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Graphical Exploration of Gene Expression Data:
A Comparative Study of Three Multivariate Methods

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SUMMARY. This article describes three multivariate projection methods and compares them for their ability to identify clusters of biological samples and genes using real-life data on gene expression levels of leukemia patients. It is shown that principal component analysis (PCA) has the disadvantage that the resulting principal factors are not very informative, while correspondence factor analysis (CFA) has difficulties regarding interpretation of the distances between objects. Spectral map analysis (SMA) is introduced as an alternative approach to the analysis of microarray data. Weighted SMA outperforms PCA and is at least as powerful as CFA in finding clusters in the samples as well as identifying genes related to these clusters. SMA addresses the problem of data
analysis in microarray experiments in a more appropriate manner than CFA and allows the application of a more flexible weighting to the genes and samples. Proper weighting is important since it enables less reliable data to be down-weighted and more reliable information to be emphasized.

KEY WORDS: Bioinformatics; Biplot; Correspondence factor analysis; Data mining; Data visualization; Gene expression data; Microarray data; Multivariate exploratory data analysis; Principal component analysis; Spectral map analysis.

1. Introduction

The advent of DNA microarray technology enabling global gene expression analysis has been a fundamental breakthrough in the life sciences. The possibility of simultaneously measuring the expression profile of thousands of genes allows for a better characterization of different types of a disease and for better insight in the underlying pathology, thus creating the possibility for identifying new therapeutic targets. In principle, DNA microarrays consist of some solid material upon which an array of spots of known DNA sequences, referred to as gene probes, is immobilized. RNA extracted from biological samples, is fluorescently labeled and applied to the array. The array is scanned and the fluorescent intensity at each position in the array is considered as a measure for the expression level of the corresponding gene. At present, a typical DNA microarray contains thousands of DNA-spots. In the near future, however, improvements to the technology will probably provide information on tens of thousands of genes, eventually encompassing entire genomes.
On the other hand, the simultaneous measurement of the expression level of thousands of genes poses an enormous task to the information processing capability of present systems. Much research is still being done in the area of statistics and data mining to provide the scientific community with better tools for pattern recognition and visualization of gene expression data. Methods of unsupervised learning, such as k-means clustering (Tavazoie et al., 1999), hierarchical clustering (Eisen et al., 1998), and self-organizing maps (Törönen et al., 1999) have found widespread application in analyzing and visualizing gene expression data. These methods however, produce results that are highly dependent on the distance-measure and clustering technique that is used and the number of clusters in a cluster analysis is often an issue of controversy. Furthermore, conventional clustering methods only allow for classification of either genes or biological samples alone, but do not allow interpretations of the association between genes and samples.

Another set of exploratory techniques is based upon projections of high-dimensional data in a lower dimensional space and plotting both genes and samples in this lower dimensional space using the biplot (Gabriel, 1971). Principal component analysis (PCA) (Pearson, 1901, Hotelling, 1933) is a well-established technique in multivariate statistics and has been applied to gene expression data (Chapman et al., 2001, Hilsenbeck et al., 1999, Landgrebe et al., 2002, Lefkovits et al., 1988). Related to PCA are procedures such as correspondence factor analysis (Benzécri, 1973) and spectral mapping (Lewi, 1976). Correspondence factor analysis (CFA) has recently been applied to microarray data by Fellenberg et al. (2001). In this paper, we propose the use of a less well-known technique,
spectral map analysis (SMA) (Lewi, 1976) for the analysis of gene expression data. In the past SMA has been successfully applied to a wide variety of problems ranging from pharmacology (Lewi, 1976), virology (Andries et al., 1990), to management and marketing research (Faes and Lewi, 1987). Thielemans, Lewi, and Massart (1988) have compared SMA with PCA and CFA, using a relatively small data set from the field of epidemiology. They concluded that the appropriate method depends upon the data to be analyzed and the features one is interested in. Up to now, the applications of SMA have always been limited to small or moderate sized data sets. The present paper illustrates the applicability of the method to large data sets and the importance of appropriate weighting in the analysis of microarray gene expression data. We will show that SMA provides the researcher with a visual data representation, useful as a tool for distinguishing patterns in the gene expression data that could be related to important biological questions.

