarding the performance of the proposed methods and its comparison with that of the benchmark methods are illustrated in Table 6 and Figure 8. It can be seen from Table 6 that the results of ACC, TNR and F1-score corresponding to the proposed methods B-MFFS, B-MPMR and B-SGFS show a better performance compared to that of the conventional methods MFFS, MPMR and SGFS. It is only in the case of TPR that the performance of the proposed methods is at a level the same as the conventional methods or is a bit lower than the performance of those techniques. Furthermore, in terms of ACC, TPR and TNR, the B-MFFS method outperforms B-SGFS and B-MPMR, while in terms of F1-score, B-SGFS performs better than the two other methods.
Table 6: Classification performance (ACC%, TPR%, TNR%, and F1-score%) results for different feature selection methods on the CCLE dataset. The number of the selected genes is set to $k = 2, 5, 10$, and the numbers in parentheses are the numbers of selected genes corresponding to the best classification results. The best results in each row are highlighted in bold and the second best results are underlined. Note that a higher ACC, TPR, TNR, and F-score value is indicative of a better classification performance.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MFFS</th>
<th>MPMR</th>
<th>SGFS</th>
<th>B-MFFS</th>
<th>B-MPMR</th>
<th>B-SGFS</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACC</td>
<td>72.68  (5)</td>
<td>71.70  (2)</td>
<td>72.59  (5)</td>
<td><strong>75.47</strong> (2)</td>
<td>72.64 (2)</td>
<td>80.19  (5)</td>
</tr>
<tr>
<td>TPR</td>
<td>98.00 (10)</td>
<td>98.00 (2)</td>
<td>98.00 (2)</td>
<td><strong>98.00</strong> (10)</td>
<td>97.35 (2)</td>
<td><strong>98.00</strong> (10)</td>
</tr>
<tr>
<td>TNR</td>
<td>20.13 (2)</td>
<td>16.67 (5)</td>
<td>12.32 (5)</td>
<td><strong>46.67</strong> (2)</td>
<td>33.56 (2)</td>
<td>43.33 (10)</td>
</tr>
<tr>
<td>F1-score</td>
<td>83.89 (5)</td>
<td>83.47 (2)</td>
<td>83.70 (5)</td>
<td>89.37 (2)</td>
<td><strong>87.86</strong> (2)</td>
<td><strong>87.27</strong> (5)</td>
</tr>
</tbody>
</table>

Figure 8 presents the results of the comparisons between each of the proposed methods and its conventional technique in terms of the average values of ACC, TPR, TNR and F1-score. It is clear from this figure that in terms of ACC, MPMR and B-MPMR have the same performance while B-MFFS and B-SGFS perform better than their conventional methods. Considering TPR, the performance of B-MFFS, B-MPMR and B-SGFS is similar to that of MFFS, MPMR and SGFS, respectively. Finally, all the proposed methods outperform their corresponding conventional technique in terms of TNR and F1-score.

Overall, the outcomes of experiments demonstrate that the incorporation of the notion of a basis into MFFS, MPMR and SGFS is completely effective.

5.3 Verification of the Feature Selections in Therapeutic Responses to Nilotinib

Oxidative response is one of the pathways activated upon Tyrosine Kinase Inhibitors (TKI) treatment in cells [66, 67, 68]. We used the gene sets assigned to Oxidative Response in cells from Molecular Biology of The Cell (MBC) ontology [69]. To find a gene signature extracted from Oxidative Stress response pathway, we used the Cancer Cell Line Encyclopedia (CCLE) data on drug treatment [60]. The dataset we used contained IC$_{50}$ for Nilotinib (a TKI inhibitor) treatment in 107 cancer cell lines IC$_{50}$ ranges of 0.0029 to 7.98 µM) and their gene expression profiles. The aim was to find gene sets that can predict the cell lines’ responses to Nilotinib in terms of IC$_{50} < 4$ µM. These gene sets were selected from oxidative stress pathway. The gene sets we discovered, had different sizes (gene numbers, $k = 2, 5, 10$). Our results demonstrate that applying B-MFFS and B-MPMR, we even can use a predictive 2-gene signatures. Table 7 shows which two genes are the most significant ones in each of these gene sets.

Figure 9 illustrates how the proposed techniques, B-MFFS, B-MPMR and B-SGFS, and their corresponding conventional methods differentiate resistant cases from sensitive ones (using IC$_{50} = 4$ µM as a threshold) with respect to two genes selected by each one of these methods as the selected features. It is apparent from Figure 9 that while MFFS, MPMR and SGFS are not able to clearly distinguish between the two categories of resistant and sensitive cases, the proposed methods carry out this task in a highly satisfying manner. All three B-MFFS, B-MPMR and B-SGFS techniques selected TNFRSF12A and YAP1 while MFFS and MPMR selected MSTN and GDF10 as the selected 2-gene signature. SGFS found GDF10 and UGT2B4 as selected features for a 2-gene signature. It has been shown that both
Figure 8: Results of comparison between “the MFFS and B-MFFS methods”, “the MPMR and B-MPMR methods”, and “the SGFS and B-SGFS methods” in respect of the average values of the metrics ACC, TPR, TNR, and F1-score.

Table 7: 2-gene signatures selected by different feature selection methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>MFFS</td>
<td>MSTN, GDF10</td>
</tr>
<tr>
<td>MPMR</td>
<td>MSTN, GDF10</td>
</tr>
<tr>
<td>SGFS</td>
<td>UGT2B4, GDF10</td>
</tr>
<tr>
<td>k = 2</td>
<td></td>
</tr>
<tr>
<td>B-MFFS</td>
<td>TNFRSF12A, YAP1</td>
</tr>
<tr>
<td>B-MPMR</td>
<td>TNFRSF12A, YAP1</td>
</tr>
<tr>
<td>B-SGFS</td>
<td>TNFRSF12A, YAP1</td>
</tr>
</tbody>
</table>

TNFRSF12A and YAP1 are involved in TKI treatment responsiveness in cancer [70, 71, 72, 73].
Figure 9: 2-gene signatures selected by different feature selection methods.

