



## A powerful and efficient multivariate approach for voxel-level connectome-wide association studies



Weikang Gong<sup>a,b,e,\*</sup>, Fan Cheng<sup>d,e</sup>, Edmund T. Rolls<sup>c</sup>, Chun-Yi Zac Lo<sup>e</sup>, Chu-Chung Huang<sup>f</sup>, Shih-Jen Tsai<sup>g</sup>, Albert C. Yang<sup>f</sup>, Ching-Po Lin<sup>f,\*\*\*</sup>, Jianfeng Feng<sup>e,c,d,\*\*</sup>

<sup>a</sup> Key Laboratory of Computational Biology, CAS-MPG Partner Institute for Computational Biology, Shanghai Institutes for Biological Sciences, Chinese Academy of Sciences, Shanghai, 200031, China

<sup>b</sup> University of Chinese Academy of Sciences, Beijing, 100049, China

<sup>c</sup> Department of Computer Science, University of Warwick, Coventry, CV4 7AL, UK

<sup>d</sup> Shanghai Center for Mathematical Sciences, Fudan University, Shanghai, 200433, China

<sup>e</sup> Institute of Science and Technology for Brain-Inspired Intelligence, Fudan University, Shanghai, 200433, China

<sup>f</sup> Institute of Neuroscience, National Yang-Ming University, Taipei, Taiwan

<sup>g</sup> Department of Psychiatry, Taipei Veterans General Hospital, Taipei, Taiwan

### ARTICLE INFO

#### Keywords:

Multivariate analysis  
Structured kernel principal component regression  
Association study  
Functional connectivity

### ABSTRACT

We describe an approach to multivariate analysis, termed structured kernel principal component regression (sKPCR), to identify associations in voxel-level connectomes using resting-state functional magnetic resonance imaging (rsfMRI) data. This powerful and computationally efficient multivariate method can identify voxel-phenotype associations based on the whole-brain connectivity pattern of voxels, and it can detect linear and non-linear signals in both volume-based and surface-based rsfMRI data. For each voxel, sKPCR first extracts low-dimensional signals from the spatially smoothed connectivities by structured kernel principal component analysis, and then tests the voxel-phenotype associations by an adaptive regression model. The method's power is derived from appropriately modelling the spatial structure of the data when performing dimension reduction, and then adaptively choosing an optimal dimension for association testing using the adaptive regression strategy. Simulations based on real connectome data have shown that sKPCR can accurately control the false-positive rate and that it is more powerful than many state-of-the-art approaches, such as the connectivity-wise generalized linear model (GLM) approach, multivariate distance matrix regression (MDMR), adaptive sum of powered score (aSPU) test, and least-square kernel machine (LSKM). Moreover, since sKPCR can reduce the computational cost of non-parametric permutation tests, its computation speed is much faster. To demonstrate the utility of sKPCR for real data analysis, we have also compared sKPCR with the above methods based on the identification of voxel-wise differences between schizophrenic patients and healthy controls in four independent rsfMRI datasets. The results showed that sKPCR had better between-sites reproducibility and a larger proportion of overlap with existing schizophrenia meta-analysis findings. Code for our approach can be downloaded from <https://github.com/weikanggong/sKPCR>.

### 1. Introduction

Functional connectivity analysis using resting-state functional magnetic resonance imaging (fMRI) data has become increasingly popular in the last few years (Smith et al., 2015; Finn et al., 2015), and the advances

have led to many investigations of functional dysconnectivity between brain areas in neurodegenerative and psychiatric brain diseases (Gong and He, 2015; Romme et al., 2017). Voxel-based functional connectivity analysis has also recently emerged (Cheng et al., 2015, 2016; Rolls et al., 2018; Satterthwaite et al., 2015; Kaczkurkin et al., 2017). However,

\* Corresponding author. Key Laboratory of Computational Biology, CAS-MPG Partner Institute for Computational Biology, Shanghai Institutes for Biological Sciences, Chinese Academy of Sciences, Shanghai, 200031, China.

\*\* Corresponding author. Institute of Science and Technology for Brain-Inspired Intelligence, Fudan University, Shanghai 200433, China.

\*\*\* Corresponding author. Institute of Neuroscience, National Yang-Ming University, Taipei, Taiwan.

E-mail addresses: [weikanggong@gmail.com](mailto:weikanggong@gmail.com) (W. Gong), [cplin@ym.edu.tw](mailto:cplin@ym.edu.tw) (C.-P. Lin), [jianfeng64@gmail.com](mailto:jianfeng64@gmail.com) (J. Feng).

<https://doi.org/10.1016/j.neuroimage.2018.12.032>

Received 27 August 2018; Received in revised form 14 November 2018; Accepted 14 December 2018

Available online 18 December 2018

1053-8119/© 2018 Elsevier Inc. All rights reserved.

designing methods to explore the associations between the whole-brain voxel-level connectome and phenotypes is a challenging task, and well-developed approaches are usually designed for parcellation-based or seed-based connectivity studies (Meskaldji et al., 2013; Bellec et al., 2015; Xia and He, 2017).

The most popular method for functional connectivity analysis is the massive univariate generalized linear model (GLM) approach. This approach uses a GLM to test the association between each voxel-voxel connectivity and the phenotype of interest, and then corrects for multiple comparison (Cheng et al., 2015, 2016) by such methods as Bonferroni correction, false-discovery rate (Benjamini and Hochberg, 1995) and random field theory (Gong et al., 2018), to locate the significant signals. The major advantage of this approach is that it can provide the exact location of the signals. However, the large number of hypothesis tests require a stringent multiple comparison threshold which usually decrease the power. In addition, univariate approach only tests the linear relationships between connectivities and phenotypes. Important higher-order information, such as the co-contribution of a set of functional connectivities and the non-linear associations, is usually ignored by this method.