The outline of this paper is as follows. In Section 2, a general framework for multivariate projection methods will be set up and the similarities and specifics of PCA, CFA, and SMA will be indicated. In Section 3, the different techniques will be compared using the gene expression profiles of leukemia patients (Golub et al., 1999). In Section 4, the advantages of weighted SMA for gene expression data will be highlighted and possible applications and limitations of the technique will be discussed.

2. Multivariate Projection Methods

The similarities and characteristics of the three multivariate projection methods, PCA, CFA, and SMA will be presented following Lewi and Moereels (1994) and Thielemans et al. (1988).
2.1 Notation

Let $X_{n \times p}$ denote the matrix containing the original expression levels $x_{ij}$ for the expression level of $n$ genes (rows) in each of $p$ different biological samples (columns). We also define two diagonal matrices with row weights $W_n$ and column weights $W_p$. The diagonal elements of $W_n$ and $W_p$ are the weight coefficients associated with the rows and columns of the matrix $X$. The weight coefficients are non-negative and normalized to unit sum. An unweighted analysis is obtained by $W_n = \text{diag}(1/n)$ and $W_p = \text{diag}(1/p)$. Alternatively, the diagonal elements of $W_n$ and $W_p$ can be set to appropriate weighting schemes such as the row and column totals, normalized to unity, i.e. $W_n = \text{diag}(X_1 / \sum_{i} X_{i1})$ and $W_p = \text{diag}(1_p / \sum_{j} X_{1j})$. There seems to be a consensus among scientists that micro-array data at lower levels of expression are less reliable, so weighing for row means seems appropriate in this context. An additional advantage of defining weights is the possibility of positioning rows and columns by setting their corresponding weights to zero. Positioning is the operation where some columns or rows of the data matrix are excluded from the actual analysis, but are still represented on the map constructed on the basis of the remaining data.

2.2 General algorithm for multivariate projection methods

In the algorithms of the three multivariate projection methods the following building blocks can be distinguished: re-expression, closure, centering, normalization, factorization, and projection. Differences between the methods are obtained by variations in these building blocks.
a. Re-expression

It is often advantageous to re-express (i.e. transform) the data as logarithms, i.e. a new matrix \( A \) is obtained whose elements \( a_{ij} = \log x_{ij} \). For this operation to be valid, measurements must be made on a ratio scale and the values must be positive. Logarithmic re-expression allows data in different physical units to be compared to one another as the logarithm of their ratios. In addition, it corrects for positive skewness and reduces the effect of large influential values. A further justification of a logarithmic re-expression is the fact that in many natural systems changes occur on a multiplicative rather than an additive scale. There is also the trivial case in which the original data are left unchanged and the elements of the re-expressed matrix \( A \) are equal to the elements of the data matrix \( X \).

b. Closure

Closure is defined as the operation of transforming the data into relative values such that they sum to unity. Closure requires the data to be non-negative and measured in the same units. From the matrix \( A \) with the re-expressed data, a new table \( B \) is obtained by either column, row, or global closure. In column-closure each element \( a_{ij} \) of \( A \) is divided by the corresponding column marginal total of \( A \), i.e. \( b_{ij} = \frac{a_{ij}}{a_{i+}} \), where \( a_{i+} = \sum_{j=1}^{n} a_{ij} \).

Column-closure imposes a linear constraint on the rows of the matrix. As a consequence, when \( n \leq p \), it reduces the rank of the data matrix by one. In row-closure the elements of the matrix \( B \) are obtained from \( A \) by dividing each element by the corresponding row marginal total, i.e. \( b_{ij} = \frac{a_{ij}}{a_{+j}} \), where \( a_{+j} = \sum_{i=1}^{p} a_{ij} \). A linear constraint is imposed on the columns of the matrix, resulting in a rank-reduction by one when \( p \leq n \). Double closure...
consists of the combined operation of dividing each element \( a_{ij} \) of the data matrix \( A \) by its corresponding row and column marginal total. The result is then multiplied by the total sum of \( A \) to yield a dimensionless number. We thus have: \( b_{ij} = \frac{a_{ij}a_{++}}{a_{rr}a_{++}} \), where 

\[
a_{++} = \sum_{j=1}^{p} \sum_{i=1}^{n} a_{ij}.
\]

