Sparse non-negative generalized PCA with applications to metabolomics

Genevera I. Allen1,2,* and Mirjana Maletić-Savatić1

1Department of Pediatrics-Neurology, Baylor College of Medicine, Jan and Dan Duncan Neurological Research Institute at Texas Children’s Hospital, 1250 Moursund St. Suite 1365, Houston, TX 77030 and 2Department of Statistics, Rice University, 6100 Main St. MS-138, Houston, TX 77005, USA

ABSTRACT

Motivation: Nuclear magnetic resonance (NMR) spectroscopy has been used to study mixtures of metabolites in biological samples. This technology produces a spectrum for each sample depicting the chemical shifts at which an unknown number of latent metabolites resonate. The interpretation of this data with common multivariate exploratory methods such as principal components analysis (PCA) is limited due to high-dimensionality, non-negativity of the underlying spectra and dependencies at adjacent chemical shifts.

Results: We develop a novel modification of PCA that is appropriate for analysis of NMR data, entitled Sparse Non-Negative Generalized PCA. This method yields interpretable principal components and loading vectors that select important features and directly account for both the non-negativity of the underlying spectra and dependencies at adjacent chemical shifts. Through the reanalysis of experimental NMR data on five purified neural cell types, we demonstrate the utility of our methods for dimension reduction, pattern recognition, sample exploration and feature selection. Our methods lead to the identification of novel metabolites that reflect the differences between these cell types.

Availability: www.stat.rice.edu/~gallen/software.html

Contact: gallen@rice.edu

Supplementary Information: Supplementary data are available at Bioinformatics online.

Received on June 27, 2011; revised on August 23, 2011; accepted on September 10, 2011

1 INTRODUCTION

Metabolomics, one of the newest fields within systems biology approaches to biomarker discovery in medicine, investigates an abundant pool of small molecules present in cells and tissues (Bollard et al., 2005; Hollywood et al., 2006; Holmes et al., 2008). One of the commonly used technologies for acquisition of this data is nuclear magnetic resonance (NMR) spectroscopy. It is a high-throughput technology for acquiring reproducible and resolved spectra that can be used to study the complete metabolic profile of a biological sample (Nicholson and Lindon, 2008). The spectra contain thousands of chemical resonances, which may belong to hundreds of metabolites (De Graaf, 2007). However, many metabolites resonate at multiple resonances and thus, unlike the typical DNA microarray data, different metabolite spectra overlap and introduce complexities that need to be addressed by signal processing and careful statistical analysis (Ebbels and Cavill, 2009; Weljie et al., 2006).

As understanding relationships between the set of biological samples and the underlying spectra is a challenge, principal components analysis (PCA) is commonly used for both dimension reduction and pattern recognition with NMR data (Coen et al., 2008; Dunn et al., 2005; Geodacre et al., 2004; Maletić-Savatić et al., 2008; Weckwerth and Morgenhal, 2005). In high-dimensional settings, however, it is well known that PCA can perform poorly due to the large number of irrelevant variables (Johnstone and Lu, 2009). Hence, many have proposed to incorporate sparsity into the principal component directions, thus selecting important features (Johnstone and Lu, 2009; Jolliffe et al., 2003; Shen and Huang, 2008; Zou et al., 2006). Non-negativity of the matrix factors, or principal component directions, has also been proposed in a number of settings to improve interpretability of the factors (Lee and Seung, 1999; Sajda et al., 2004). Several recent papers have combined these concepts to encourage both sparsity and non-negativity into the model (Hoyer, 2004; Kim and Park, 2007; Zass and Shashua, 2007).

In this article, we make the following statistical contributions: (i) propose a framework for incorporating sparsity, known structural dependencies and non-negativity into the principal component (PC) loadings and (ii) develop a fast, computationally efficient algorithm to compute these in high-dimensional settings. This work is presented in Section 2. Then, in Section 3, we evaluate the performance of our methods on real NMR data. We also demonstrate how to interpret the PC loadings to understand important biological patterns and identify candidate metabolites. In Section 4, we conclude with a summary of the implications of our work and future areas of research.