In recent years, many improvements over the univariate method have been proposed. These approaches usually adopt global association tests to achieve higher power. In other words, they test whether the signal is present somewhere in a set of functional connectivities rather than localizing it. We briefly review some of them here. First, in the brain-wide association study (BWAS) approach (Gong et al., 2018), the authors proposed to test whether the observed cluster size of the supra-threshold functional connectivities is larger than that by chance. The BWAS is a generalization of the traditional cluster-size inference approach (Friston et al., 1994) and popular network-based statistic (NBS) approach (Zalesky et al., 2010) to voxel-level connectivity studies. Second, the multivariate distance matrix regression (MDMR) (Shehzad et al., 2014) is a nonparametric multivariate approach, which directly tests the association between a phenotype of interest and a between-subject distance matrix estimated using the functional connectivity data. Third, in Pan et al. (2014); Kim et al. (2014, 2015), the authors proposed the adaptive sum of powered score (aSPU) test and its extensions. This approach first assigns a score to measure the association between a phenotype and an individual connection. It then combines all the individual scores into a summary statistic and uses a permutation test to access the significance. Other related approaches include (Simpson and Laurienti, 2015; Chen et al., 2015b; Fiecas et al., 2017; Meskaldji et al., 2015; Belilovsky et al., 2016). However, in the context of voxel-level connectivity analysis, the above approaches have three major drawbacks: First, they do not explicitly model the spatial structure of the voxel-level connectome, which is structurally and smoothly correlated. Therefore, as shown in our analysis, the unmodelled spatial noise usually decreases their power. Second, they use the computationally expensive nonparametric permutation to get voxel/connectivity-wise p-values. Third, they can only detect linear association signals, whereas important nonlinear signals may be missed.

In this paper, we propose a novel multivariate approach specifically designed to detect associations in the voxel-level connectome and overcome the above mentioned drawbacks of previous methods. This approach, termed ‘structured kernel principal component regression’ (sKPCR), is specifically designed for the voxel-level connectome, and it can be applied to both volume-based and surface-based fMRI data. The sKPCR evaluates the simultaneous contribution of the whole-brain connectivities of each voxel to a phenotype of interest. We have designed this method to perform three steps: (1) extract important features from the data using a newly developed structured kernel principal component analysis (sKPCA) approach; (2) test the association between the low-dimensional features (principal components) and the phenotype of interest using an adaptive regression approach; and (3) control the voxel-wise family-wise error rate (FWER), using an efficient nonparametric permutation procedure. Methodologically, we make three contributions to the science of voxel-level

connectome analysis. First, we developed sKPCA as the first step of sKPCR, which is an extension of the widely used principal component analysis (PCA) (Jolliffe, 2002) and probabilistic PCA methods (Tipping and Bishop, 1999). However, unlike the PCA method, which assumes independent and identically distributed noise structure, sKPCA assumes a more realistic, spatially correlated noise structure among functional connectivities, leading to superior performance in dimension reduction. A nonlinear extension is also developed based on the idea of kernel principal component analysis (Schölkopf et al., 1997). Second, we proposed a new adaptive linear regression approach as the second step of sKPCR to test the association between a set of principal components and phenotypes of interest. The model can adaptively choose the optimal number of principal components, and its performance is robust, even if many noise components are wrongly included in the model. Third, we developed a highly efficient permutation approach which can simultaneously estimate voxel-wise p-values and correct for multiple comparisons. Other attractive features of sKPCR include 1) applicability for both categorical (e.g., disease status) and continuous variables (e.g., IQ, symptom scores), as well as 2) covariate effects (e.g., age, gender, motion).

The remainder of the paper is organized as follows. We begin by reviewing PCA and its probabilistic model. Next, we describe the details of sKPCR, including sKPCA and adaptive regression (see Fig. 1 for a graphical overview). We then conduct comprehensive simulations to compare the proposed sKPCR approach with several state-of-the-art methods discussed above, such as the univariate approach, MDMR, aSPU, in terms of their false-positive rate, power of detecting signals and computation time. Finally, using four schizophrenia datasets, we evaluate and compare their between-sites reproducibility and the proportion of overlaps with existing schizophrenia meta-analysis findings. The code for our approach can be downloaded from <https://github.com/weikanggong/sKPCR>.

## 2. Method

### 2.1. Structured kernel principal component analysis (sKPCA)

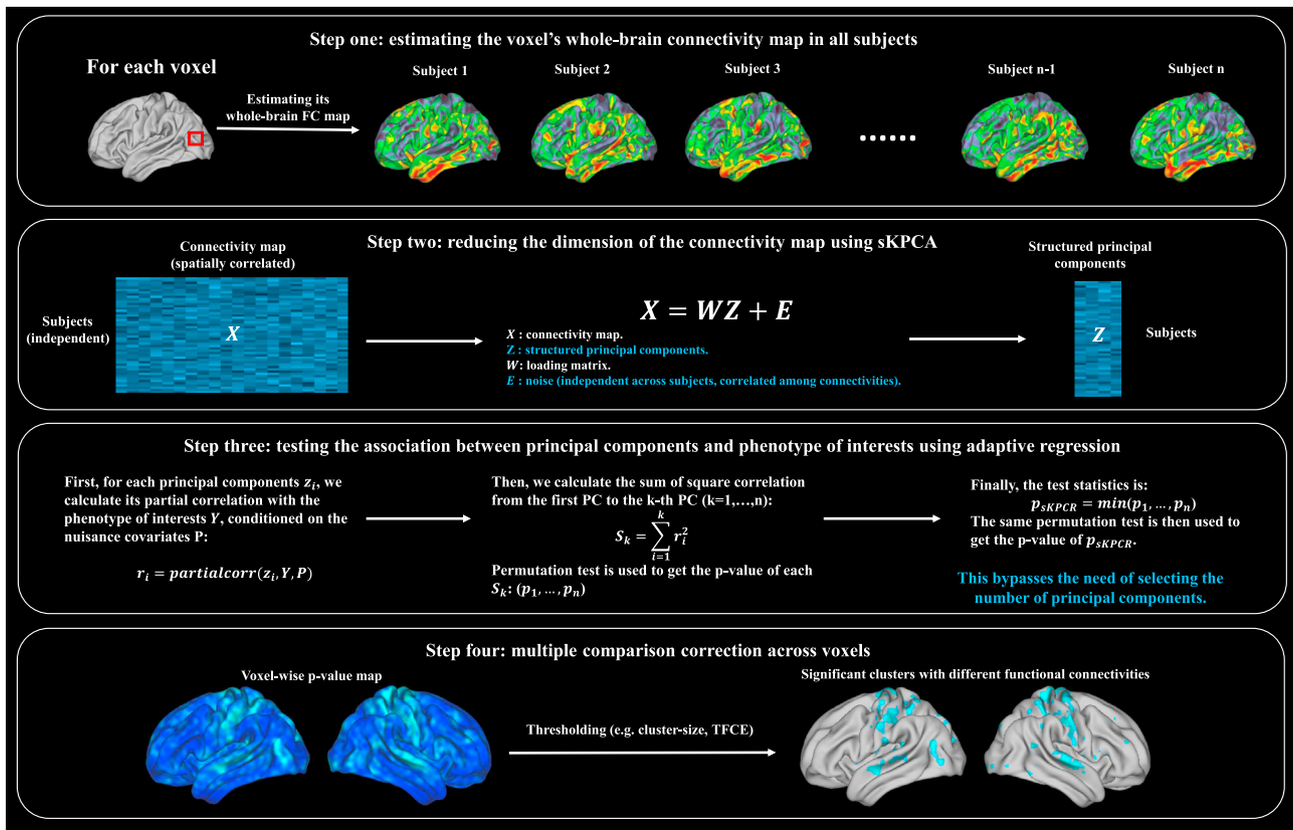
#### 2.1.1. Background: principal component analysis and its probabilistic model