Double closure always involves a reduction of the rank of the original data matrix by one. The operation of double closure combined with weighting of rows and columns by their corresponding marginal totals forms the core of CFA. Of course, one should also consider the trivial case of no closure in which \( b_{ij} = a_{ij} \).

c. Centering

Centering is defined as a correction of \( B \) for a mean value to yield the centered matrix \( Y \). There are different ways to derive mean values from a matrix, each resulting in a different way to center the data. Geometrically, centering involves a translation to the origin of the data in the column-space, the row-space, or in both. In column centering, the matrix \( Y = B - 1_n m_p^T \) contains deviations from the weighted column means \( m_p^T = 1_n^T W_n^T B \). In row centering, \( Y = B - m_n 1_p^T \) is the matrix with deviations from the weighted row means \( m_n^T = BW_p 1_p \). In global centering, the matrix \( Y = B - m \) contains the deviations of the elements \( B \) from the global weighted mean \( m = 1_n^T W_n^T BW_p 1_p \).

Simultaneous centering by rows and columns yields the double-centered matrix of deviations from row and column means \( Y = B - 1_n m_p^T - m_n 1_p^T + m_n 1_p^T \). The operation of double-centering involves a projection of the data matrix on a hyperplane that runs through the origin and is orthogonal to the line of identity. The result is a reduction by
one of the rank of the original matrix. The dimension that is lost is related to a component of “size” that is common to all elements of the data table and often obscures important information that is present in the data. Applying double-centering after logarithmic re-expression is the very essence of SMA. It is interesting to note the close analogy between double closure and double centering after logarithmic re-expression. For the centering part of the algorithm, we also define the trivial case of no centering with \( Y = B \).

d. Normalization

Normalization or standardization is the operation of dividing \( Y \) by the square root of the mean sums of squares or norm, yielding a normalized matrix \( Z \). There are several ways to compute the norm of a matrix each resulting in a different method of normalization. In column-normalization the normalized results is obtained as \( Z = YD_p^{-1} \), with the weighted column-norm \( D_p \) defined as \( D_p = \text{diag} \left( \left( Y^T \right)^{1/2} W_p 1_p \right)^{\frac{1}{2}} \). The effect of column-normalization in the column-space is to weight each column-dimension proportional to the inverse of its mean sum of squares. In the row-space, the effect is a sphericization, such that the points are forced to lie on a hypersphere. Column-normalization after column-centering is a standard operation in PCA. In row-normalization \( Z = D_c^{-1} Y \) with the weighted row-norm \( D_c \) defined as \( D_c = \text{diag} \left( Y^T W_p 1_p \right)^{\frac{1}{2}} \). The geometric interpretation of row-normalization is similar to that of column-normalization with the row and column spaces interchanged. Normalization for the weighted global norm \( d = 1_n W_n Y^2 W_p 1_p \) yields the global-normalized matrix \( Z = \frac{1}{d} Y \). For the sake of completeness, we also have the case of no normalization where \( Z = Y \).
e. Factorization

Factorization of $\mathbf{Z}$ yields factors that are orthogonal to one another and account for a maximum of the variance of the data. For a weighted analysis, the multivariate projection methods under consideration rely on the generalized singular value decomposition as factorization method. The generalized singular value decomposition of $\mathbf{Z}$ is defined as:

$$\mathbf{W}_n^{\frac{1}{2}} \mathbf{Z} \mathbf{W}_p^{\frac{1}{2}} = \mathbf{U} \mathbf{A} \mathbf{V}^T \quad [1]$$

where $\mathbf{A}$ is an $r \times r$ matrix of singular values, $r$ being the rank of $\mathbf{W}_n^{\frac{1}{2}} \mathbf{Z} \mathbf{W}_p^{\frac{1}{2}}$. In addition, we have $\mathbf{U}^T \mathbf{U} = \mathbf{I}_r$ and $\mathbf{V}^T \mathbf{V} = \mathbf{I}_r$. Consequently, $(\mathbf{W}_n^{-\frac{1}{2}} \mathbf{U})^T \mathbf{W}_n (\mathbf{W}_n^{-\frac{1}{2}} \mathbf{U}) = \mathbf{I}_r$ and $(\mathbf{W}_p^{-\frac{1}{2}} \mathbf{V})^T \mathbf{W}_p (\mathbf{W}_p^{-\frac{1}{2}} \mathbf{V}) = \mathbf{I}_r$.