2 METHODS

We introduce a framework for PCA that incorporates structural dependencies, sparsity and non-negativity to better understand relationships between the samples and recognize patterns among the variables. 2.1 Review: generalized PCA

Recently, Allen et al. (2011) introduced a new matrix decomposition, the Generalized Least Squares Matrix Decomposition (GMD), and showed how this decomposition can be used to generalized PCA by directly incorporating known structural information or dependencies. Here, we...
review the Generalized PCA (GPCA) problem and specifically discuss its utility in the context of spectroscopy data.

We observe data, \( X \in \mathbb{R}^{n \times p} \), for \( n \) samples and \( p \) variables that has previously been normalized. (With NMR data, this includes baseline correction, normalizing by the integral of the spectrum and standardizing the variables at each ppm.) Let \( \mathbb{R}^{n \times p} \) be a positive semi-definite matrix called the quadratic operator that captures the noise structure in the data. Then, GPCA seeks the linear combination of variables maximizing the sample variance in the inner product space induced by \( \mathbf{R} \):

\[
\begin{align*}
\text{maximize} & \quad v_i' \mathbf{R} X' \mathbf{R} v_i \\
\text{subject to} & \quad v_i' \mathbf{R} v_i = 1 \quad \text{and} \quad v_i' \mathbf{R} v_1 = 0 \quad \forall k' < k. \\
\end{align*}
\]

(1)

The \( k \)-th GPC is \( z_k = \mathbf{R} v_k \). If \( \mathbf{R} = \mathbf{I} \), then we have the standard PCA optimization problem. Additionally, Allen et al. (2011) have shown that an extension of the power method for computing eigenvectors can be used to calculate these GPCs.

GPCA can be used to directly account for dependencies between adjacent variables in the spectra. The quadratic operator, \( \mathbf{R} \), behaves like an inverse covariance matrix of multivariate normal data (Allen et al., 2011). We can let \( \mathbf{R} \) encode the inverse covariance of dependencies or structure in the data that do not contribute, and are independent of the signal of interest. The resulting GPCA solution can be interpreted as a decomposition of the resulting GPCA solution can be interpreted as a decomposition of the

2.1 Sparse non-negative GPCA

We introduce the single-factor sparse non-negative GPCA problem. \( \mathbf{R} \) is biconcave, meaning that it is concave in \( u \) with \( \mathbf{R} \) fixed and in \( \mathbf{R} \) with \( u \) fixed. This leads to a simple maximization strategy that is guaranteed to increase the objective function while converging to a minimum value: alternate maximizing with respect to \( u \) and \( \mathbf{R} \). These coordinate-wise maximization problems turn out to have a simple solution.

Proposition 1. Let \( k \) be the minimizers of the following:

\[
\begin{align*}
\text{minimize} & \quad \frac{1}{2} \| \mathbf{X} u - \hat{v}_j \|_2^2 - \lambda \| u \|_1 \quad \text{subject to} \quad v_j \geq 0. \\
\end{align*}
\]

(3)

Then, the coordinate updates, \( u^* \) and \( v^* \), maximizing the single-factor sparse non-negative GPCA problem, (2), are given by:

\[
\begin{align*}
& v^* = \begin{cases} 
\hat{v}_j / \| u^* \|_2 & \text{if } |\hat{v}_j| / \| u^* \|_2 > \lambda / \| u^* \|_1 \\
0 & \text{otherwise,}
\end{cases} \\
& u^* = \mathbf{R} v^* / \| \mathbf{R} v^* \|_2
\end{align*}
\]

(All proofs are given in the Supplementary Materials).

The solution to the single-factor sparse non-negative GPCA problem, (2), can be obtained by solving a simple lasso penalized non-negative regression optimization problem. This non-negative regression problem in turn can be solved via a fast coordinate descent algorithm:

Proposition 2. The solution to (3) can be obtained via coordinate descent with updates \( \hat{u}_j = \begin{cases} u_j & \text{if } u_j > 0 \\
\hat{v}_j / \| u^* \|_2 & \text{otherwise} \end{cases} \), where \( \hat{u}_j \) denotes the row elements of column \( j \) of \( \mathbf{R} \) and \( \hat{v}_j \) denotes the positive part.