Principal component analysis (PCA) is one of the most popular dimension reduction and feature extraction approaches (Jolliffe, 2002). It is usually defined as the orthogonal projection of a data matrix  $X \in \mathbb{R}^{p \times n}$  onto a low-dimensional space which maximizes the projection variance, where  $n$  is the number of subjects and  $p$  is the number of features. Specifically, suppose that  $X = (x_1, x_2, \dots, x_n)$  has been centered to zero mean by rows. Let  $u_j \in \mathbb{R}^{1 \times p}$  be a  $p$ -dimensional vector that projects each data point  $x_i$  to a scalar value  $t_i = u_j x_i$ , then  $t_i^2 = (u_j x_i)^2$  is proportional to the variance of the projection, and PCA seeks such  $u_j$  that maximizes it. As an optimization problem, this can be written as:

$$\begin{aligned} & \max_{u_j \in \mathbb{R}^p} && u_j X X^T u_j^T \\ \text{subject to} &&& u_j u_j^T = 1, \quad j = 1, 2, \dots, k \\ &&& u_j u_{j'}^T = 0, \quad \forall j' < j \end{aligned} \quad (1)$$

This constrained optimization problem can be solved by an eigenvalue decomposition of the sample covariance matrix  $XX^T \in \mathbb{R}^{p \times p}$ , in which the first  $k$  principal components are exactly the first  $k$  eigenvectors  $U = (u_1, u_2, \dots, u_k)$  of  $XX^T$ . When the number of features  $p$  is larger than the number of subjects  $n$ , note that we can equivalently perform an eigenvalue decomposition on  $X^T X \in \mathbb{R}^{n \times n}$ . We can only extract a maximum of  $\min(n, p)$  number of principal components.

Probabilistic PCA reformulates PCA as the maximum likelihood solution of a probabilistic latent variable model (Tipping and Bishop, 1999; Bishop, 2006), which is closely related to factor analysis (Bartholomew and Knott, 1999; Bishop, 2006) and probabilistic canonical correlation analysis (Bach and Jordan, 2005). In probabilistic PCA, the generative model of the  $i$ -th data point  $x_i \in \mathbb{R}^{p \times 1}$  of a data matrix  $X \in \mathbb{R}^{p \times n}$  is



**Fig. 1.** An overview of the structured kernel principal component regression (sKPCR) in a voxel-level connectome-wide association study. First, for each voxel and each subject, the whole-brain functional connectivity map is computed. Second, a dimension reduction technique, termed structured kernel principal component analysis (sKPCA), is applied to extract important features in this connectivity map, which utilizes the spatial information of functional connectivities. Third, an adaptive regression is fitted to test the association between a phenotype of interest and principal components of this voxel, which automatically selects the optimal number of components. Finally, voxel-wise multiple correction is performed to identify significant clusters.

assumed to be a projection of latent variable  $z_i \in \mathbb{R}^{k \times 1}$  using weight matrix  $W \in \mathbb{R}^{k \times p}$  plus isotropic Gaussian noises  $\varepsilon$  among features:

$$\begin{aligned} z_i &\sim \mathcal{N}(0, I) \\ \varepsilon &\sim \mathcal{N}(0, \sigma^2 I) \\ x_i &= W^T z_i + \varepsilon \end{aligned} \quad (2)$$

To estimate the model parameter  $W$ ,  $\sigma^2$  and latent variable matrix  $Z = (z_1, z_2, \dots, z_n) \in \mathbb{R}^{k \times n}$ , [Tipping and Bishop \(1999\)](#) proposed to maximize the (complete-data) log likelihood with respect to the parameters:  $\max_{W, Z, \sigma^2} \log p(X, Z|W, \sigma^2)$ . They showed that the solution of the above maximum likelihood problem can be obtained by using an expectation-maximization (EM) algorithm or in closed-form as:

$$\begin{aligned} W_{MLE} &= U(\Lambda - \sigma_{MLE}^2 I)^{1/2} \\ \sigma_{MLE}^2 &= \frac{1}{p-k} \sum_{j=k+1}^p \lambda_j \\ Z_{MLE} &= (W_{MLE} W_{MLE}^T + \sigma_{MLE}^2 I)^{-1} W_{MLE} X \end{aligned} \quad (3)$$

where  $\Lambda \in \mathbb{R}^{k \times k}$  is a diagonal matrix, the diagonal elements of which are the first  $k$  eigenvalues of sample covariance matrix  $XX^T$ , i.e.,  $\lambda_j, j = 1, 2, \dots, p$ , and  $U \in \mathbb{R}^{k \times p}$  is the corresponding eigenvectors.

The equivalence between conventional PCA and probabilistic PCA can be seen from the closed-form solution, as demonstrated in Eq. (3). That is, when  $k \rightarrow p$ , then  $\sigma^2 \rightarrow 0$ , the maximum-likelihood estimation (MLE) of latent variable  $Z_{MLE} = (W_{MLE} W_{MLE}^T)^{-1} W_{MLE} X$ , which is an orthogonal projection of the data onto the latent space, enabling recovery

of the standard PCA model ([Tipping and Bishop, 1999](#); [Bishop, 2006](#)).

From the probabilistic interpretation of PCA, we can see that the noise is assumed to be independent between features (and subjects). However, the assumption may break down when analyzing voxel-based functional connectivity data because the noise terms among the connectivities are spatially correlated. In addition, PCA can only perform linear dimension reduction and feature extraction, and many important non-linear factors may be missed by this method. Therefore, we propose a structured kernel principal component analysis (sKPCA) approach in the next section, which can model the spatial structure of the noise and extract both linear and non-linear features.

### 2.1.2. Structured PCA

We propose a framework for structured PCA (sPCA) in this section. It allows structured noise among features and subjects to be modelled. We first describe sPCA from a probabilistic perspective by modelling the noise as a multivariate Gaussian distribution, and then provide an efficient algorithm for estimating the principal components.

