Modeling Multiple Correlated Functional Outcomes with Spatially Heterogeneous Shape Characteristics

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Summary: We propose a copula-based approach for analyzing functional data with multiple outcomes exhibiting spatially heterogeneous shape characteristics. To accommodate the possibly large number of parameters in multiple outcome data, parameter estimation is performed in two steps: first, the parameters for the marginal distributions are estimated using the skew t family, and then the dependence structure both within and across outcomes is estimated using a Gaussian copula. We develop an estimation algorithm for the dependence parameters based on the Karhunen-Loeve expansion and an EM algorithm that significantly reduces the dimension of the problem and is computationally efficient. We also demonstrate prediction of an unknown outcome when the other outcomes are known. We apply our methodology to diffusion tensor imaging (DTI) data for multiple sclerosis (MS) patients with three outcomes, and identify differences in both the marginal distributions and the dependence structure between the MS and control groups. ROC curves show that the cross-correlations between DTI outcomes are predictive of MS status. Our proposed methodology is quite general and can be applied to other functional data with multiple outcomes in biology and other fields.

Key words: Diffusion Tensor Imaging; Gaussian copulas; Multiple sclerosis; Skewed functional data; Tractography data

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1. Introduction

Functional data analysis (FDA) is a powerful tool for modeling data observed at various time points or locations to uncover the underlying features that are assumed to vary over a continuum of time points or locations. Ramsay and Silverman (2005) and Ferraty and Romain (2010) provide a comprehensive review of functional data analysis. Often we obtain multiple functional observations and we wish to study them simultaneously. In this case, it is important to have tools to study the dependence structure between different outcomes. The methodology should be efficient in reducing the dimension of the data to accommodate the possible large size of functional data, especially in the case of multiple outcomes.

[Figure 1 about here.]

This study proposes a method for studying functional data with multiple outcomes that exhibits some shape characteristics that vary with spatial or temporal location. Figure 1 illustrates an example of data that motivated this study. The data were obtained from a diffusion tensor imaging (DTI) study of neuronal tracts in 162 multiple sclerosis (MS) patients and 42 healthy controls. DTI is a magnetic resonance imaging technique for measuring the diffusion of water that can be used to detect abnormalities in brain tissue (see Basser et al., 1994, 2000). DTI tractography can be summarized by 3 functional outcomes, fractional anisotropy, parallel diffusivity and perpendicular diffusivity, each along the normalized tract distance. As seen in Figure 1, for each outcome, the pointwise mean and variance of both groups seem to vary along the tract location and are different between the two groups. Another interesting characteristic of this data is that the amount of skewness seems to vary along the tract location as well. An appropriate methodology to study the data should be able to capture this variation in the skewness, especially because skewness is related to disease status and could provide insights into the disease process.
Much of the FDA literature assumes Gaussian marginal distributions, but non-Gaussian functional data is not uncommon. Fortunately, models for non-Gaussian data are now being developed. Staicu et al. (2011) developed a copula-based approach for analyzing functional data with one outcome and applied their study to the parallel diffusivity measurement from this DTI data. The data were assumed to have pointwise marginal distributions in a parametric family with shape parameters such as a skew normal or skew $t$ family (Azzalini, 1985; Azzalini and Capitanio, 1985). The mean, standard deviation and shape parameter functions were modeled nonparametrically as functions of location. The dependence structure was estimated assuming an underlying Gaussian or $t$ copula using methods based on principal component analysis (PCA) or the sample Kendall’s tau matrix. Their results showed that the marginal mean, variance, and skewness functions were different between the healthy and MS groups, while the correlation within the outcome is remarkably similar between the two groups. In contrast, we found the interesting result that the correlation between two different outcomes can differ between the MS patients and controls; see Section 5.

There have been many other studies using the DTI tractography for MS patients. For instance, Reich et al. (2005) analyzed various indices obtained from DTI tractography to identify abnormalities in MS patients. Goldsmith et al. (2011a,b, 2012) studied DTI data using penalized functional regression. Longitudinal functional PCA (LFPCA) was introduced by Greven et al. (2010) to account for the longitudinal variability of tractography data in MS. LFPCA extended the MFPCA approach of Di et al. (2009) who focused on replicated functional data. See Staicu et al. (2003) for the application of PCA for complex multilevel spatially correlated functional data. McLean et al. (2012) introduced a functional generalized additive model and applied it to the relationship between a cognitive test score and DTI tractography.
There are relatively few previous studies that analyze the relationship between multiple functional outcomes. Ramsay and Silverman (2005) explained and demonstrated a PCA methodology to study the simultaneous variation of more than one functional outcomes. Zhou et al. (2008) used a functional data approach to analyze paired longitudinal data based on a mixed-effect model framework. One of the main challenges in analyzing functional data with multiple outcomes is the number of parameters that need to be estimated. As the number of outcomes grows, the traditional method of moment estimators might not be practical. We overcome this challenge with two important tools: 1) using the copula approach introduced by Staicu et al. (2011) so that the marginal distributions of the data and its dependence structure can be modeled separately, and, 2) under the Gaussian copula assumption, applying the Karhunen-Loeve (KL) expansion for Gaussian processes to write the data processes as a sum of a small number of components. The dependence between outcomes can be modeled through the dependence between their components. The resulting covariance structure of these components is similar to those in the study by Zhou et al. (2008) under a mixed-effect model framework. We develop a similar EM algorithm as done by Zhou et al. (2008) to calculate maximum likelihood estimators of the copula parameters. Our algorithms are fast and does not require separated steps to model the dependence within each outcome and between different outcomes. This allows the use of a bootstrapping technique to obtain confidence intervals for the estimates. We also explain the steps needed to predict an outcome when all of the other outcomes are known, along with its prediction intervals.