f. Projection

Projection of the results of the generalized singular value decomposition along the first few common factors yields the biplot (Gabriel, 1971). Different biplots, with characteristic geometric properties, can be constructed using combinations of two factor-scaling coefficients $\alpha$ and $\beta$, set to either 0, 0.5, or 1, where the weighted factor scores $\mathbf{S}$ and factor loadings $\mathbf{L}$ are obtained from [1] by $\mathbf{S} = \mathbf{W}_n^{\frac{1}{2}} \mathbf{U} \mathbf{A}^\alpha$ and $\mathbf{L} = \mathbf{W}_p^{\frac{1}{2}} \mathbf{V} \mathbf{A}^\beta$. It is easy to show that the above expressions for $\mathbf{S}$ and $\mathbf{L}$ can also be written as $\mathbf{S} = \mathbf{Z} \mathbf{W}_p^{\frac{1}{2}} \mathbf{V} \mathbf{A}^{\alpha-1}$ and $\mathbf{L} = \mathbf{Z} \mathbf{W}_n^{\frac{1}{2}} \mathbf{U} \mathbf{A}^{\beta-1}$. The latter form, though more complex, is required for positioning supplementary rows or columns by setting their respective weights to zero. The following cases of factor scaling can be distinguished:
− \( \alpha = 1, \beta = 1 \) referred to as eigenvalue scaling. This type of symmetric scaling is customary in CFA. Distances of points in row-space as well as in column-space are reproduced in the plot, as well as the correlation structure of the column-variables.

− \( \alpha = 1, \beta = 0 \) referred to as unit column-variance scaling, is customary in PCA. Only distances between row-points are preserved in this asymmetric type of scaling. In full factor-space the distances of the column-items from the origin are constant and the correlation structure between column-variables is not reproduced.

− \( \alpha = 0, \beta = 1 \) referred to as unit row-variance scaling, is also customary in PCA. Only distances between the column-items and the correlation structure between column-variables are preserved. In full factor-space the distances of the row-points from the origin are constant.

− \( \alpha = 0.5, \beta = 0.5 \) referred to as singular value scaling, is customary in SMA. This type of factor-scaling is a compromise between the versions given above. Distances between row-points and the correlation structure of the column-variables are not fully reproduced. The distortion is most pronounced when the ratios between the eigenvalues \( \Lambda^2 \) associated with the axes of the biplot are very large or very small.

Having defined a general framework encompassing the three projection methods, we will now discuss their different characteristics.

2.1 Principal component analysis

Historically, PCA dates back to Pearson (1901) and Hotelling (1933). In the algorithm described above it is defined as: constant weighting of rows and columns, optional re-expression, column-centering, column-normalization, and factor scaling with either symmetric eigenvalue scaling with \( \alpha = 1, \beta = 1 \), asymmetric unit column-variances with
\( \alpha = 1, \beta = 0 \), or asymmetric unit row-variances with \( \alpha = 0, \beta = 1 \). Note that PCA makes a clear distinction between row- and column-items in the centering and normalization procedure.

2.2 Correspondence factor analysis

CFA has been developed by Benzécri (1973) and is adequately described by Greenacre (1984). This multivariate projection method was originally developed for the analysis of contingency tables but has also been applied to other tables with non-negative values (Fellenberg et al., 2001). CFA involves the following steps: weighting of rows and columns by marginal row and column totals, no re-expression, double-closure, double-centering, global normalization, and symmetric eigenvalue factor-scaling (\( \alpha = 1, \beta = 1 \)). The double-closure and double-centering transformations are symmetric with respect to the rows and columns of the data table. In the CFA-biplot distances of the row- and column-items from the center of the biplot are interpreted as chi-square values.

