Research Article

Group Factor Analysis for Alzheimer’s Disease

Wei-Chen Cheng, Philip E. Cheng, and Michelle Liou

Institute of Statistical Science, Academia Sinica, Taipei 11529, Taiwan

Correspondence should be addressed to Wei-Chen Cheng; wccheng@stat.sinica.edu.tw

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For any neuroimaging study in an institute, brain images are normally acquired from healthy controls and patients using a single track of protocol. Traditionally, the factor analysis procedure analyzes image data for healthy controls and patients either together or separately. The former unifies the factor pattern across subjects and the latter deals with measurement errors individually. This paper proposes a group factor analysis model for neuroimaging applications by assigning separate factor patterns to control and patient groups. The clinical diagnosis information is used for categorizing subjects into groups in the analysis procedure. The proposed method allows different groups of subjects to share a common covariance matrix of measurement errors. The empirical results show that the proposed method provides more reasonable factor scores and patterns and is more suitable for medical research based on image data as compared with the conventional factor analysis model.

1. Introduction

Modern medical imaging techniques are capable of measuring human brain in vivo [1]. For instance, magnetic resonance (MR) imaging measures nuclei of atoms, and positron emission tomography detects the positron-emitting radionuclides to construct three-dimensional images. The imaging procedures are designed and settled before medical or cognitive experiments. Once the protocol is established, the laboratory and the hospital begin to recruit a variety of subjects of interest into experimental sessions. Errors resulting from individual scans are actually generated from common sources, such as the scanner, protocol, and software. Initial classification of subjects into groups can be realized by using clinical diagnosis, which may be uncertain to some extent, provided by physicians along with subjects’ anamnesis.

Conventional factor analysis [2] models reduce high-dimensional data into a few latent variables and assume that data $\mathbf{x}$ were generated by a set of unobserved independent unit-variance Gaussian source $\mathbf{f}$ plus uncorrelated zero-mean Gaussian random noise $\mathbf{u}$, $\mathbf{x} = \mathbf{Lf} + \mathbf{u}$, where $\mathbf{L}$ is the factor loading matrix. The sample covariance of $\mathbf{x}$ can be expressed as $LL^T + \Psi$, where $\Psi$ is a diagonal covariance matrix of random noises. The goal of factor analysis is to find $\mathbf{L}$ and $\Psi$ that maximally fit the sample covariance [3–5]. The EM algorithm was proposed to estimate the matrices [6]. Factor analysis is commonly applied to the dataset as a whole or to different groups of data separately, which may result in factor patterns hard to interpret and limit the potential use of the method in a wider range of medical applications. In this study, we propose a mixture factor analysis model (MFAM) to assign a common covariance matrix of noises or measurement errors to different groups of subjects but to allow individual groups having their own latent structures. In the empirical application, we analyzed an Alzheimer’s disease (AD) dataset by first extracting the volumetric information from MR anatomical images for both healthy controls and the patients suffering either AD or mild cognitive impairment, followed by applying the proposed MFAM to the volumetric data.

2. Material and Method

2.1. The Model. Let $M$ be the number of subject groups. To find multiple sets of factor loadings, $\{L_j; j = 1, \ldots, M\}$, with
Figure 1: The plots of means and standard deviations for the three groups in different subcortical structures.
the $f$ scores distributed as Gaussian within each group, the data vector can be decomposed into a linear combination of factor loadings for each group \[7, 8\], that is, $L_j \in \mathbb{R}^{D \times K}$,

$$
x = \sum_{j} \pi_j (\mu_j + L_j x | w_j) + u, 
$$

where $x$ is $D$-dimensional and each factor scores $f | w_j$ has $K$ variables, that is, $f \in \mathbb{R}^K$. The parameter $\pi$ is associated with the proportion of subjects in the $j$th group, $\pi_j = p(w_j)$. The indicator variable $w$ is one, $w_j = 1$, when the data belongs to $j$th group, otherwise $w$ is set to zero, $w_j = 0$. The formula (1) using $\pi$ introduces the main difference from previous mixture models of factor analysis. The data vector $x$ need not be centered and the mean of the $j$th group data is $\mu_j$. The covariance matrix of residuals $u$ is a diagonal matrix $\Psi = \text{diag}[\Psi_1, \Psi_2, \ldots, \Psi_D]$. The data distribution can be expressed as