We demonstrate our methodology using the DTI data from the healthy and MS groups. The objective is to identify the locations where the abnormalities occur and also explain the characteristics of the abnormalities. The intention is to use differences between controls and diseased subjects in any DTI characteristics to help understand the disease process. These differences should be useful for detecting MS and for monitoring the progress of a treatment.
We found the same result as in Staicu et al. (2011) for parallel diffusivity. Each of the mean, variance, and skewness functions differs between the two groups. For fractional anisotropy and perpendicular diffusivity, only the mean and variance functions are significantly different. The correlations within each outcome are similar for the two groups. Our most important finding is that the cross-correlation between fractional anisotropy and parallel diffusivity is different between the MS and control groups in the middle part of the tract (roughly from tract locations 0.2 to 0.8). The cross correlation in this tract section is slightly positive in the control group, while it is negative in the MS group. Bootstrapping confirms that the difference is significant. In Section 5 we develop a novel method for predicting disease status using the difference in cross-correlation structure between MS cases and controls. ROC curves with confidence intervals show that there is statistical significant predictive power.

The difference between the two groups in the cross-correlation between the fractional anisotropy and the perpendicular diffusivity is less apparent, while the cross-correlation between the perpendicular and parallel diffusivities is similar in most parts of the tract.

The article is organized as follows. We briefly explain the process for estimating marginal distributions and present the model for the dependence structure in Section 2. In Section 3, the EM and prediction algorithms are presented. In Sections 4 and 5, the methodology is applied to simulated data sets and the DTI data, respectively. Section 6 concludes with a discussion including possible extensions.

2. Modeling Multiple Outcomes
Let \( \{ Y_{ip}(t); t_{ij} \in \mathcal{T} \} \) with \( j = 1, 2, \ldots, m_i \) be the data from outcome \( p, p = 1, 2, \ldots, P \) for subject \( i, i = 1, 2, \ldots, N \) observed at a grid points \( \{ t_{i1}, \ldots, t_{i m_i} \} \in \mathcal{T} \). Following the copula approach introduced by Staicu et al. (2011), we suppose that

\[
Y_{ip}(t) = \mu_p(t) + \sigma_p(t)G^{-1}\{W_{ip}(t); \alpha_p(t)\}
\]
where $\mu_p(t)$ is the mean function and $\sigma_p(t)$ is the standard deviation function of outcome $p$. Here $W_{ip}(t)$ is a latent process such that for each $t$, $W_{ip}(t)$ is uniformly (0,1) distributed, and $G^{-1}(\cdot, \alpha)$ is the inverse of $G$ in a parametric family of distribution functions with zero mean, unit variance, and shape parameter $\alpha$. For example, $G$ can be a skew normal or skew t distribution. In the case of the skew normal, the shape parameter is the skewness parameter, while in the case of the skew t, the shape parameter includes the skewness parameter and the degree of freedom (Azzalini, 1985; Azzalini and Capitanio, 1985). We assume that $\mu_p(t)$, $\sigma_p(t)$, and $\alpha_p(t)$ vary smoothly with $t$. The main objective is to estimate the marginal parameter functions and the dependence structure within each outcome and across different outcomes. We take a two-step approach. In the first step, we estimate the marginal parameter functions $\mu_p(t)$, $\sigma_p(t)$, and $\alpha_p(t)$ for each outcome $p$. In the second step, we estimate the correlation function of $W_{ip}(t)$.

2.1 Modeling Marginal Distribution

This section summarizes the method used to estimate the marginal parameter functions. For details, see Staicu et al. (2011). The estimation is done in two steps.

**step 1:** For simplicity, suppose the data are observed on a common dense grid of points so that $t_{ij} = t_j$ for all $i$ and $j = 1, 2, \ldots, m$. In this first step, we obtain an undersmoothed estimates $\tilde{\mu}_p(t)$, $\tilde{\sigma}_p(t)$ and $\tilde{\alpha}_p(t)$ for $\mu_p(t)$, $\sigma_p(t)$, and $\alpha_p(t)$ by maximizing the pointwise likelihood function

$$
\ell \{\mu_p(t_j), \sigma_p(t_j), \alpha_p(t_j)\} = \sum_{i=1}^{N} \log \left[ g \left\{ \frac{Y_{ip}(t_j) - \mu_p(t_j)}{\sigma_p(t_j)} \right\} ; \alpha_p(t_j) \right] - \log \{\sigma_p(t_j)\}
$$

where $g(y; \alpha) = \partial G(y; \alpha)/\partial y$ is the density function corresponding to the distribution function $G$. When $G$ is assumed to be a skew normal or skew-t distribution, the estimates $\tilde{\mu}_p(t)$, $\tilde{\sigma}_p(t)$ and $\tilde{\alpha}_p(t)$ can be computed using the functions *sn.mle* or *st.mle* from R package
step 2: The estimates from step 1 are smoothed further using penalized splines. Write
the mean function as $\mu_p(t) = B(t)^T \beta_{\mu,p}$, where $B(t)$ is a vector of spline functions
evaluated at $t$ and $\beta_{\mu,p}$ is a vector of spline coefficients. The estimates $\hat{\beta}_{\mu,p}$ for $\beta_{\mu,p}$
are obtained by minimizing the penalty criterion

$$PL_{\mu}(\beta_{\mu,p}) = \sum_{j=1}^{m} \left\{ \mu_p(t_j) - B_j^T \beta_{\mu,p} \right\}^2 + \lambda_{\mu,p} \Omega_{\mu,p}(\beta_{\mu,p}),$$

where $\Omega_{\mu,p} = \beta_{\mu,p}^T D_{\mu,p} \beta_{\mu,t}$, $B_j = B(t_j)$, and $D_{\mu,p}$ is a penalty matrix. See Ruppert et al.
(2003) for a discussion of penalized splines and penalty matrices. The final estimates $\hat{\mu}_p$ is
$B(t)^T \hat{\beta}_{\mu,p}$. The estimates for the standard deviation function can be computed similarly. For
the shape parameter function, Staicu et al. (2011) proposed a penalized marginal pseudo-
likelihood criterion where the mean and variance parameter functions are fixed at estimates.