2.3 Spectral map analysis

SMA was originally developed for the display of activity spectra of chemical compounds (Lewi, 1976). The algorithm for spectral mapping is characterized by: constant weighting of rows and columns or weighting by some properly chosen weighting factor, logarithmic re-expression, double-centering, global normalization, and factor scaling using either symmetric scaling with singular values (\( \alpha = 0.5, \beta = 0.5 \)) or asymmetric scaling with unit column-variance (\( \alpha = 1, \beta = 0 \)). A further characteristic of SMA is that in the biplot the areas of the symbols are made proportional to a selected column, or to marginal row- and column-totals.
The double-centering transformation in SMA is symmetric with respect to the rows and columns of the data table. As a result of the double-centering, all absolute aspects of the data are removed. What remains are contrasts between the different rows (genes) and contrasts between the different columns (samples) of the data table. These contrasts can be expressed as ratios due to the logarithmic transformation. The contrasts can be understood as specificities of the different genes for the different samples. Conversely, they refer also to the specificities or preferences of the different samples for some of the genes. Therefore, one could state that SMA provides a visualization of the interactions between genes and samples. An advantage of SMA over CFA is that the scope of SMA is not limited to contingency tables and cross-tabulations. In addition, SMA offers the possibility to use other weighting factors than the marginal totals.

2.4 Implementation

The general algorithm as described above has been implemented in the open source language R and is available under the terms of the GNU Public License (GPL) from http://alpha.luc.ac.be/~lucp1456/.

3. Application

In a recent study, Golub et al. (1999) obtained gene expression profiles of 38 patients suffering from acute leukemia. In the following, we will refer to this data set as MIT1. Patients were diagnosed as suffering from either acute myeloid leukemia (18 patients) or acute lymphoblastic leukemia (20 patients). The latter class could further be subdivided in B-cell and T-cell classes. In addition to the initial 38 patients Golub et al. also considered a second validation sample (MIT2) of 34 patients for which the gene expression profile was determined. The original data were preprocessed as follows: genes that were
called "absent" in all samples were removed from the data sets, since these measurements are considered unreliable by the manufacturer of the technology. Negative measurements that were present in the data were set to 1. The resulting data set contained 5327 genes of the 6817 originally reported by Golub and co-workers.

The MIT1 data set will be used to compare the PCA, CFA, SMA, and weighted SMA methods with one another with regard to their ability to discover the three pathological classes and to identify genes that are related to these classes. The MIT2 data set will then be used to validate the findings that were obtained with the MIT1 data.

3.1 Principal component analysis

PCA was carried out after logarithmic re-expression of the gene expression profiles in MIT1. Since gene expression data are positively skewed and can contain large influential values, we considered a logarithmic re-expression appropriate. For the construction of the biplot (Figure 1), an asymmetric scaling with unit column-variance \((\alpha = 1, \beta = 0)\) was used to allow better visual discrimination between the different samples. This special type of factor scaling was considered optimal for extreme rectangular matrices of microarray data where variability between the genes (average variance log transformed data = 6.4) is much higher than between the different samples (average variance = 2). A consequence of unit column-variance factor scaling is that correlations and distances between samples are not represented in the biplot. However, in exploring gene expression data only patterns in the distribution of the biological samples are of direct interest. In Figure 1, the horizontal axis of the biplot, represents the first principal component that accounts for 71% of the total variance in the data. The second principal component is represented by the vertical axis of the biplot and explains only 3% of the total variance. The remaining
principal components were considered to reflect random disturbances. The horizontal axis is dominated entirely by a global component related to the size of the measurements and does not contribute any information about the differential expression of genes in the samples. Differences between biological samples are found only along the vertical axis. Only a difference between the ALL and AML groups is eminent, while data from ALL B-lineage and ALL T-lineage completely overlap one another. Furthermore, it is impossible to use the biplot for selecting genes that discriminate best between the ALL and AML classes.