This coordinate descent algorithm is related to the fast shrinking algorithms of Friedman et al. (2010), and the speed can be further improved by employing active set learning and warm starts. We note that this algorithmic approach is a major improvement in terms of computational efficiency over the least angle-based approach to the non-negative lasso of Renard et al. (2008).

2.2.2 Algorithm

We have presented an optimization problem and solution to the single-factor sparse non-negative GPCA problem, and we are also interested in extracting multiple components. Then, we employ a greedy approach to estimating multiple components that is closely related to the power method algorithm for computing eigenvectors. This algorithm is summarized in Algorithm 1.

The sparse non-negative GPCA algorithm begins with the standardized data and computes the first component by solving the single-factor problem via coordinate descent. Subsequent components are calculated by solving the single-factor problem for the residual where the previously computed outer product has been removed. Each component is calculated in a greedy manner and is hence conditional on the previously estimated components. Thus, the components are not necessarily ordered in terms of the amount of variance they explain. This approach is common among existing methods for sparse PCA (Allen et al., 2011; Les et al., 2010; Shen and Huang, 2008; Witten et al., 2009; Zou et al., 2006). As the dominant operation in our algorithm is solving a non-negative lasso problem, the computational complexity is \( \mathcal{O}(n^2) \). While traditional PCA methods may be faster to compute, our algorithm requires comparable computational time to existing sparse and non-negative PCA methods (Shen and Huang, 2008; Zass and Shashua, 2007).
We seek a data-driven mechanism for selecting the amount of sparsity in the variance explained by each component are altered. When using PCA methods for differing penalty parameters for each component.

### 2.2.4 Amount of variance explained

...the formulas for calculating the proportion of variance explained by the k-th GPC is 

\[
\hat{v}_k = \frac{v_k^T R v_k}{v_k^T v_k}.
\]

Then, the cumulative proportion of variance explained by the k-th sparse non-negative GPC is 

\[
\sum_{i=1}^{k} \frac{v_i^T R v_i}{v_i^T v_i}.
\]

Note that the proportion of variance explained by individual sparse non-negative GPCs can be found by taking the differences of the cumulative proportion explained. Thus, the proportion of variance explained by our methods can be interpreted as the ratio of the R-norm projected sample variance of the k-th linear projection relative to the total variance of the data in the R-norm. Notice that as the sparse non-negative GPC factors are not constrained to be orthogonal, the sample variance explained must be adjusted for possible correlations among the factors as discussed in Shen and Huang (2008). Given these results, we can compare our methods to traditional PCA and sparse PCA methods in terms of the variance explained and dimension reduction.

### 3 RESULTS

We evaluate the utility of GPCA and Sparse Non-Negative GPCA for metabolomics through comparisons on real NMR data. (Simulation studies are given in the Supplementary Materials.) We use a dataset with 27 samples acquired by in vitro 1D H-NMR on five neural cell types: neurons, neural stem cells, microglia, astrocytes and oligodendrocytes (Manganas et al., 2007). [For methodology used on cell culturing, see Manganas et al. (2007)] The data are preprocessed in the traditional manner (Dunn et al., 2005): after acquisition, functional spectra is discretized by binning variables into bins of size 0.04 ppms yielding a total of 2394 variables. For each sample, the spectra are baseline corrected and normalized to their integral. Before applying multivariate techniques, the variables are standardized to have mean zero and variance one.

We compare our GPCA method to traditional PCA and our sparse non-negative GPCA method to sparse non-negative PCA. The later is implemented via Algorithm 1 by setting \( R = I \). The BIC method is used to select penalty parameters for both sparse PCA methods and the first 15 PCs are calculated for all methods. For the GPCA methods, the quadratic operator, \( R \), was taken to be a Gaussian kernel smoother with smoothing parameter, \( \sigma = 20 \). Five possible values of \( \gamma \) were considered, \( \gamma = 5, 10, 15, 20, 25 \). With \( \gamma \) chosen to explain the most sample variance.