$$
P(x) = \sum_{j=1}^{M} P(x | f, w_j) P(f | w_j) P(w_j) df. 
$$

In this work, capitalized $P$ denotes the probability function of a vector or a matrix and lowercase $p$ denotes the probability function of a scalar. The factor scores are assumed to be distributed as Gaussian

$$
P(f | w_j) = N(0, I), \quad \forall j. 
$$

The notation $I$ is the identity matrix of order $D$. The distribution of data $x$ in each group is given by

$$
P(x | f, w_j) = N(\mu_j + L_j f_j, \Psi). 
$$

Based on the MFAM (2), the likelihood function $Q$ is as follows:

$$
Q = E \left[ \prod_{i=1}^{N} \prod_{j=1}^{M} \left\{ (2\pi)^{-D/2} |\Psi|^{-1/2} \exp \left\{ -\frac{1}{2} (x_i - \mu_j - L_j f_j)^T \Psi^{-1} \right\} \right\} \right]^{w_i}, 
$$

where $E$ denotes the expectation. The $N$ is the number of data vectors (subjects) with subscript $i$ for the $i$th subject. We need to compute the expectation of the variables,

$$
E (w_j | x_i) = E (w_j | x_i) E (f_j | w_j, x_i). 
$$

To estimate $Q$ in (5), the posterior probability of the $j$th group is calculated as

$$
P(w_j | x) = \frac{P(x | w_j) P(w_j)}{P(x)} 
$$

$$
= \frac{\pi_j N(x - \mu_j, L_j L_j^T + \Psi)}{\sum_{i} \pi_i N(x - \mu_i, L_i L_i^T + \Psi)}, 
$$

where the probability of $x$ given $w_j$ is

$$
P(x | w_j) = N(x - \mu_j, L_j L_j^T + \Psi). 
$$
where
\[
E(\tilde{f}_i | \tilde{f}_i, x_i) = R_i^{-1} L_j^T \Psi^{-1} (x_i - \mu_j),
\] (13)
and \( \tilde{L}_i = [L_j \mu_j] \). The expected log likelihood function can be expressed as
\[
E[\log Q] = \exp \left( \sum_{i,j} \int f_i^T L_j^T \Psi^{-1} x_i \right) \frac{1}{2} \left( x_i - \tilde{L}_j \tilde{f}_j \right)^T \Psi^{-1} \left( x_i - \tilde{L}_j \tilde{f}_j \right)
\] (14)
To maximize \( Q \) with respect to \( \tilde{L}_j \), we equate the derivative of (14) to zero,
\[
\frac{\partial \log E[Q]}{\partial \tilde{L}_j} = - \sum_i h_{ij} \Psi^{-1} x_i E[\tilde{f}_i | x_i, w_j]^T + h_{ij} \Psi^{-1} \tilde{L}_j E[\tilde{f}_i | x_i, w_j]^T = 0
\] (15)
where
\[
E[\tilde{f}_i | x_i, w_j] = \left[ E[\tilde{f}_i | x_i, w_j] \right]^T.
\] (16)
Equation (1) can be written
\[ x = \sum_{j} \pi_j (\mu_j + L_j \times (HH^T)f | w_j) + u = \sum_{j} \pi_j (\mu_j + L^*_j \times f^* | w_j) + u, \quad (21) \]

Scores in (6) and the second moment of the scores, the E-step, the algorithm computes the expectation of the factor loadings and the factors scores can capture the latent factors of different disease groups. The proposed model also carries the same indeterminacy problem associated with factor patterns; that is there exist numerous orthogonal transformations to rotate the matrix of factor loadings without changing the maximum of Q [9]. Considering H be any K x K orthogonal matrix, \( HH^T = H^TH = I \). Equation (1) can be written

Substituting (15) for \( \bar{L}_j \) and making constraints on the diagonal of \( \Psi \), we obtain

\[ \Psi = \frac{1}{N} \text{diag} \left( \sum_{i,j} h_{ij} (x_i - \bar{L}_j E \left( f_i \mid x_i, w_j \right)) x_i^T \right). \quad (20) \]