That is, the criterion to be minimized are

$$PL_{\alpha}(\beta_{\alpha,p}) = -2 \sum_{i=1}^{N} \sum_{j=1}^{m} \ell_{ij}(\beta_{\alpha,p}; \tilde{Y}_{ip}(t_j)) + \lambda_{\alpha,p} \Omega_{\alpha,p}(\beta_{\alpha,p})$$

where $\tilde{Y}_{ip}(t_j) = \{ Y_{ip}(t_j) - \hat{\mu}_p(t_j) / \hat{\sigma}_p(t_j) \}$ are the standardized observations and $\ell_{ij}(\beta_{\alpha,p}; \tilde{Y}_{ip}(t_j)) = \log [g(\tilde{Y}_{ip}(t_j); B_j^T \beta_{\alpha,p})]$. In some cases, it might be better to model a transformed parameter
$h(\alpha)$ instead of $\alpha$. Some possible criteria for choosing smoothing parameters are the restricted
maximum likelihood (REML) (Wood, 2006), AIC, corrected AIC (Ruppert et al., 2003) and
cross-validation (CV). See Staicu et al. (2011) for details and a discussion of smoothing
parameter selection and an alternative method.

2.2 Modeling the Dependence across Different Outcomes

We assume Gaussian copulas for simplicity and because they seem adequate for our purposes.
After the estimates of the parameters of marginal distributions for each outcome $\hat{\mu}_p(t), \hat{\sigma}_p(t)$
and $\hat{\alpha}_p(t)$ are obtained, we transform the observed outcome by

$$R_{ip}(t) = \Phi^{-1}\left[G\left\{\frac{Y_{ip}(t) - \hat{\mu}_p(t)}{\hat{\sigma}_p(t)}; \hat{\alpha}_p(t)\right\}\right],$$

(5)

where $\Phi$ is the cumulative distribution function of the standard normal distribution, and $G(\cdot; \alpha)$ is the distribution function of a distribution with mean 0, variance 1, and shape parameter $\alpha$. We model $R_{ip}$ as the sum of two independent components: 1) a finite Karhunen-Loeve (KL) expansion with a small number of components, and 2) a white noise process with variance $\sigma_{\epsilon p}^2$. That is,

$$R_{ip}(t) = \sum_{k=1}^{K_p} Z_{ipk}f_{kp}(t) + \epsilon_{ip}(t),$$

(6)

where $\{Z_{ikp}, k = 1, 2, \ldots, K_p\}$ are normally distributed with mean 0 and are independent across $k$, and $f_{kp}$ are the eigenfunctions of the covariance function of the process $R_{ip}(t)$ (Levy, 2008). To have the marginal variance of $R_{ip}(t)$ equal to 1, we require that for all $t$,

$$\sum_{k=1}^{K_p} f_{kp}^2(t) \text{var}(Z_{ipk}) + \sigma_{\epsilon p}^2 = 1.$$  

(7)

For identifiability purpose, we assume that the variances of $\{Z_{ikp}, k = 1, 2, \ldots, K_p\}$ are in decreasing order.

The eigenfunctions $f_{kp}$ can be estimated using splines as follows. Let $b(t) = \{b_1(t), \ldots, b_q(t)\}^T$ be an orthonormal spline basis, i.e. $\int_T b_k(t)b_l(t)dt = \delta_{kl}$ where $\delta_{kl}$ is the delta function. We estimate $f_p = (f_{1p}, \ldots, f_{K_p p})^T$ using $b(t)$ as

$$f_p(t)^T = b(t)^T\Theta_p$$

(8)

where $\Theta_p$ is a matrix of spline coefficients with dimension $q \times K_p$ with orthogonal columns. The orthogonality of $b(t)$ and $\Theta_p$ implies the orthogonality of the principal component curves $f_{kp}$.

Let $Z_{ip} = (Z_{ip1}, \ldots, Z_{ipK_p})^T$ and $Z_i = (Z_{i1}^T, \ldots, Z_{iK_p}^T)^T$. Let $D_p = \text{cov}(Z_{ip})$, which is diagonal since the $Z_{ipk}$ are independent across $k$. To have identifiability, we also have that
the diagonal elements of $D_p$ are in a decreasing order. Let $C_{pp'} = \text{cov}(Z_{ip}, Z_{ip'})$. Then we can write the covariance matrix $\Sigma$ of $Z_i$ as

$$
\Sigma = \text{cov}(Z_i) = 
\begin{pmatrix}
D_1 & C_{12} & \cdots & C_{1P} \\
C_{21} & D_2 & \cdots & C_{2P} \\
\vdots & \vdots & \ddots & \vdots \\
C_{P1} & C_{P2} & \cdots & D_P \\
\end{pmatrix}
$$

(9)

Suppose that we have the same observation times $\{t_j, j = 1, 2, \ldots, m\}$ for every subject $i$. Let $R_{ip} = \{R_{ip}(t_1), \ldots, R_{ip}(t_m)\}^T$, $B = \{b(t_1), \ldots, b(t_m)\}^T$ and $\epsilon_{ip} = \{\epsilon_{ip}(t_1), \ldots, \epsilon_{ip}(t_m)\}^T$. Putting everything together, we have the following reduced-rank model for $R_{ip}$,

$$
R_{ip} = B\Theta_p Z_{ip} + \epsilon_{ip}
$$

(10)

$$
\epsilon_{ip} \sim \mathcal{N}(0, \sigma^2_{\epsilon_p} I_m), \quad Z_{ip} \sim \mathcal{N}(0, D_p), \quad \text{cov}(Z_{ip}, Z_{ip'}) = C_{pp'}, \quad \text{for} \quad p \neq p'.
$$

Our reduced-rank model for the latent processes (10) has a similar dependence structure as the model for two outcomes introduced by Zhou et al. (2008). Here we adopt a similar approach using an iterative EM algorithm for estimating the parameters.