To accomplish this, we first introduce the matrix normal distribution  $\mathcal{M}\mathcal{N}_{n \times p}(M, Q, R)$ , which is a generalization of the multivariate normal distribution to matrix-valued random variables  $X \in \mathbb{R}^{p \times n}$ , the probability density function of which is:  $p(X|M, Q, R) = \frac{\exp[-\frac{1}{2} \text{tr}(Q^{-1}(X-M)^T R^{-1}(X-M))]}{(2\pi)^{np/2} |Q|^{n/2} |R|^{p/2}}$ , where  $M \in \mathbb{R}^{p \times n}$  is the mean, and  $R \in \mathbb{R}^{p \times p}$  and  $Q \in \mathbb{Q}^{n \times n}$  are the covariance matrix of the rows and columns of  $X$ . The connection between the matrix normal distribution and multivariate normal distribution is:  $\text{vec}(X) \sim \mathcal{N}(\text{vec}(M), Q \otimes R)$ , where  $\otimes$  denotes the Kronecker product and  $\text{vec}$  denotes the vectorization  $M$ .

We can represent probabilistic PCA using the matrix normal distribution as:

$$X = W^T Z + E; E \sim \mathcal{M}\mathcal{N}_{p,n}^r(0, \sigma^2 I, I) \tag{4}$$

Here, both the rows (e.g. features) and columns (e.g. subjects) of the noise matrix are assumed to be independent from each other. However, for neuroimaging data, noises among voxels/vertices are known to be smoothly correlated with each other. Therefore, our sPCA model is a generalization of the probabilistic PCA model, which allows two-way dependence between noise terms:

$$X = W^T Z + E; E \sim \mathcal{M}\mathcal{N}_{p,n}^r(0, R, Q) \tag{5}$$

where  $W$  is the weight matrix and  $Z$  is the latent variable matrix in accordance with probabilistic PCA.

In this model, the  $Q$  and  $R$  do not need to be estimated from the data; however, they do need to be prespecified based on the known topological structure of the data (see section 2.2 for details). Then, we can still use the MLE approach to estimate the  $W$  and  $Z$  in (5):

$$\begin{aligned} \max_{W,Z} \log P(X, Z|W) &= \max_{W,Z} -\frac{1}{2} \text{tr} [Q^{-1}(X - W^T Z)^T R^{-1}(X - W^T Z)] - \frac{1}{2} \text{tr}(Z^T Z) + \text{Const} \\ &= \max_{\tilde{W}, \tilde{Z}} -\frac{1}{2} \text{tr} [(\tilde{R} X \tilde{Q} - \tilde{R} W^T Z \tilde{Q})^T (\tilde{R} X \tilde{Q} - \tilde{R} W^T Z \tilde{Q})] - \frac{1}{2} \text{tr}(Z^T Z) + \text{Const} \\ &= \max_{\tilde{W}, \tilde{Z}} -\frac{1}{2} \text{tr} [(\tilde{X} - \tilde{W}^T \tilde{Z})^T (\tilde{X} - \tilde{W}^T \tilde{Z})] - \frac{1}{2} \text{tr}(Z^T Z) + \text{Const} \end{aligned} \tag{6}$$

where we have decomposed  $Q^{-1} = \tilde{Q}\tilde{Q}^T$  and  $R^{-1} = \tilde{R}\tilde{R}^T$ , and  $\tilde{X} = \tilde{R}X\tilde{Q}$ ,  $\tilde{W} = \tilde{R}W$  and  $\tilde{Z} = Z\tilde{Q}$ . Therefore, the sPCA problem (6) is equivalent to the standard PCA or probabilistic PCA problems using the ‘weighted’ data matrix  $\tilde{X}$  (Escoufier, 1977; Allen et al., 2014; Zhu et al., 2017), where the weights are learned from the external information, i.e., the spatial, temporal or population structures of data, as:

$$\begin{aligned} \max_{u_j \in \mathbb{R}^{1 \times p}} \quad & u_j R^{-1} X Q^{-1} X^T R^{-1} u_j^T \\ \text{subject to} \quad & u_j R^{-1} u_j^T = 1, j = 1, 2, \dots, k \\ & u_j R^{-1} u_j^T = 0, \forall j' < j \end{aligned} \tag{7}$$

Therefore, the principal components  $U$  (or  $W$ ) can be obtained by a simple eigenvalue decomposition on the matrix  $R^{-1}XQ^{-1}X^TR^{-1} \in \mathbb{R}^{p \times p}$ , or when  $n < p$ , on the matrix  $Q^{-1}X^TR^{-1}XQ^{-1} \in \mathbb{R}^{n \times n}$ .

### 2.1.3. Structured kernel PCA

The sPCA model can be further generalized to perform non-linear dimension reduction by using kernel tricks. Specifically, let  $\tilde{x}_i$  be the  $i$ -th ‘weighted sample’, i.e., the  $i$ -th column of  $\tilde{X} = \tilde{R}X\tilde{Q}$ , we first perform a non-linear mapping of the sample  $\tilde{x}_i$  to the high dimensional feature space as  $\tilde{x}_i \rightarrow \Phi(\tilde{x}_i)$ . Now, we assume that each  $\Phi(\tilde{x}_i)$  has been mean centered in the feature space and we will return to this point later. We can perform a PCA in the mapped high-dimensional feature space by maximizing the projected variance as:

$$\max_{u_j \in \mathbb{R}^{1 \times n}} u_j \Phi(\tilde{X})^T \Phi(\tilde{X}) u_j^T, j = 1, 2, \dots, n \tag{8}$$

Similar to the kernel principal component analysis (Schölkopf et al., 1997), the optimization problem (8) can be solved by first performing a mean normalization of the kernel matrix  $K \in \mathbb{R}^{n \times n}$ , where  $K_{ij} =$

$\Phi(\tilde{x}_i)^T \Phi(\tilde{x}_j)$ :  $\tilde{K} = K - I_n K - K I_n + I_n K I_n$ , where  $I_n$  is an  $n \times n$  matrix in which each element takes the value  $1/n$ . Then, we solve the eigenvalue problem:

$$n^{-1} \tilde{K} u_j^T = \lambda u_j^T, j = 1, 2, \dots, n$$

and obtain  $n$  eigenvalues in a descending order as  $(\lambda_1, \dots, \lambda_n)$  and the corresponding eigenvectors  $(u_1, \dots, u_n)$ . The  $k$ -th principal component is the  $k$ -th eigenvector  $u_k$ .