The prior probability \( p(w_j) \) should be proportional to the clinical diagnosis such that the estimation of the factor loadings and the factor scores can capture the latent factors of different disease groups. The proposed model also carries the same indeterminacy problem associated with factor patterns; that is there exist numerous orthogonal transformations to rotate the matrix of factor loadings without changing the maximum of Q [9]. Considering H be any K x K orthogonal matrix, \( HH^T = H^TH = I \). Equation (1) can be written

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\[ x = \sum_{j} \pi_j (\mu_j + L_j \times (HH^T) \times f \mid w_j) + u \]

All the variables are estimated by the EM algorithm. In the E-step, the algorithm computes the expectation of the factor scores in (6) and the second moment of the scores,

\[ E \left[ w_j f_i^T \mid x_i \right] = E \left[ w_j \mid x_i \right] E \left[ f_i^T \mid w_j, x_i \right], \quad (17) \]

by

\[ E \left[ w_j f_i^T \mid x_i \right] = h_{ij} \text{Cov} \left( f_i \mid w_j, x_i \right) + h_{ij} E \left[ f_i \mid w_j, x_i \right] E \left[ f_i \mid w_j, x_i \right]^T. \quad (18) \]

The covariance matrix of residual, \( \Psi \), can be estimated by its inverse matrix,

\[ \frac{\partial Q}{\partial \Psi^{-1}} = -\frac{N}{2} \Psi - \sum_{i,j} h_{ij} x_i x_i^T + h_{ij} E \left[ f_i \mid x_i, w_j \right] E \left[ f_i \mid x_i, w_j \right]^T \]

\[ = \frac{1}{2} h_{ij} E \left[ f_i^T \mid x_i, w_j \right] E \left[ f_i \mid x_i, w_j \right]^T L_j = 0. \quad (19) \]
where $L^*_j = L_jw_j$, and $\mathbf{f}^* | w_j = H^T \times f | w_j$. The assumption, $\mathbf{f}^* | w_j \sim N(0, I)$, is kept. The covariance of $\mathbf{x}$ is $L_j(L_j)^T + \Psi = L_jH^TH^T + \Psi = L_jL_j^T + \Psi$, which remains the same. Therefore, there are infinite equivalent solutions to satisfy the maximum of (5). Imposing reasonable constraints to identify a set of model parameters can make the factor loadings scientifically interpretable. A widely used approach for a simple factor structure is realized by setting some factor loadings to hypothetical values such as zeros.

The permutation and changing the sign of columns in the factor loading matrix with factor scores does not affect the model at all and the algorithm will yield the same solution. In order to realize consistent, interpretable, and comparable results, we suggest to recursively test all combinations to find the one of them that has the highest similarity among results, we suggest to recursively test all combinations to find a coherent factor loading matrix with factor scores does not affect the matching and the Hungarian algorithm can find the match in time.

<table>
<thead>
<tr>
<th>Left thalamus proper</th>
<th>NL</th>
<th>vAD</th>
<th>AD</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.22</td>
<td>-0.86</td>
<td>-0.87</td>
<td></td>
</tr>
<tr>
<td>Left caudate</td>
<td>0.13</td>
<td>-0.36</td>
<td>-0.50</td>
</tr>
<tr>
<td>Left putamen</td>
<td>0.23</td>
<td>0.60</td>
<td>1.01</td>
</tr>
<tr>
<td>Left pallidum</td>
<td>0.04</td>
<td>0.01</td>
<td>-0.45</td>
</tr>
<tr>
<td>Brain stem/ventricle</td>
<td>-0.04</td>
<td>0.20</td>
<td>-0.00</td>
</tr>
<tr>
<td>Left hippocampus</td>
<td>0.25</td>
<td>-0.67</td>
<td>-1.03</td>
</tr>
<tr>
<td>Left amygdala</td>
<td>0.10</td>
<td>-0.24</td>
<td>-0.47</td>
</tr>
<tr>
<td>Left accumbens area</td>
<td>0.16</td>
<td>-0.44</td>
<td>-0.67</td>
</tr>
<tr>
<td>Right thalamus proper</td>
<td>0.23</td>
<td>-0.63</td>
<td>-0.93</td>
</tr>
<tr>
<td>Right caudate</td>
<td>0.14</td>
<td>-0.42</td>
<td>-0.50</td>
</tr>
<tr>
<td>Right putamen</td>
<td>0.20</td>
<td>-0.55</td>
<td>-0.84</td>
</tr>
<tr>
<td>Right pallidum</td>
<td>0.11</td>
<td>-0.28</td>
<td>-0.54</td>
</tr>
<tr>
<td>Right hippocampus</td>
<td>0.25</td>
<td>-0.68</td>
<td>-1.08</td>
</tr>
<tr>
<td>Right amygdala</td>
<td>0.11</td>
<td>-0.27</td>
<td>-0.58</td>
</tr>
<tr>
<td>Right accumbens area</td>
<td>0.22</td>
<td>-0.60</td>
<td>-0.95</td>
</tr>
</tbody>
</table>

**Figure 6:** The cluster means of the three groups.

Table 1: The ANOVA results for the three groups in different subcortical structures.