Figure 1. About here

3.2 Correspondence factor analysis

The biplot obtained from CFA on the original data in MIT1 is depicted in Figure 2. The same asymmetric unit column-variance scaling was used as in PCA, to allow optimal visual discrimination of the different samples. While distances between samples are not represented in this type of scaling, the weighted distances of genes from the center are interpreted as chi-square values. In CFA sums of squares are expressed as chi-square values and the global weighted sum of squares is defined as the global chi-square. The horizontal axis of the biplot in Figure 2 accounts for 17% of the global chi-square, while the vertical axis accounts for an additional 10%. In contrast to PCA the first dominant component is not related to size. CFA highlights the differential genetic profiles of the different samples, an approach that is much more relevant to the problem. In Figure 2, genes are distributed in a funnel-like pattern and there is a clear separation between ALL and AML patients with only 2 patients that overlap one another. In contrast to PCA,
B-lineage and T-lineage classes within the ALL group are also separated from one another. It is tempting to identify a few genes that could be used in characterizing the three pathological classes. Gene probes located at the poles of the triangular-like shape should be characteristic for a given class of leukemia. However, for the two gene probes identified as the top left and right pole only a few valid measurements were made and the results depended largely on the expression level obtained in a single patient. This underscores the, in this case, less desirable sensitivity of CFA to single large values. There is also a problem with the interpretation of the numerical value of the distances between genes. Since in CFA, distances refer to chi-square values that have a meaning only for contingency tables and not for continuous data as is the case in gene expression experiments, one could seriously question the applicability of CFA in microarray data analysis.

=== Figure 2. About here ===

3.3 Spectral map analysis

In carrying out SMA, we used variable weighting for the genes and samples, with weights proportional to the mean expression levels of genes and samples respectively. Asymmetric unit column-variance was used as factor scaling in the construction of the biplot depicted in Figure 3. Genes located near the center of the map are displayed as dots, while the 0.5% (27) most distal genes are displayed as circles with areas proportional to their marginal row mean. In addition, some of these genes were labeled with their accession number. SMA, like CFA, stresses the differential genetic profile of the different samples, but in contrast to CFA relative distances can be interpreted and quanti-
fied as ratios. The three classes of leukemia are completely separated from one another and cluster around the three poles of a triangle. The horizontal axis of the map is dominated by the ratio in gene expression between the AML and ALL class and accounts for 14% of the total interaction variance. The vertical axis is dominated by the contrast between the ALL T-cell and ALL B-cell group and accounts for an additional 12% of the interaction. All of the genes that are located distal from the center could have a physiological meaning. It is noteworthy to mention that only 4 of the 27 most distal genes were among the 50 genes selected by Golub et al. (1999) to discriminate between the different classes of disease.

In a subsequent analysis (Fig. 4), we carried out a weighted SMA using the 27 genes identified in Figure 3. Since row and column variances are now comparable, the biplot was constructed using singular values \((\alpha = 0.5, \beta = 0.5)\) as the method for factor scaling. The horizontal and vertical axes explain 43% and 32% of the global interaction variance. Using only this small subset of 27 genes allows complete separation of the three pathological classes. To validate this finding, we positioned the samples obtained in the second data set (MIT2) on the biplot based on MIT1 (results not shown). AML and ALL-B classes could clearly be distinguished from one another without any overlap. There was only one possible mismatch, namely the single sample in MIT2 that was identified as ALL-T.
c. SMA as a tool to quantify differential gene expression

The maps shown in Figures 3 and 4 suggest an even further reduction of the data. Indeed, the genes located at the poles of the triangle formed by the three pathological classes almost completely represent the interaction that is present in the first factorial plane. To emphasize this point we constructed the biplot in Figure 5 using only the expression profile of the genes with accession numbers X82240, X76223, and M84526. This case of spectral map analysis where only three rows or columns are considered is also referred to as multivariate ratio analysis (MRA) and has found applications in the field of ecology (Hermy and Lewi, 1991). MRA differs from conventional SMA only by the application of asymmetric unit column-variance ($\alpha = 1, \beta = 0$) as the method for factor scaling. All 72 samples present in MIT1 and MIT2 data sets were plotted on the plane determined by these three gene-poles. In addition axes were drawn through the poles of the triangle. These axes allow quantification of the different ratios in gene expression that can be calculated from the data. A major advantage of SMA as compared to CFA is that the map permits directly the reading of the different genetic profiles of each of the samples with respect to the three characteristic genes. Figure 5 shows that samples whose expression profile is not specific for any of the genes and consequently are located at the center of the map, also have a low level of expression for all three genes, as is indicated by the extreme small areas of the corresponding squares. Genetic specificity as expressed by the differential ratio between any two genes can be substantial and can amount to a factor of 10000 or more, as is illustrated by the axis M84526 versus X76223. Furthermore, it is shown that, using only three genes the three pathological classes can be discriminated to a substantial extent.
4. Discussion