Similar to most other kernel-based approaches, all the computations can be expressed in the form of the kernel matrix. When using the linear kernel, the sKPCA is exactly the same as sPCA. In addition, for many commonly used kernels, we do not even need to estimate  $\tilde{Q}$  and  $\tilde{R}$ . For example, we can calculate  $X^* = Q^{-1}X R^{-1}X^T Q^{-1}$ , and the polynomial kernel can be calculated as  $K_{ij} = (aX_{ij}^* + b)^c$ , the sigmoid kernel can be calculated as  $K_{ij} = \tanh(aX_{ij}^* + b)$ , and the Gaussian kernel can be calculated as  $K_{ij} = \exp(-\frac{\|\tilde{x}_i - \tilde{x}_j\|^2}{2\sigma^2}) = \exp[-(X_{ii}^* - 2X_{ij}^* + X_{jj}^*)/2\sigma^2]$ .

### 2.2. The choice of sKPCA parameters

Many methods have been developed to determine the number of principal components for conventional PCA, such as the ratio estimator (Lam and Yao, 2012; Li et al., 2017), the information criteria approaches (Bai and Ng, 2002, 2007), the distribution-based approach (Choi et al., 2014) or just by the amount of variance explained (e.g., 90%). Although these methods can be easily extended to the current sKPCA framework, we have found that none of them works optimally in the subsequent association tests, which is our main goal in this paper. Therefore, we developed a novel adaptive regression approach in the next section, in order to address the problem of selecting the number of principal components in the association study.

Many possible choices are available for the covariance matrix (Allen et al., 2014; Ramsay, 2006), but we will introduce and compare just three in this paper. The first one is the Graph Laplacian (GL) operator, which has been widely used in Bayesian task-activation studies (Penny et al., 2005; Flandin and Penny, 2007; Sidén et al., 2017). It is also known as the inverse covariance operator (Allen et al., 2014; Ramsay, 2006). To define the GL operator  $G$ , we first define the feature-feature adjacency matrix  $A$  as a binary matrix such that  $a_{ij} = 1$  if the spatial distance between feature  $i$  and  $j$  ( $i \neq j$ ) equals one (or feature  $i$  and  $j$  are spatial neighbours) and  $a_{ij} = 0$  otherwise. Based on  $A$ , we can define  $G$  as, for feature  $i$  and  $j$ ,  $G_{ii} = \sum_{i \neq j} a_{ij}$  and  $G_{ij} = -a_{ij}$  if  $i \neq j$ . The second one is the normalized Graph Laplacian (NGL) operator  $G^*$ . Based on  $A$ , it is defined as  $G_{ii}^* = 1$  and  $G_{ij}^* = -\frac{1}{\sqrt{\sum_{l \neq i} a_{il} \sum_{l \neq j} a_{jl}}}$  if  $i \neq j$ . The third one is called the

Gaussian random field operator. It assumes that the noise covariance between two features is a functional of their spatial distance:  $\Sigma_{ij} = \exp\left(-\frac{\|X_i - X_j\|_2}{2\sigma^2}\right)$  where  $\|X_i - X_j\|_2$  represents the spatial Euclidian

distance between feature  $i$  and feature  $j$  in the volume space, and  $\|X_i - X_j\|_2$  can also be the geodesic distance in surface space. The  $\sigma$  can be specified based on the estimated Full Width at Half Maximum (FWHM) of the images using the relationship  $\text{FWHM} = 2\sqrt{2 \log 2} \sigma$ . We will show that sKPCR is very stable for selecting different covariance operators and that the GL operator has a slightly higher power. Therefore, it is used in all our analyses.

### 2.3. Structured kernel principal component regression (sKPCR) for identifying connectome-wide associations

#### 2.3.1. Overview

We extend sKPCA to structured kernel principal component regression (sKPCR) to identify connectome-wide associations. In our study, the individual-level brain functional network is estimated by the Fisher's Z transformed Pearson correlation coefficient between every pair of voxels' BOLD signal time series. Let  $n$  be the number of subjects in a study, and  $p$  be the number of voxels; as such, the total number of functional connectivities in each individual's brain network is  $p(p-1)/2$ , and there are  $p-1$  functional connectivities connecting a voxel to all other voxels across the whole brain. Let  $Y \in \mathbb{R}^{n \times 1}$  be the phenotype of interest of  $n$  subjects (e.g. disease status, clinical symptoms) and  $Z \in \mathbb{R}^{n \times q}$  be the nuisance covariates (e.g. age, gender, motion terms). Our aim is to test, for each voxel, whether the phenotype of interest  $Y$  is associated with the voxel's whole-brain functional connectivity pattern  $X \in \mathbb{R}^{n \times (p-1)}$ , conditioned on the nuisance covariates  $Z$ . Since connectivity is of ultra-high dimensionality (e.g. for each voxel, there are  $10^4$  to  $10^5$  whole-brain functional connectivities, but only a few hundred samples), the basic idea of sKPCR is to (1) extract important low-dimensional features (principal components) in the data  $X$  by sKPCA, and then (2) test the association between the extracted principal components  $U = (u_1, \dots, u_k)$  and the phenotype of interest  $Y$ .

#### 2.3.2. An adaptive regression model

To test associations, we propose a novel adaptive regression approach which can estimate a single p-value per voxel to summarize the overall significance of the association. Traditionally, we would manually select the top  $k$  principal components and then use a general linear model with  $F$  statistic for statistical testing. However, we found the pre-specification of  $k$  to be very difficult, and the top principal components may not always explain the phenotype of interest. Therefore, we propose a new approach which is able to adaptively choose the optimal number of principal components to include in the model, one that is sufficiently robust to include noise components. The idea is similar to many other adaptive test approaches widely used in the neuroimaging (Kim et al., 2015) and genetics fields (Pan et al., 2014; Lee et al., 2012).