<table>
<thead>
<tr>
<th>Structure</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left thalamus proper</td>
<td>$2.0183 \times 10^{-15}$</td>
</tr>
<tr>
<td>Left caudate</td>
<td>$1.3736 \times 10^{-5}$</td>
</tr>
<tr>
<td>Left putamen</td>
<td>$6.9145 \times 10^{-18}$</td>
</tr>
<tr>
<td>Left pallidum</td>
<td>0.0351</td>
</tr>
<tr>
<td>Left hippocampus</td>
<td>0.1707</td>
</tr>
<tr>
<td>Brain stem and ventricle</td>
<td>0.1351</td>
</tr>
<tr>
<td>Left amygdala</td>
<td>$9.5999 \times 10^{-4}$</td>
</tr>
<tr>
<td>Left accumbens area</td>
<td>$1.0958 \times 10^{-4}$</td>
</tr>
<tr>
<td>Right thalamus proper</td>
<td>$2.6864 \times 10^{-17}$</td>
</tr>
<tr>
<td>Right caudate</td>
<td>$1.6556 \times 10^{-8}$</td>
</tr>
<tr>
<td>Right putamen</td>
<td>$1.6447 \times 10^{-13}$</td>
</tr>
<tr>
<td>Right pallidum</td>
<td>$8.8771 \times 10^{-3}$</td>
</tr>
<tr>
<td>Right hippocampus</td>
<td>$3.5498 \times 10^{-22}$</td>
</tr>
<tr>
<td>Right amygdala</td>
<td>$6.5971 \times 10^{-5}$</td>
</tr>
<tr>
<td>Right accumbens area</td>
<td>$8.7323 \times 10^{-17}$</td>
</tr>
</tbody>
</table>

2.2. Data Description. The T1-weighted MR images of 416 subjects were downloaded from the Open Access Series of Imaging Studies [10], which is publically available for analysis. All the T1-weighted images were acquired on a 1.5-T Siemens Vision scanner. Among all 416 subjects, there are 316 normal subjects (average age: 45.09 ± 23.90), 70 subjects who have been clinically diagnosed with very mild AD (average age: 76.21 ± 7.19), and 30 are with moderate AD (average age: 78.03 ± 6.91). The proportions of each type of subject are $\pi = [75.96\%, 16.83\%, 7.21\%]^T$. Multiple intrasession acquisitions provide extremely high signal-to-noise ratio, making the data amenable to our analysis. The available images were provided skull stripped, gain field corrected, and registered to the atlas space of Talairach and Tournoux [11] with a 12-parameter rigid affine transform. The resolution of the images is $176 \times 208 \times 176$. The number of voxels, which is more than six million, is much larger than the number of subjects. We extracted the clinically and psychologically interested regions instead of processing whole voxels in the image. The subcortical structures are extracted by the segmentation method [12] which uses manually labeled image data as priori information for a Bayesian framework that utilizes the principles of the active shape and appearance models. The size of a subcortical region was calculated by multiplying the voxel size and the number of voxels in the region. Fifteen subcortical regions were successfully extracted.

According to a demographic study by the National Institute on Aging and Alzheimer’s Association based on the data collected in the Chicago Health and Aging Project, the prevalence of dementia among individuals aged 71 and older was 13.9%, and AD (Alzheimer’s disease) was 9.7% [13]. The study was based on a sample of 856 individuals. The $\pi$ was estimated to be $[76.4\%, 13.9\%, 9.7\%]^T$ which is close to the statistics in our empirical data. The data vector of each subject had fifteen dimensions, each corresponding to the volume size of a subcortical structure divided by the estimated total intracranial volume. The average size of all of the intracranial volume is 1480.5 cm$^3$. The intracranial volume is estimated by the linear registration from a manually measured intracranial volume of a standard brain to the individual brain [14].