To have identifiability, it is sufficient to require that the first nonzero elements of each column of $\Theta_p$ are positive. With finite samples, it is best to determine the sign using the elements with the largest magnitude in each column of $\Theta_p$, since this choice is least influenced by finite-sample random fluctuation (for details, see Zhou et al., 2008). In our estimation algorithm, we require that, in every iteration, the element with the largest magnitude in each column of $\Theta_p$ is positive. In the next section, we develop an algorithm for estimating parameters in model (10).
3. Estimation Algorithms

This section explains the criterion and algorithm for estimating parameters, along with a discussion of tuning parameter specification. We also explain how to predict an outcome after we have observed each of the other outcomes.

3.1 Parameter Estimation Algorithm

The estimates of the dependence parameters can be obtained by maximizing the pseudo-likelihood, with $R_{ip}$ obtained by transforming the observed data using the estimates of the marginal parameters. Let $L_i(\Lambda)$ denote the contribution to the likelihood from individual $i$, where $\Lambda$ is the covariance matrix of $R_i = (R_{i1}^T, \ldots, R_{iP}^T)^T$. The loglikelihood for individual $i$ is given as

$$\log L_i(\Lambda) = -\frac{Pm}{2} \log(2\pi) - \frac{1}{2} \log |\Lambda| - \frac{1}{2} R_i^T \Lambda^{-1} R_i, \quad (11)$$

where $\Lambda$ can be computed in terms of $(\{\Theta_p\}, \Sigma, \{\sigma_{ep}^2\})$ as

$$\text{cov}(R_i) = \Lambda = \begin{pmatrix} \Lambda_{11} & \Lambda_{12} & \cdots & \Lambda_{1P} \\ \Lambda_{21} & \Lambda_{22} & \cdots & \Lambda_{2P} \\ \vdots & \vdots & \ddots & \vdots \\ \Lambda_{P1} & \Lambda_{P2} & \cdots & \Lambda_{PP} \end{pmatrix}$$

where

$$\Lambda_{pp} = B\Theta_p D_p \Theta_p^T B^T + \sigma_{ep}^2 I_m \quad \text{and} \quad \Lambda_{pp'} = B\Theta_p C_{pp'} \Theta_p^T B^T \quad (12)$$

Even though the main interest here is to estimate the covariance matrix $\Lambda$ of $R_i$, not the principal component functions which can obtained from $\Theta_p$, we use the reduced-rank model for $R_{ip}$ so that our estimation algorithm involves estimating $D_p$, $C_{pp'}$ and $\Theta_p$ which are of much smaller size than $\Lambda$. The KL expansion also provides a nice diagonal structure for $D_p$. Our iterative algorithm estimates $D_p$ and $C_{pp'}$ sequentially. Once the estimates for all of the
parameters are obtained, the estimated covariance of $R_{ip}(t)$ can be computed by plugging in the estimates to equation (12).

Extending the approach as in Zhou et al. (2008), we use the penalized pseudo-likelihood and minimize

\[-2 \sum_{i=1}^{N} \log L_i(\Lambda) + \sum_{p=1}^{P} \lambda_p \sum_{k=1}^{K_p} \Theta_{pk}^{T} \int b''(t)b''(t)^{T} dt \Theta_{pk}\]

\[= \sum_{i=1}^{N} \left\{ P m \log(2\pi) + \log |\Lambda| + R_i^{T} \Lambda^{-1} R_i \right\} + \sum_{p=1}^{P} \sum_{k=1}^{K_p} \lambda_p \Theta_{pk}^{T} \int b''(t)b''(t)^{T} dt \Theta_{pk}, \tag{13} \]

where $\Theta_{pk}$ denotes the $k^{th}$ column of $\Theta_p$, and $\lambda_{pk}$ are the smoothing parameters. In general, we can have different $\lambda_{pk}$ for each $f_{pk}$. Since smoothing parameters are mostly determined by the scale of data, to simplify the computation, we assume that $\lambda_{pk} = \lambda_{p}$ for all $k = 1, \ldots, K_p$. Minimizing this expression can be complicated. Instead, we treat the $Z_i$ as missing values and use the EM algorithm (Dempster et al., 1977). The joint loglikelihood is given as

\[
\log L(R_i, Z_i) = \log \left\{ f(R_{i1}|Z_{i1}) \ldots f(R_{ip}|Z_{ip})f(Z_i) \right\} \\
= \sum_{p=1}^{P} \left\{ -\frac{m}{2} \log(\sigma^2_{\epsilon_p}) - \frac{1}{2\sigma^2_{\epsilon_p}} (R_{ip} - B\Theta_{p}Z_{ip})^{T}(R_{ip} - B\Theta_{p}Z_{ip}) \right\} \\
- \frac{1}{2} \log(|\Sigma|) - \frac{1}{2} Z_{i}^{T}\Sigma^{-1}Z_{i}. \tag{14} \]

The EM algorithm is performed as follows:

**The E-step:** Compute the conditional distribution of $Z_i$ given $R_i$

**The M-Step:** Update the parameter estimates by minimizing

\[-2E \left\{ \sum_{i=1}^{N} \log L(R_i, Z_i) \bigg| R_i \right\} + \sum_{p=1}^{P} \lambda_p \sum_{k=1}^{K_p} \Theta_{pk}^{T} \int b''(t)b''(t)^{T} dt \Theta_{pk} \]

The details for each step are given in Web Appendix A.
3.2 Specification of Splines and Tuning Parameters

Recall that we require the splines \( b(t) \) used to estimate the eigenfunctions \( f_p(t) \) to be orthonormal; see equation (8). This implies that the matrix \( B = \{b(t_1), \ldots, b(t_m)\}^T \) has to be orthogonal. As in Zhou et al. (2008), to obtain an orthogonal matrix \( B \), we start by choosing any spline basis \( \tilde{b}(t) \), such as a truncated power basis, and then we evaluate these functions at the observation points to obtain the matrix \( \tilde{B} = \{\tilde{b}(t_1), \ldots, \tilde{b}(t_m)\}^T \). Let \( \tilde{B} = QR \) be the QR decomposition of \( \tilde{B} \), where \( Q \) is an orthogonal matrix and \( R \) is an upper triangular matrix. Then the orthogonal matrix \( B \) is obtained by \( B = \tilde{B}R^{-1} \). R package \texttt{orthogonalsplinebasis} (Redd, 2011) provides functions for orthogonalizing a spline basis.