In detail, let  $r_i$  be the partial correlation between a principal component  $u_i$  and a phenotype  $Y$  conditioned on covariates  $Z$ , then we define a score  $S_k$  to measure the overall correlations between  $k$  ( $k = 1, 2, \dots, K$ ) extracted components and the phenotype as:

$$S_k = \sum_{i=1}^k r_i^2$$

We can get a score vector  $S = (S_1, S_2, \dots, S_K)$ . Using a non-parametric permutation approach, we can get the p-value of each score by permuting the phenotype  $Y$  and recalculating the 'null' score vector  $M$  times. That is, let  $\tilde{Y}^j$  be a randomly permuted phenotype in the  $j$ -th permutation. We first calculate the partial correlation between  $\tilde{Y}^j$  and  $u_i$  conditioned on covariates  $Z$  to get the permuted coefficient  $\tilde{r}_i^j$ . Then, as above, we calculate the score as

$$\tilde{S}_k^j = \sum_{i=1}^k (\tilde{r}_i^j)^2$$

With a total of  $M$  permutations, we can get a vector of null score  $\tilde{S}_k = (\tilde{S}_k^1, \tilde{S}_k^2, \dots, \tilde{S}_k^M)$ . Therefore, the p-value of score  $S_k$  can be estimated non-parametrically as:

$$p_k = \frac{\#\{\tilde{S}_k \geq S_k\} + 1}{M + 1}$$

where  $\#\{A \geq a\}$  denotes the number of times the elements in vector  $A$  is larger than a number  $a$ . After the above steps, we can get the p-values of the  $K$  scores  $S$ , denoting them as  $(p_1, p_2, \dots, p_K)$ . The above steps can be computationally very efficient by using a simple matrix computation strategy.

Now, we define our test statistic as the smallest p-values in  $(p_1, p_2, \dots, p_K)$ :

$$T_{\text{sKPCR}} = \min(p_1, p_2, \dots, p_K)$$

Note that  $T_{\text{sKPCR}}$  is a *not* a p-value, because its null distribution is no longer subjected to a uniform distribution. Therefore, its statistical significance should be estimated using non-parametric permutation again. However, it is interesting to note that we do not need to run another set of permutations, but rather simultaneously estimate the null distribution of  $T_{\text{sKPCR}}$  using the above permutations, which have been used to calculate  $(p_1, p_2, \dots, p_K)$ . Specifically, the permutation empirical p-value of the  $k$ -th score in the  $j$ -th permutation is:

$$\tilde{p}_k^j = \frac{\#\{\tilde{S}_k \geq \tilde{S}_k^j\}}{M}$$

Therefore, the most significant p-value across  $K$  scores in the  $j$ -th permutation is:

$$\tilde{T}_{\text{sKPCR}}^j = \min(\tilde{p}_1^j, \tilde{p}_2^j, \dots, \tilde{p}_K^j)$$

Thus, for all the  $M$  permutations, we get  $\tilde{T}_{\text{sKPCR}} = (\tilde{T}_{\text{sKPCR}}^1, \tilde{T}_{\text{sKPCR}}^2, \dots, \tilde{T}_{\text{sKPCR}}^M)$ . Finally, the p-value of our test statistic  $T_{\text{sKPCR}}$  can be estimated as:

$$p(\tilde{T}_{\text{sKPCR}}) = \frac{\#\{\tilde{T}_{\text{sKPCR}} \geq T_{\text{sKPCR}}\} + 1}{M + 1}$$

Note also that this approach can provide an exact control of false-positive rate. However, for the general linear model approach, the p-value of a  $F$ -test or likelihood-ratio test may not provide a valid p-value when the number of components is comparable to the number of subjects (Sur et al., 2017).

#### 2.3.3. Multiple comparison correction

After getting the voxel-wise p-values, we can use a nonparametric permutation approach (Nichols and Holmes, 2002) or false-discovery rate method (Benjamini and Hochberg, 1995) to perform multiple comparison correction. For permutation-based approaches, we can still use the same set of permutations above to perform topological inference, including peak-level inference (Worsley et al., 1996), cluster-size inference (Friston et al., 1994), cluster-mass inference (Zhang et al., 2009), and threshold-free cluster enhancement (Smith and Nichols, 2009).

### 3. Evaluating sKPCR using simulation studies: false-positive rate, power and robustness

#### 3.1. Data

We use two resting-state fMRI datasets to evaluate different methods, including 281 subjects from the Southwest University (SWU) dataset in the International Data-sharing Initiative (IDNI, [http://fcon1000.projects.nitrc.org/indi/retro/southwestuni\\_qiu\\_index.html](http://fcon1000.projects.nitrc.org/indi/retro/southwestuni_qiu_index.html)), and 150 subjects from the Human Connectome Project (HCP\_REST1\_LR, <https://www.humanconnectome.org/>). All subjects were healthy adults with

similar demographic information.

The data from SWU were preprocessed using a standard volume-based fMRI pipeline (code can be download from <https://github.com/weikangong/Resting-state-fMRI-preprocessing>). For each individual, the preprocessing steps include: slice timing correction (FSL slicetimer), motion correction (FSL mcflirt), spatial smoothing by a 3D Gaussian kernel (FWHM = 6 mm), despiking motion artifacts using the BrainWavelet Toolbox (Patel et al., 2014), registering to  $4 \times 4 \times 4 \text{ mm}^3$  standard space by first aligning the functional image to the individual T1 structural image using boundary based registration (Greve and Fischl, 2009), and then to standard space using FSL's linear and non-linear registration tool (FSL flirt and fnirt), regressing out nuisance covariates including 12 head motion parameters (6 head motion parameters and their corresponding temporal derivatives), white matter signal, cerebrospinal fluid signal and global signal, band-pass filtering (0.01–0.1 Hz) using AFNI (3dTproject). All the images were manually checked to ensure successful preprocessing. Finally, 14364 grey matter voxels located in each subject's cerebrum were extracted for the subsequent analysis.

The data from HCP-S900 were preprocessed using the *fMRISurface* minimal preprocessing pipeline (Glasser et al., 2013; Smith et al., 2013). The basic steps included: correcting for spatial distortions caused by gradient nonlinearity, correcting for head motion by registration to the single band reference image, correcting for  $B_0$  distortion, and registering to the T1w structural image. The global intensity was normalized. Then, independent component analysis (ICA) was run using MELODIC with automatic dimensionality estimation (Beckmann and Smith, 2004). These components were fed into FIX (Salimi-Khorshidi et al., 2014), which classified components into 'good' vs. 'bad'. Bad components were removed from the data. From this resulting volume time-series, the data were mapped onto the standard 32k Conte69 cortical surface using the Multimodal Surface Matching approach (MSMALL pipeline (Robinson et al., 2014)). Finally, the Gaussian spatial smoothing was carried out on the cortical surface with a Full-Width at Half Maximum of 4 mm. In our analysis, BOLD time series of 32492 cortical vertices from each subject's left cortical surface were used.