In Figure 1, we compare scatter plots of the normalized sample PCs for the four methods. Notice that the scatter plots of all methods exhibit clustering of the neuron and neural stem cell samples, while the other cell types are more scattered. Sparse methods and especially sparse non-negative GPCA, however, cluster the remaining cell types better, illustrating the utility of incorporating sparsity in high-dimensional data analysis.

Next, we compare the methods in terms of dimension reduction in Figure 2. As sparse PCA methods naturally explain less sample variance than PCA methods, we compare the two sets of methods separately. Also note that as sparse PCA methods calculate components in a greedy manner, they are not necessarily ordered in terms of how much variance they explain. Overall, by incorporating the known structure of spectroscopy data into the PCA problem, the GPCA methods explain a larger portion of the sample variance. Thus, the reduction of dimensions for GPCA methods is greater. This behavior is especially pronounced for the sparse non-negative methods where seven PCs explain over 90% of the variance for sparse non-negative GPCA, while 15 PCs are needed to explain the same amount of variance for sparse non-negative PCA. Thus, sparse non-negative GPCA provides over 50% more dimension reduction than sparse non-negative PCA. GPCA methods demonstrate a clear advantage over traditional PCA methods in terms of variance explained and dimension reduction.
Fig. 1. Scatter plots of normalized sample PCs for the neural cell types data. Results from PCA, GPCA, Sparse Non-Negative PCA (SPCA) and Sparse Non-Negative GPCA (SGPCA) are compared for the five neural cell types. Sparse methods (bottom rows) demonstrate clearer separation of samples from different cell types.

Fig. 2. Amount of variance explained by the PCs for the five neural cell type data. Comparison of the percentage of variance explained by individual PCs (top panel) and cumulative percentage of variance explained (bottom) between PCA and GPCA (left), and sparse non-negative PCA and sparse non-negative GPCA (right). GPCA methods explain larger proportions of the sample variance.

A major motivation of our work is to incorporate feature selection into the traditional PCA framework and assess its utility for NMR data. We compare the degree of sparsity seen in the PCs for the sparse non-negative PCA and GPCA methods in Figure 3. By directly accounting for the dependencies at adjacent chemical shifts, sparse non-negative GPCA gives a greater degree of sparsity, yielding a more parsimonious model. The GPCA method also explains more of the variance in the data with fewer features selected, an important attribute. Greater sparsity means that one needs to consider fewer peaks when explaining the patterns in the data. Also, a parsimonious
Sparse non-negative GPCA loadings and sample PC heatmaps for the first seven PCs, which explain over 90% of the sample variance. Scaled PC loadings are superimposed on the average scaled spectra of neural stem cells, neurons, microglia and ‘Glia’, which includes oligodendrocytes and astrocytes. Sparse non-negative GPCA loadings reveal important patterns across the samples and spikes in the loadings denote the location of peaks that vary greatly across the samples. For example, PC3 exhibits peaks that have higher intensities in neural stem cells, while the peaks selected by PC5 have higher concentrations in microglia.

PC loading vector indicates that more irrelevant variables have been discarded from the model. As sparse non-negative PCA does not incorporate structural information, many more variables are selected as the method tries to explain both the dependencies between neighboring chemical shifts and the biological variation. By directly accounting for these spatial dependencies, however, sparse non-negative GPCA is free to select features that explain the biological variation in the samples. Overall, these results indicate that sparse non-negative GPCA outperforms PCA, GPCA and sparse non-negative PCA in terms of sample exploration, dimension reduction and feature selection.