The two sets of tuning parameters that need to be specified are the spline smoothing parameters, \( \lambda_p \), and the number of components in the KL expansion, \( K_p \). For penalized splines, the knots are typically placed at fixed quantiles of the observation times or locations, and the number of knots are not critical provided that it is sufficient large. The smoothness of the curves is mainly determined by the smoothing parameters \( \lambda_p \) (Ruppert, 2002; Ruppert et al., 2003). We start by performing the analysis for each individual outcome. For each outcome \( p \), we specify a range of \( \lambda_p \) for the grid search. For each \( \lambda_p \) fixed at a value in the grid search range, we vary the number of components starting with only one component. The EM algorithm (as explained in Section 3.1) is used to fit each 1-outcome curve. We keep adding another component and performing the EM algorithm until the variance of the newly added principal component score is less than a prespecified small fraction of that of the previously added component. The number of components before adding this last component is \( K_p \) chosen for this \( \lambda_p \). The corrected AIC (Ruppert et al., 2003) is computed for this \( (\lambda_p, K_p) \). The pair \( (\lambda_p, K_p) \) with the minimum corrected AIC is selected for this outcome.

Other criteria such as CV and AIC can also be used. Once we analyze multiple outcomes, the number of components can be fine tuned further. We do not expect the values of \( \lambda \) to
change from the individual to the multiple outcome case, as $\lambda$ should depend on the scale of individual outcome data more than the dependence structure between outcomes.

3.3 Confidence Intervals for Estimates

Bootstrapping can be easily applied to obtain pointwise confidence intervals for the estimates for the marginal distribution parameters and the correlations. Bootstrap samples are obtained by resampling the subjects from the original data $\{Y_{ip}\}$. For each bootstrap sample, we calculate estimates for the marginal distribution parameters (as explained in section 2.1), use them to transform the data as in equation (5), and then perform the EM algorithm to obtain estimates for the correlations, by plugging in the parameter estimates to equation (12). The pointwise confidence intervals for the estimates of both the marginal distribution parameters and the correlations are computed based on sample quantiles of the estimates for the bootstrap samples.

3.4 Prediction of an Outcome Using all the Other Outcomes

In addition to simplifying the computation for parameter estimation, our reduced-rank model for multiple outcomes also provides a straightforward way of predicting an outcome for an individual when all the other outcomes of the same subject are known. This can be done using the conditional distribution property of the multivariate normal random variables as follows. Suppose we have the observation $Y_{ip}$ of a subject $i$ for all outcomes $p \neq 1$ and we want to predict outcome 1 for this individual. Let $R_{ip}$ denote the transformed process as in equation (5). Since we have that $(R_{i1}, \ldots, R_{iP}) \sim N(0, \Lambda)$, we can compute the conditional distribution of $R_{i1}$ as

$$R_{i1}|R_{i2}, \ldots, R_{iP} \sim N(\bar{\mu}_1, \bar{\Sigma}_1)$$ (16)
where
\[ \bar{\mu}_1 = \begin{pmatrix} \Lambda_{12} & \ldots & \Lambda_{1P} \end{pmatrix} \begin{pmatrix} \Lambda_{22} & \ldots & \Lambda_{2P} \\ \vdots & \ddots & \vdots \\ \Lambda_{2P} & \ldots & \Lambda_{PP} \end{pmatrix}^{-1} \begin{pmatrix} R_{i2} \\ \vdots \\ R_{iP} \end{pmatrix}, \]

and
\[ \bar{\Sigma}_1 = \Lambda_{11} - \begin{pmatrix} \Lambda_{12} & \ldots & \Lambda_{1P} \end{pmatrix} \begin{pmatrix} \Lambda_{22} & \ldots & \Lambda_{2P} \\ \vdots & \ddots & \vdots \\ \Lambda_{2P} & \ldots & \Lambda_{PP} \end{pmatrix}^{-1} \begin{pmatrix} \Lambda_{21} \\ \vdots \\ \Lambda_{P1} \end{pmatrix}. \]

Then \( \bar{\mu}_1 \) can be used as a predictor for \( R_{i1} \). A prediction \( \hat{Y}_{i1} \) for \( Y_{i1} \) is obtained by transforming \( \bar{\mu}_1 \) back as
\[ \hat{Y}_{i1}(t) = G^{-1} \{ \Phi(\bar{\mu}_1); \bar{\sigma}_1(t) \} \bar{\sigma}_1(t) + \hat{\mu}_1(t), \] (17)

using the estimates \( \hat{\mu}_1(t), \bar{\sigma}_1(t), \bar{\alpha}_1(t) \) obtained earlier. The standard deviation of the prediction error can be computed using bootstrapping and used for a prediction interval. The prediction for other outcomes can be done similarly.

### 4. Simulation Studies

In this section, we demonstrate our methodology using simulated data. As far as we are aware, there exist no other estimators for multiple functional outcomes of the type we are considering, that is, with spatially varying non-Gaussian characteristics. Therefore, there are no competing estimators to compare with ours.