### 3.2. Type I error rate evaluation

To evaluate whether the proposed sKPCR approach could control the type I error rate, we evaluated whether it had a nominal false-positive rate when comparing the connectome of two groups of healthy subjects with similar demographic information (Eklund et al., 2016). If a method can provide a valid control of type I error rate, the observed false-positive rate will be around its nominal level (e.g. 0.05). Specifically, first, a voxel was randomly selected, and functional connectivities between it and all other voxels across the whole brain were estimated for every subject. Second, subjects were randomly divided into two groups, and sKPCR with 5 different types of kernel was then applied to test whether this voxel showed differences between the two randomly assigned groups. This step resulted in one p-value for sKPCR per kernel. Third, the above two steps were repeated for 1000 times, and the observed false-positive rate was estimated as the proportion of times the p-value was below 0.05. Some commonly used kernels we evaluated here were a linear kernel, polynomial kernel with degree 2,3,4,5 and a Gaussian kernel with  $\sigma$  parameter equaling the middle of the Euclidian distance among data points (Brown et al., 2000).

### 3.3. Comparing the statistical power of detecting linear signals with other methods

In this simulation, we considered methods for detecting linear signals. We compared the linear sKPCR with 5 other approaches, including a connectivity-wise general linear model (GLM) approach controlling the family-wise error rate (i.e., SPU(Inf) approach (Kim et al., 2014)); multivariate distance matrix regression (MDMR) (Shehzad et al., 2014); and adaptive sum of powered score (aSPU) and extensions (i.e., SPU(1),

SPU(2), aSPU (Kim et al., 2014)). All of these approaches can produce voxel-wise p-value maps based on the rsfMRI data. A brief description of these methods and their parameter settings are shown in the Supplementary Material.

In our simulation, first, one voxel was randomly selected, and functional connectivities between it and all other voxels across the whole brain were estimated for every subject, and they were normalized to zero mean unit variance. Second, signals were then randomly added to a subset of functional connectivities (proportion of null functional connectivity  $\rho$ ). For linear signals, we simulated a phenotype of interest  $y$  which was linearly correlated with some functional connectivities  $x$ . This was achieved by first simulating new functional connectivities as  $x_{new} = \gamma y + x$ , and then normalizing them again to zero mean unit variance. The signal-to-noise ratio  $\gamma^2$ , which was defined as the ratio of signal variance and noise variance, varied from 0 (no signal) to 0.25 in our simulation. Each method was then applied to test whether the overall connectivity pattern was associated with the signal  $y$ . Third, the above two steps were repeated 1000 times, and the empirical power was estimated by the proportion of times the p-value was below 0.05.

### 3.4. Comparing the statistical power of detecting nonlinear signals with other methods

For non-linear signals, we compared our approach with 2 other methods, including kernel principal component regression (KPCR) (Schölkopf et al., 1997) and least-squares kernel machine (LSKM) (Liu et al., 2007; Ge et al., 2012). A brief description of these methods and their parameter settings are shown in the Supplementary Material.

This simulation is similar to the above one. The only difference is the way of generating the nonlinear signals. This was achieved by simulating new functional connectivities  $x_{new}$  as  $x_{new} = \gamma y^d + x$  where  $d$  is the polynomial degree. Theoretically, therefore, methods using a polynomial kernel with corresponding degree should achieve the highest power.

### 3.5. The robustness of sKPCR when the kernel is misspecified

To evaluate whether sKPCR is robust when the kernel was misspecified, we also used sKPCR, KPCR and LSKM with linear kernels for signal detection. The simulation procedures were exactly the same as those described in Section 3.4, but we used the wrong kernels to detect signals.

### 3.6. The robustness of sKPCR when the number of principal components is misspecified

As illustrated in section 2.3, the proposed adaptive regression approach, which was used to detect association after sKPCA dimension reduction, was robust to the misspecification of number of principle components. To demonstrate this, we conducted a simulation study to compare it with the traditional general linear model approach.

We assumed a total of 200 subjects and we totally extracted 100 principal components. As the components are not correlated with each other, we first simulated independent Gaussian white noise data, which formed a  $200 \times 100$  matrix  $X$ . The first 10 components were assumed to be correlated with a phenotype  $y$ , and the 11-th to 100-th components were assumed to be noise components. Therefore, we added  $\beta y$  to the first 10 components, where  $\beta$  could be treated as the effect size. In our experiment,  $\beta = 0, 0.03, 0.06, 0.09, 0.12, 0.15$ . We tested the association between the first  $i$  columns of  $X$  and  $y$  using either the adaptive regression or the linear regression model with the  $F$  statistic. The above procedures were repeated 1000 times for different effect sizes and numbers of components. We compared the power of the two methods when the number of principal components were misspecified by adding more noise components to the model.

### 3.7. The robustness of sKPCR under different covariance operators

Following the same simulation procedures as those detailed in section 3.3, we compared the power of linear sKPCR when using different covariance operators. The covariance operators we tested included GL, NGL, Gaussian with  $\sigma^2 = 2, 4, 6$  voxels (Gaussian (2), Gaussian (4), Gaussian (6)).

## 4. Evaluating sKPCR in real data: brain-wide associations of the schizophrenic connectome

### 4.1. Data

Four resting-state fMRI datasets were used here: Taiwan, Centers of Biomedical Research Excellence (COBRE [http://fcon\\_1000.projects.nitrc.org/indi/retro/cobre.html](http://fcon_1000.projects.nitrc.org/indi/retro/cobre.html)), BrainGluSchi (<http://schizconnect.org/>), and NMorphCH (<http://schizconnect.org/>). All of them are datasets with schizophrenia patients and matched healthy controls. The demographic information is shown in Table 1. Resting-state fMRI data were preprocessed using the same pipeline as the SWU dataset (code can be download from <https://github.com/weikangong/Resting-state-fMRI-preprocessing>). Finally, 18757 voxels located in each subject's cerebral regions were extracted for the subsequent analysis.

### 4.2. Between-sites reproducibility

We applied sKPCR with a linear kernel to identify voxels with significantly altered connectivities in each schizophrenia dataset separately. At the same time, five other methods, including the univariate approach, MDMR, SPU(1), SPU(2) and aSPU, were used for comparison. To compare the between-sites reproducibility of each method, they were applied to analyse the four schizophrenia datasets separately, and the Dice Coefficient (DC) between the resulting voxel-wise p-value maps of two sites was calculated. DC is defined as

$$DC = \frac{2|E_1 \cap E_2|}{|E_1| + |E_2|}$$

where  $|E_i|$  is the number of significant voxels in the  $i$ -th experiment. We will mainly report the DC with  $p = 0.001$  as the voxel-wise significance threshold, as it is widely used (Woo et al., 2014). We expect that a better approach has a larger overlap.