Sparse non-negative GPCA can be used to understand important biological patterns in the NMR data. Figure 4 gives the sparse non-negative GPCA loadings for the first seven sparse non-negative GPCs which explain over 90% of the variance in the data. Along with these loadings, we give heatmaps of the sample PCs to show how each of the samples contribute to the patterns seen in the loading vectors. The loading vectors are scaled and superimposed on the mean spectra from neurons, neural stem cells, microglia and ‘Glia’, which includes astrocytes and oligodendrocytes. (Plots of the loading vectors for PCA, GPCA and sparse non-negative PCA are given in the Supplementary Materials.)
Table 1. Locations in parts per million (ppm) of the most important peaks identified by the first seven sparse non-negative GPCA loadings

<table>
<thead>
<tr>
<th>Peak location (ppm)</th>
<th>Cell types</th>
<th>Metabolites</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.96</td>
<td>Neuron, microglia</td>
<td></td>
</tr>
<tr>
<td>1.19</td>
<td>Microglia, neuron</td>
<td></td>
</tr>
<tr>
<td>1.28</td>
<td>Neural stem cell, oligodendrocyte</td>
<td>Lipid moiety</td>
</tr>
<tr>
<td>1.48</td>
<td>Oligodendrocyte</td>
<td></td>
</tr>
<tr>
<td>2.02</td>
<td>Neuron</td>
<td>NAA</td>
</tr>
<tr>
<td>2.65</td>
<td>Astrocyte</td>
<td></td>
</tr>
<tr>
<td>3.01</td>
<td>Neuron</td>
<td></td>
</tr>
<tr>
<td>3.04</td>
<td>Oligodendrocyte</td>
<td>Creatine</td>
</tr>
<tr>
<td>3.23</td>
<td>Oligodendrocyte, neuron, astrocyte</td>
<td>Choline</td>
</tr>
<tr>
<td>3.43</td>
<td>Oligodendrocyte</td>
<td></td>
</tr>
<tr>
<td>3.66</td>
<td>Microglia</td>
<td></td>
</tr>
</tbody>
</table>

Boldfaced locations denote peaks with especially strong signals as indicated by the loading vector. As the dependencies of the noise must be known to construct the quadratic operator, our methods can be used to find patterns in multidimensional metabolomics data. Not only does our approach provide more feature selection than competing methods, but also yields easily interpretable results that lead to understanding of important biological patterns in the spectra.

4 DISCUSSION

We have presented a framework for incorporating structural dependencies, sparsity and non-negativity into PCA. By comparing our techniques to traditional PCA methods on real NMR data, we have demonstrated the many advantages of our methods. Future areas of research are to extend our framework to supervised multivariate analysis techniques such as partial least squares and linear discriminant to better classify NMR samples.

While we have demonstrated our methods on 1D H-NMR spectroscopy, our approach can be applied to many other high-throughput metabolomics technologies. Mass spectrometry and other spectroscopy techniques also produce a spectrum of non-negative variables. Additionally, many researchers employ multidimensional spectroscopy to further identify metabolites in a sample (De Graaf, 2007). In this data, each sample consists of a matrix of spectroscopy variables. Sparse non-negative GPCA can be applied to this multidimensional data in a straightforward manner by vectorizing the matrix of variables and employing a 2D kernel smoother over the lattice of variables. As a future area of research, one can also extend our methods to tensors or higher order PCA to find patterns and achieve dimension reduction for this multidimensional metabolomics data.

In addition to metabolomics data, our methods are general and hence applicable to a variety of other structured biomedical data. As the dependencies of the noise must be known to construct the quadratic operator, our methods can be used to find patterns in data where these noise dependencies are well established. Possible further applications of our methods then include copy number variation and methylation data in which variables strongly depend on known chromosomal location, and microscopy, neuroimaging and other bio-medical imaging data in which pixels are spatially correlated with adjacent pixels.

In conclusion, we have developed a novel modification of PCA particularly suited to the challenges associated with analyzing NMR data. While our methods show numerous advantages in the analysis of metabolomics data, there are still many open research problems and potential extensions related to our work.

ACKNOWLEDGEMENTS

The authors would like to thank Han Xu, Yanli Chen, Dr Li-Hua Ma, Dr Marina Vannucci and Dr Juan Botas for helpful discussions related to this work.

Funding: National Institute of Neurological Disorders and Stroke (R21NS058785-1 and K08NS0044276); McKnight Endowment Fund; DANA Foundation; Lisa and Robert Lourie Foundation and...
the NIH Intellectual and Developmental Disabilities Research Grant (P30HD020641) to M.M.-S.

Conflict of Interest: none declared

REFERENCES