The simulated data contains 100 datasets, each containing \( P = 3 \) outcomes from \( N = 200 \) subjects. Each observation is taken at the common time points \( t_j, j = 1, 2, \ldots, 80 \). The data are generated using the finite KL expansion for Gaussian processes as follows:
(1) For $p = 1, 2, 3$, generate

$$Q_{ip}(t) = \sum_{k=1}^{K_p} Z_{ipk} h_{kp}(t) + \epsilon_{ip}(t),$$  \hspace{1cm} (18)

where $K_1 = K_2 = 2, K_3 = 3, \epsilon_{ip}(t) \sim N(0, \tau_p^2)$, $\tau_1^2 = 0.2, \tau_2^2 = 0.3, \tau_3^2 = 0.4$, $Z_i = (Z_{i11}, Z_{i12}, Z_{i21}, Z_{i22}, Z_{i31}, Z_{i32}, Z_{i33})$ is normal with mean 0 and covariance matrix

$$
\begin{pmatrix}
6.47 & 0 & -4.46 & 1.53 & 2.93 & 0.31 & 1.76 \\
0 & 4.08 & 0.02 & 0.89 & 0.99 & -2.39 & -1.14 \\
-4.46 & 0.02 & 20.03 & 0 & -3.52 & 2.61 & -4.05 \\
1.53 & 0.89 & 0 & 7.46 & 3.51 & -2.01 & -1.43 \\
2.93 & 0.99 & -3.52 & 3.51 & 16.34 & 0 & 0 \\
0.31 & -2.39 & 2.61 & -2.01 & 0 & 7.63 & 0 \\
1.76 & -1.14 & -4.05 & -1.43 & 0 & 0 & 2.98
\end{pmatrix}
$$

The functions $h_{kp}(t)$ are the normalized version of $\sin(\pi t)$ and $\cos(\pi t)$ for $p = 1$, $t^2 - 13t + 43/6$ and $t + 0.5$ for $p = 2$, and $e^t, t - (e - 1)^{-1}$ and $t^2 - 1.0025t + 0.1654$ for $p = 3$. Then we transform $Q_{ip}(t)$ to

$$R_{ip}(t) = \frac{Q_{ip}(t)}{\sigma_p(t)}, \text{ where } \sigma_p^2(t) = \sum_{k=1}^{K_p} \text{var}(Z_{ipk}) h_{kp}^2(t) + \tau_p^2,$$  \hspace{1cm} (19)

so that $R_{ip}(t)$ has a marginal $N(0,1)$ distribution.

(2) Generate the observations

$$Y_{ip}(t) = \mu_p(t) + \sigma_p(t) G^{-1} [\Phi \{ R_{ip}(t) \} ; \alpha_p(t)],$$  \hspace{1cm} (20)

where $\Phi(\cdot)$ and $G(\cdot; \alpha)$ denote the cumulative distribution functions of the standard normal and skew normal distribution with mean 0, variance 1 and shape parameter $\alpha$, respectively. The mean and shape parameter functions are
\[\begin{align*}
\mu_1(t) &= 5, \quad \mu_2(t) = -3t^5 + 4.5t^3 - 1.5t + 15, \quad \mu_3(t) = 10e^{-t^2/2} + 20t^2 + 10, \\
\alpha_1(t) &= 0, \quad \alpha_2(t) = 10 \sin(2\pi t), \quad \alpha_3(t) = 48t^2 - 48t + 6
\end{align*}\] (21)

The plots of simulation results are shown in Figures S1-S4 in Web Appendix B. The estimates of the mean, standard deviation and shape parameter functions of each outcome from all 100 datasets are shown in Web Figure S1. We assumed the skew normal distribution. These estimates were obtained by maximizing the pointwise likelihood at each point \(t_j\) using the R package \texttt{sn} (Azzalini, 2011). The estimates were then smoothed further using penalized truncated cubic polynomial splines. The smoothing parameters were selected using the REML criterion implemented in R package \texttt{mgcv} (Wood, 2006). The black lines indicate the true functions. Even though there seems to be high variation in the estimates when \(\alpha \approx 10\), the density of skew normal for \(\alpha\) equal to 10 is not much different from when \(\alpha\) is much higher. In fact, as \(\alpha \to \infty\) the skew normal distribution converges to the half normal distribution, and when \(\alpha\) is 10 it is already similar to the half normal. This means that even though the estimate of \(\alpha\) in this range is not very accurate, we can still obtain a good estimate for the distribution.

The dependence among the Gaussian copula were estimated using the orthonormal transform of cubic splines with 9 knots. The number of components and smoothing parameters were chosen based on AIC. The transformation and parameter selection are explained in Section 3.2. Web Figure S2 shows the true and estimates of the covariance from 2 datasets. The true covariances within an outcome and cross covariances between outcomes are shown in the first and fourth rows, respectively. The estimated covariances within an outcome from 2 datasets are shown in rows 2 and 3. The estimated cross covariance between outcomes from the same 2 datasets are shown in rows 5 and 6.

Estimated variances of each latent process are shown in Web Figure S3. The values
close to 1 indicate the closeness to our assumption that the latent processes have marginal variance of 1. The square root of the integrated mean square error (IMSE), integrated square bias (IBIAS) and integrated variance (IVAR) for the marginal parameter functions and covariance parameters are shown in Table S1 in Web Appendix B. These quantities are calculated, for example, for the mean function by \( \text{IMSE} = \int_0^1 \hat{E} \{ \hat{\mu}(t) - \mu(t) \}^2 dt \), \( \text{IVAR} = \int_0^1 \hat{\text{Var}} \{ \hat{\mu}(t) \} dt \) and \( \text{IBIAS} = \int_0^1 \left[ \hat{E} \{ \hat{\mu}(t) - \mu(t) \} \right]^2 dt \). Here \( \hat{E} \) and \( \hat{\text{Var}} \) denote the sample mean and sample variance, respectively. Overall, the bias and variance are low for the mean and standard deviation estimates. The bias for the shape parameter estimates are somewhat high, especially, as seen from Web Figure S1, at locations with high \( \alpha \) (in absolute value). This should not affect our estimated marginal distributions too much as explained earlier. The covariance estimates have small IMSE. The contour plots of pointwise square root of the mean square error for the covariance estimates are shown in Figure S4.

5. Application to Diffusion Tensor Imaging (DTI)

In this section, we apply our methodology to the DTI data. A subset of our data set is freely available as the DTI data set in R’s refund package Crainiceanu et al. (2012).

DTI is a magnetic resonance imaging technique that measures the diffusion of water in tissue. The anisotropy of water diffusion allows images of the white matter in the brain to be generated. White matter tracts are made up of axons that transmit signals between different regions of the brain. These axons are surrounded by and insulated by a fatty substance called myelin which helps the signal to be carried quickly. Multiple sclerosis is an autoimmune disease associated with damage to myelin and can lead to significant disabilities in patients (Goldsmith et al., 2012).