6 Conclusion

High-dimensional data analysis is a foundation for systems approaches in medicine [74]. The applicability of these approaches in the real-world settings requires high dimensionality reduction [74]. This reduction helps to build predictive models in systems pharmacology of cancer to optimize therapeutic regimens proposed by in vivo and in vitro experiments [75]. Our results showed that our modified
version of feature selection methods based on matrix factorization, we call B-MFFS, B-MPMR, and B-SGFS, could reduce the dimensionality of different datasets and provide feature signatures with higher accuracy than conventional methods. This is very important in the practice of quantitative and systems pharmacology, where we need to build both mechanistic and phenomenological models that emerge from omic data [76, 77]. To translate big complex data of omics and multi-omics into applied models for in vitro, in vivo, and clinical validation, we need such methodologies. This high dimensionality reduction can pave the path to establishing integrated systems pharmacology platforms that can predict drug responses in cancer cell lines and tumor models using small sets of bio-molecular markers [76].

One of the most potent applications of systems pharmacology is using dynamical models of signaling networks. These signaling networks consist of complex interactions among a large number of proteins or genes [78]. One of the primary practices to build and tune these models is to use prior biological knowledge to construct a reduced version of these interactions [78]. Our proposed methods can reduce these feature spaces quantitatively and select the most significant nodes in the interaction networks for more accurate and realistic dynamical models. The huge parameter spaces of these models also need reduction to represent a robust, optimized model [9]. Our dimensionality reduction methods can be applied to solve this problem as well.

Besides dynamic models, finding molecular sets to predict therapeutic responses in cancer models is an unmet need, and quantitative modalities like our proposed methodologies can provide the principles for this complex challenge. To improve the theoretical foundation of the proposed frameworks, it is important to determine that within a category of appropriate bases, which one is the most well-suited basis of features for a given unsupervised feature selection problem. This issue can be considered as a potential direction for future research.

Author Contributions


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Abbreviations

ACC, Clustering Accuracy; CCLE, Cancer Cell Line Encyclopedia; DLBCL, Diffuse large B-cell lymphoma; GB, Glioblastoma Multiforme; LS, Laplacian Score; MBC, Molecular Biology of The Cell; MFFS, matrix factorization; MPMR, Maximum Projection and Minimum Redundancy; NMF, Non-negative Matrix Factorization; NMI, Normalized Mutual Information; PCA, Principal Components Analysis; SGFS, Subspace learning-based Graph Regularized Feature Selection; SVD, Singular Value
Decomposition; TKI, Tyrosine Kinase Inhibitor; TPR, True Positive Rate; TNR, True Negative Rate; VS, Variance Score.

References


Appendix

A Global Conjugate Gradient Method

The global conjugate gradient method was developed as an extension of the conjugate gradient technique [54, 79]. Over the past recent years, it has been demonstrated that comparing to direct techniques such as the Gaussian elimination method, the iterative technique of global conjugate gradient enjoys a far better performance to deal with large matrix equations [80]. In such cases, it is usually supposed that constraints such as being both sparse and symmetric positive definite are imposed on the coefficient matrix corresponding to the given matrix equation [52]. A basic form of matrix equations whose exact solution can be estimated by this method is

$$A_1XB_1 + A_2XB_2 + \cdots + A_pXB_p = C,$$

(27)

in which the matrices $A_j \in \mathbb{R}^{n \times n}$ and $B_j \in \mathbb{R}^{m \times m}$ are symmetric positive definite (or symmetric semi-positive definite) for $j = 1, \ldots, p$ and $C \in \mathbb{R}^{n \times m}$. The unknown matrix is also denoted by $X \in \mathbb{R}^{n \times m}$.

Algorithm A1 presents the global conjugate gradient framework.

Algorithm A1 The global conjugate gradient method.

**Input:** The matrices $A_j \in \mathbb{R}^{n \times n}$, $B_j \in \mathbb{R}^{m \times m}$, for $j = 1, \ldots, p$, the right-hand side $C \in \mathbb{R}^{n \times m}$, an initial guess $X_0 \in \mathbb{R}^{n \times m}$, a tolerance $\epsilon$ and the maximum number of iterations $\text{maxIter}$.

**Output:** Return $X_k$ as the approximate solution of (27).

1: Set $\mathcal{M}(X) = A_1XB_1 + A_2XB_2 + \cdots + A_pXB_p$.
2: Set $R_0 = C - \mathcal{M}(X_0)$, and $P_0 = R_0$.
3: for $k = 1, \ldots, \text{maxIter}$ do
4: \quad Compute $\alpha_{k-1} = \frac{\|R_{k-1}\|_F^2}{\mathcal{M}(P_{k-1})^T(P_{k-1})}$.
5: \quad Compute $X_k = X_{k-1} + \alpha_{k-1}P_{k-1}$, and $R_k = R_{k-1} - \alpha_{k-1}\mathcal{M}(P_{k-1})$.
6: \quad If $\|R_{k+1} - R_k\|_F < \epsilon$ then return $X_k$.
7: \quad Compute $\beta_k = \frac{\|R_k\|_F^2}{\|R_{k-1}\|_F^2}$, and $P_k = R_k + \beta_kP_{k-1}$.
8: end for