The results obtained in the previous section illustrate the impact of the different building blocks introduced in Section 2. The characteristic difference between conventional PCA on the one hand and CFA and SMA on the other hand are the operations of double-closure and double-centering. The double-closure operation in CFA eliminates the size factor that is related to the first dominant component in PCA and stresses differences among the genes and among the samples. The same effect is obtained by double-centering after logarithmic re-expression in SMA. Although, mathematically, these two operations are related, the results can differ substantially as is illustrated by the differences in the biplots obtained from CFA and SMA, respectively. Re-expressing the data to logarithms downplays very large contrasts that result from extreme outcomes. This is a desirable property for the analysis of gene expression data that typically suffer from the presence of severely outlying measurements. A drawback of the logarithmic re-expression is that contrasts at a less reliable level of gene expression are considered of equal importance as are contrasts at a more reliable level. This phenomenon can be counteracted by incorporating weights proportional to the marginal totals in the centering, normalization, and factorization building blocks leading to weighted SMA.

Our results indicate that weighted SMA is a valuable tool for the analysis of gene expression microarray data. Weighted SMA and CFA outperform conventional PCA in visualizing the data, determining clusters of samples and genes, correlating samples with gene
expression profiles, and reducing the data. An advantage of SMA over CFA is the possibility of interpreting distances as ratios, while CFA does not allow such an intuitive approach. A limitation with regard to interpretation of the spectral map would be the abundance of groupings in the different samples as is the case in some data mining applications. However, for such applications one could consider exploring subsets of the data instead of entire data sets.

Apart from the data analytic aspects of this report, it is noteworthy to mention that the three genes selected in the construction of Figure 5, could be related to leukemia. Only “Adipsin” (M84526) was also present in the set of 50 genes used by Golub et al. (1999) for class determination. This gene was also identified by an alternative analysis of the same data set reported by Chow, Moler, and Mian (2001). The second gene “T-cell leukemia/lymphoma 1A” (X82240) is reported to be involved in T-cell malignancies (Virgilio et al., 1994). The last gene probe (X76223) measures the presence of exon 4 of the gene “MAL”, which encodes a human T-cell specific proteolipid protein (Rancano et al., 1994).

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Figure 1. Biplot of the results of PCA with unit column-variance scaling ($\alpha = 1$, $\beta = 0$).

First principal component (horizontal axis) and second principal component (vertical axis) account for 71% and 3% respectively of the global variance. Small dots represent genes. The three classes of leukemia are identified by squares (AML), triangles with top up (ALL B-Cell), or triangles with top down (ALL T-cell).
Figure 2. Biplot of the results of CFA with unit column-variance scaling ($\alpha = 1, \beta = 0$).

Horizontal axis represents 17% and vertical axis 10% of the global chi-square. The three classes of leukemia are identified by squares (AML), triangles with top up (ALL B-Cell), or triangles with top down (ALL T-cell).
Figure 3. Biplot of the results of SMA weighted for marginal row and column means with unit column-variance scaling ($\alpha = 1$, $\beta = 0$). Horizontal axis represents 14% and vertical axis 12% of the total interaction variance. The three classes of leukemia are identified by squares (AML), triangles with top up (ALL B-Cell), or triangles with top down (ALL T-cell). The 0.5% most distal genes are represented as circles with areas proportional to the marginal mean, a measure of overall gene intensity. Some of the genes are labeled with their accession number.
Figure 4. Biplot of the results of SMA weighted for marginal row and column means using only the 0.5 % (27) genes most distant from the center and singular value factor scaling ($\alpha = 0.5$, $\beta = 0.5$). Horizontal axis represents 43 % and vertical axis 32 % of the total interaction variance. The three classes of leukemia are identified by squares (AML), triangles with top up (ALL B-Cell), or triangles with top down (ALL T-cell). Genes are represented as circles with areas proportional to the marginal mean, a measure of overall gene intensity. Some of the genes are labeled with their accession number.
Figure 5. All 72 samples plotted on the plane determined by the ratios between the three most extreme genes X82240, X76223, and M82546, of Figure 4. Differential gene expressions of individual patients can be read from the calibrated axes. Samples are represented as squares, with size proportional to the marginal mean and coded as: M for AML, B for ALL B-cell, and T for ALL T-cell class patients.