DTI provides many measurements of water diffusion. In this study, we consider 3 measurements: 1) fractional anisotropy, 2) parallel diffusivity and 3) perpendicular diffusivity. Parallel and perpendicular diffusivities are diffusion parallel and perpendicular to the long
axis of a fiber bundle, respectively, while fractional anisotropy measures the difference of
diffusion in the two directions. Mathematically, at each location, DTI tractography can be
described as a $3 \times 3$ symmetric, positive definite matrix. Suppose the eigenvalues of the
matrix are $\lambda_1 > \lambda_2 > \lambda_3$. The parallel and perpendicular diffusivities are given by $\lambda_1$ and
$\frac{1}{2} (\lambda_2 + \lambda_3)$, respectively. The fractional anisotropy is calculated as

$$ \left[ \frac{3 \{(\lambda_1 - \bar{\lambda})^2 + (\lambda_2 - \bar{\lambda})^2 + (\lambda_3 - \bar{\lambda})^2\}}{2(\lambda_1^2 + \lambda_2^2 + \lambda_3^2)} \right]^{1/2}, \tag{22} $$

where $\bar{\lambda} = (\lambda_1 + \lambda_2 + \lambda_3)/3$ (McLean et al., 2012).

Our data consists of measurements along the corpus callosum from 42 healthy controls
and 162 MS patients, measured at 93 locations along the tract. Figure 1 displays the 3
outcomes for both groups. Our goal is to estimate for each group the marginal distributions
the outcomes and the dependence structure across locations within and between outcomes.
We also wish to identify differences between the two groups.

5.1 Marginal Parameter Estimation for DTI Data

We assumed that each measurement from each group follows a skew-t marginal distribution.
We assumed that the mean, standard deviation, and skewness parameter vary along the tract,
while the degrees of freedom are constant along the tract locations. (There were insufficient
observations to estimate spatially-varying degrees of freedom parameters.) Using R package
sn (Azzalini, 2011), we obtained the maximum likelihood estimates for the degree of freedom
for the fractional anisotropy, parallel diffusivity and perpendicular diffusivity equal to 24.93,
11112 and 16.30, respectively for the control group, and 11732, 5.98 and 8.47 for the MS
group. We used the skew normal distribution to model any outcome with the estimated
degrees of freedom $> 100$. The undersmoothed estimates for the mean, standard deviation,
and skewness parameters were obtained via pointwise maximum likelihood estimation using
function st.mle in the R package sn (Azzalini, 2011). The mean and standard deviation
estimates were smoothed further by penalized spline fitting using generalized additive model implemented in package \texttt{mgcv} in \texttt{R} (Wood, 2006). The skewness parameters were smoothed further by fitting the penalized spline to maximize the penalized pseudo likelihood computed using the estimates of the mean and standard deviation as in equation (4).

The estimated mean, variance and skewness functions for the three outcomes for both groups are shown in Figure S5 in Web Appendix C. The bootstrap 90\% pointwise confidence intervals for the differences of the marginal parameters between the two groups are shown in Figure 2. The mean, variance, and skewness functions are all statistically significantly different between the two groups for parallel diffusivity, while only the mean and variance are significantly different for fractional anisotropy and perpendicular diffusivity. The mean of fractional anisotropy for the MS group is higher than the control group, but the means of parallel and perpendicular diffusivities are lower in the MS group. The variances for the MS group are higher in all of the three outcomes. The skewness of the fractional anisotropy is close to zero for both groups. The parallel diffusivity of the MS group is positively-skewed while the skewness is close to zero in most part of the tract in the control group. The difference in the skewness between the two groups is significant. The perpendicular diffusivity is positively-skewed in most part of the tract in both groups. There is no significant difference in the skewness between the two groups.

[Figure 2 about here.]

5.2 Dependence Structure of DTI Data

After the estimates $\hat{\mu}_p, \hat{\sigma}_p, \hat{\alpha}_p$ for the mean, standard deviation and skewness parameters for the outcome $p = 1, 2, 3$ for each group were obtained, the data were transformed using equation (5). Here, $G$ is the skew-t distribution with mean 0, variance 1, skewness parameter $\hat{\alpha}_p$ and degree of freedom for each outcome and each group as stated earlier. The method explained in Section 3 was applied to the transformed data. The transformation implies that
the marginal variance of the transformed process is 1, and, hence, the covariance function
for the transformed process is also its correlation function. In our study, estimates of the
marginal variance vary from 0.6 to 1.3 for the MS group, and 0.6 to 1.5 for the control
group, due to the small sample sizes. We verified this by a simple simulation as follows. We
generated data with 42 observations for control group, and 162 observations for MS group
according to the skew t distribution using the estimated parameters obtained earlier. Then
we transformed the simulated data as in Equation (5) and computed the marginal sample
variances. The variances vary similarly to those of the DTI data. The sample variances are
noticeably closer to 1 as the number of observations for each group increases to 200. Most of
the sample variances lie well within [0.9,1.1] as the number of observations increases to 500.

[Figure 3 about here.]

[Figure 4 about here.]

The estimated correlations for the DTI data were obtained by converting the estimated
covariances for the transformed process (i.e., scale the covariances by the marginal variances).
Figure S6 in Web Appendix C displays the estimated correlations within each of the three
outcomes for the healthy and MS groups. Figure 3 displays the estimated cross-correlations
between different outcomes for the healthy and MS groups. The correlations within the
outcomes look somewhat similar for the two groups, while there is a noticeable difference
between the two groups for the cross-correlations, especially for the cross-correlation between
parallel diffusivity and fractional anisotropy. The pointwise confidence intervals for the
estimated correlations can be computed using bootstrapping. Figure 4 shows the significance
levels of the difference of the correlations between the two groups calculated from the 90%
and 95% bootstrap pointwise confidence intervals based on 1000 samples.