The analysis of variance (ANOVA) of the data for each structure were calculated and shown in Table 1 and Figure 1. The smaller $p$ value indicates high probability of inequality of the structure size among the three groups.

We subtracted the mean from the data and used the remainder for analysis. Using the covariance matrix of the data to estimate the factor scores would cause that a few structures dominate the factor loadings; therefore, we divided each dimension by its standard deviation to compel each of them to have unit variance. After the algorithm converged, we used varimax rotation [15], which transforms the loadings
### Table 2: The normality test of factor score by Kolmogorov-Smirnov test.

<table>
<thead>
<tr>
<th></th>
<th>Proposed method</th>
<th>Traditional method</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Factor 1</td>
<td>Factor 2</td>
</tr>
<tr>
<td>NL</td>
<td>0.20</td>
<td>0.53</td>
</tr>
<tr>
<td>vAD</td>
<td>0.00</td>
<td>0.29</td>
</tr>
<tr>
<td>AD</td>
<td>0.14</td>
<td>0.81</td>
</tr>
</tbody>
</table>

3. Results

Figure 3 shows the trend of the likelihood climbs as adding the number of factors in the analysis. In the scree plot in Figure 2, three eigenvalues of the covariance matrix of the whole dataset are greater than one and the cumulative percentage of variance from the largest three eigenvalues reaches 78%. Thus we set $K = 3$ in this analysis.

The factor loadings for the three groups are shown in Figure 4, in which the vertical axis marks the fifteen regions. The vAD denotes the group of very mild AD. The log likelihood in (5) after the algorithm converges is $-5475.814$. The loading of structures has symmetric property and usually the right and left structures have similar loadings. Using the factor loadings to estimate $\pi$ and the expected group information given $x$ by (9), we obtain the adjusted and turned proportions as $[0.8289, 0.7976, 0.1491]^T$. This may suggest the underlying variation among different groups of subject and need further investigation. Note that the reestimated proportion $h_{ij}$ is not binary anymore.

We show the results of conventional factor analysis in Figure 5 as a comparison. The program run on the mild AD patients in the dataset cannot achieve reproducible results; therefore, the quantity of mild AD’s results in Figure 5 varies from time to time. The analysis for the AD group cannot converge, however the factor loadings are reproducible. The distance of whole factor loading matrices among the three groups for conventional factor analysis is 5.28 while the proposed method is 4.54. The correlation of the three-factor loading matrix estimated by conventional factor analysis methods is $[C_{12}, C_{13}, C_{23}] = [0.3879, 0.1698, 0.3633]$. The correlation by proposed method is $[0.5388, 0.5564, 0.4986]$. Table 2 lists the $p$ values for all factors by the Kolmogorov-Smirnov test [16] on the factor scores against a Gaussian distribution. The test examines the difference between input distributions and a Gaussian distribution. The smaller the $p$ values, the more strongly the test rejects the Gaussian assumption. The algorithm tries to find factors with normally distributed factor score, hence large $p$ indicates the factor fit well to Gaussian distribution.

The means (centers) of the clusters are shown in Figure 6. The means are near the origin and include negative value because the data are standardized by the subtraction of the overall mean of the data in the preprocess stage. The yellow color in the first column indicates that healthy controls have larger sizes in subcortical structures, and the second and the third column indicate that the patients have smaller sizes in different subcortical regions in general. The AD patient has very small thalamus, putamen, and hippocampus. The hippocampus is related to memory and learning. The putamen is a structure involved in the regulation of voluntary movement. The abnormal pallidum in Figure 4 can cause movement disorders. Figure 7 shows the associations of first factor scores with the score of minimental state examination (MMSE) by both methods.

4. Conclusions

The proposed method finds closer and more correlated factor loadings than the conventional method because it considers the same error matrix for different groups of data. The result of conventional factor analysis having higher normality.
for AD patients than normal subjects is less convincing. Conventional factor analysis that decomposes the observed data together intermixes the latent factors. Taking the data apart will misseparate the noise. This work proposed using a mixture model of factor analysis method for neurodegenerative disease research by showing highly correlated factor loading across different groups of subjects and together with proper normality of the factor scores.

Acknowledgment

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References