The bootstrapping results confirm that there is no significant difference in the correlations
within fractional anisotropy and parallel diffusivity. The correlation within perpendicular
diffusivity is different in various small regions of the tract. There is a significant difference between the two groups in the cross-correlation between the fractional anisotropy and the parallel diffusivity in a large middle part of the tract (at around locations 0.2 to 0.8; see the bottom, left panel of Figure 4). The cross-correlation is slightly positive in the control group, while it is negative in the MS group. The difference in the cross-correlation between the fractional anisotropy and the perpendicular diffusivity is significant in a smaller part of the tract (at around locations 0.4 to 0.8; see the bottom, middle panel of Figure 4). The cross-correlation is more negative in the MS group in this region. The cross-correlation between the parallel and perpendicular diffusivities is somewhat similar between the two groups with the parallel diffusivity at around location 0.2 being more positively-correlated with the overall perpendicular diffusivity in the MS group.

5.3 Case Status Prediction for DTI Data

In this section, we demonstrate an application of the prediction introduced in Section 3.4 to predict subjects' case status. For each subject with unknown case status, we use the parallel and perpendicular diffusivities to obtain two predictions for the fractional anisotropy using 2 sets of parameters: one for the MS group and the other for the control group. If this subject is a MS patient, the prediction using the MS parameters should be more similar to the real observed curve than the prediction using the control parameters. One possible criterion for measuring the similarity between curves is to compare the areas between each predicted curve and the observed curve. The prediction can be similarly applied to the parallel and perpendicular diffusivities.

We studied the discriminating power of our prediction using a leave-one-out analysis. To predict case status for subject $i$, we obtained estimates for marginal and dependence parameters using all of the subjects but subject $i$. Then we performed prediction of an outcome using the other two outcomes as explained in Section 3.4 to subject $i$ using the
obtained parameter estimates for MS and control groups. Let $\text{Area}_{i,\text{control},p}$ and $\text{Area}_{i,\text{MS},p}$ denote the areas between the observed outcome $p$ of subject $i$ and the predicted outcome $p$ for subject $i$ using the control and MS parameters, respectively. To predict the case status, we use a test statistic

$$T_{i,p} = \text{Area}_{i,\text{control},p} - \text{Area}_{i,\text{MS},p},$$

and classify this subject to be MS patient if $T_{i,p} \geq \tau_p$ for some threshold value $\tau_p$. Figure 5 depicts receiver operating characteristic (ROC) curves with confidence intervals as the threshold values vary using predicted curves for each of the 3 outcomes. The vertical axis shows the true positive rate which is the proportion of MS patients that are correctly classified, and the horizontal axis shows the false positive rate which is the proportion of healthy subjects that are incorrectly classified. The ROC curves and their confidence intervals were produced by the R package \texttt{pROC} (Robin et al., 2011). The confidence intervals lie almost entirely above the 45° line, which indicates that the differences in correlation structure between the MS and control groups have statistically significant predictive power.

[Figure 5 about here.]

6. Discussion

We have developed a copula-based approach for modeling functional data with multiple outcomes. The copula approach, together with the KL expansion, significantly reduces the dimension of the model and allows fast estimation of the marginal distributions and the dependence structure. This is essential in the analysis of functional data, especially in the case of multiple outcomes where the data set can be very large.

Our methodology performed very well in our simulation study. We extended the study done by Staicu et al. (2011) to include three outcomes from DTI study and were able to identify the differences in the pointwise marginal distributions between the control and MS groups.
Our dependence structure study shows that the correlations within the same outcome are not different between the two groups, while the cross correlation between outcomes, especially between the parallel diffusivity and fractional anisotropy, are different. ROC curves show that the cross-correlations between DTI outcomes are predictive of MS status.

One extension to our Gaussian assumption would be to a t-copula. The t-copula covers a larger class of models than Gaussian copula and is better in capturing tail dependence behavior (McNeil et al., 2005). The KL expansion used in our model does not apply directly to the t-copula. Another efficient alternative methodology will need to be developed.

Another possible extension is to study functional data in a longitudinal setting. For example, the data used in this study, which contains only the data from the first visit of each subject, was taken from a larger dataset that contains measurements from subjects’ multiple visits. This data set was studied by, for example, Goldsmith et al. (2012) and Greven et al. (2010). Goldsmith et al. (2012) studied the relation between cognitive disability in MS patients by incorporating DTI tractography measurement from multiple visits as functional covariates in a longitudinal penalized functional regression setting. Greven et al. (2010) developed a longitudinal functional principal component analysis approach to decompose the longitudinal DTI data into a time-dependent population average, baseline subject-specific variability, longitudinal subject-specific variability, subject/visit-specific variability and measurement error. A longitudinal extension to our methodology would allow ones to assess how the pointwise marginal distributions of each outcome and/or the dependence within and across the outcomes across tract locations of the subjects from each group change over time.

7. Supplementary Materials

Web Appendices, Table and Figures referenced in Section 4 and Section 5 are available with this paper at the Biometrics website on Wiley Online Library.
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REFERENCES


Multiple Sclerosis 16, 166–177.


**Figure 1.** Plot of DTI data. The top row shows the three outcomes from 42 healthy controls. The bottom row shows the same three outcomes from 162 MS patients. The tract distance is normalized to [0,1]
Figure 2. Estimated differences (black solid lines) in the mean, log-variance and skewness between the healthy and MS groups. The black dashed lines show 90% confidence intervals computed using bootstrapping with 1000 samples. The gray dashed horizontal lines are through 0.
Figure 3. Estimated cross-correlations between different outcomes for the control group (left panel) and the MS group (right panel). This figure appears in color in Figure S7 in Web Appendix C.
Figure 4. The differences of the correlations within each outcome (top panel) and between outcomes (bottom panel) between the healthy and MS groups. The gray and black regions are the areas at which the differences are statistically significant at 90% and 95% levels, respectively. The computation is based on bootstrap pointwise confidence intervals using 1000 samples.
Figure 5. Receiver operating characteristic curves with 95% confidence intervals for case status prediction using the predicted fractional anisotropy, parallel diffusivity and perpendicular diffusivity.