### 4.3. Evidence from the literature: overlaps with existing meta-analysis findings

We also evaluated whether the above results were consistent with existing meta-analysis findings reported in the Neurosynth database (Yarkoni et al., 2011). We searched for the term 'schizophrenia' on the Neurosynth website (<http://neurosynth.org/analyses/terms/schizophrenia/>), and downloaded the default forward inference map (FDR < 0.01). We calculated the DC between the p-value map of each approach in each dataset and the Neurosynth schizophrenia forward inference map. We will mainly report the DC with  $p = 0.001$  as the voxel-wise significance threshold, as it is widely used (Woo et al., 2014).

## 5. Results

### 5.1. Evaluating sKPCR using simulation studies

#### 5.1.1. Type I error rate evaluation

We first evaluate whether sKPCR could control the type I error rate using different kernels in different types of data. We simulated a case-control study in the absence of any group difference (see Section 3.2), and sKPCR was applied to detect signals. The results show that the proposed approach can control the false-positive rate appropriately using a wide range of kernels (linear, polynomial and Gaussian) in both volume-based and surface-based fMRI data (Fig. 2), because the observed false-positive rates are similar to their theoretical nominal level at 0.05.

#### 5.1.2. Comparing the statistical power of detecting linear signals with other methods

Fig. 3 shows the results of comparing the power of different methods when the true signal is linear, i.e. some functional connectivities are linearly correlated with a simulated phenotype of interest, in volume-based fMRI data (see Section 3.3). Similarly, Fig. 4 shows the results of the same simulation using surface-based fMRI data.

For both volume-based and surface-based data, it can be clearly seen that the proposed sKPCR method with a linear kernel always has the highest power in different situations (different signal-to-noise ratios and proportions of non-null connectivities). The univariate approach and aSPU have similar power in different situations. The performance of MDMR and SPU(2) are similar to that of aSPU and the univariate method when the number of non-null functional connectivities is large (e.g., 20% non-null). However, the power of MDMR and SPU(2) decreases dramatically when only a few non-null connectivities exist (e.g., 1% non-null). SPU(1) performs the worst in these simulations. In addition, the power of sKPCR displays a larger gap with other approaches in surface-based fMRI data compared to volume-based fMRI data.

To get a more intuitive understanding of why sKPCR had a better performance, we simulated a case-control study with some of the functional connectivities in one group having a higher mean than another group with the same strategies as the above simulation. We applied sKPCA with a linear kernel and PCA to the simulated data and extracted the top 4 principal components. For each method, we plotted each pair of principal components in a 2D figure and used different colors to distinguish the two groups. As can be clearly observed from Fig. 5, sKPCA (top row) has much better performance because the case and control groups are better separated than with PCA dimension reduction (bottom row).

**Table 1**

Demographic information of subjects used in simulations (Southwest University and Human Connectome Project datasets) and real data analysis (4 schizophrenia datasets: COBRE, Taiwan, NMorphCH and BrainGluSchi).

Dataset	Group	# Subjects	Age (mean $\pm$ std)	Gender (M/F)	mean FD
Southwest University	Control	281	19.7 $\pm$ 0.85	0/281	0.09 $\pm$ 0.03
	HCP	150	28.8 $\pm$ 3.71	64/86	0.08 $\pm$ 0.02
COBRE	Control	72	35.6 $\pm$ 11.7	50/22	0.21 $\pm$ 0.11
	Patient	58	36.7 $\pm$ 13.5	49/9	0.24 $\pm$ 0.11
	Statistic (p-value)		0.61	0.07	0.07
Taiwan	Control	136	44.1 $\pm$ 12.0	57/79	0.11 $\pm$ 0.05
	Patient	123	44.0 $\pm$ 11.3	51/72	0.10 $\pm$ 0.07
	Statistic (p-value)		0.79	1	0.66
NMorphCH	Control	39	30.6 $\pm$ 8.1	19/20	0.13 $\pm$ 0.07
	Patient	42	32.8 $\pm$ 6.9	30/12	0.18 $\pm$ 0.11
	Statistic (p-value)		0.20	0.04	0.01
BrainGluSchi	Control	76	38.7 $\pm$ 12.4	49/27	0.23 $\pm$ 0.11
	Patient	60	34.5 $\pm$ 13.6	55/5	0.21 $\pm$ 0.11
	Statistic (p-value)		0.06	1.9e-4	0.54

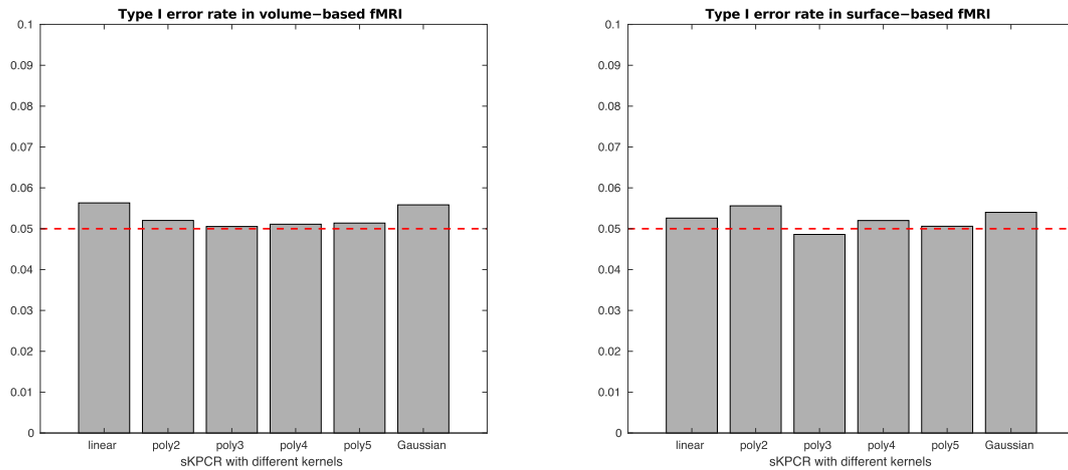


Fig. 2. Type I error rate of structured kernel principal component regression (sKPCR) estimated in the volume-based and surface-based fMRI data using different kernels (linear, 2,3,4,5-degree polynomial and Gaussian). The results show that the method can control the type I error rate at its nominal level (approximately 5%, dashed red line).

### Volume-based fMRI with linear signal

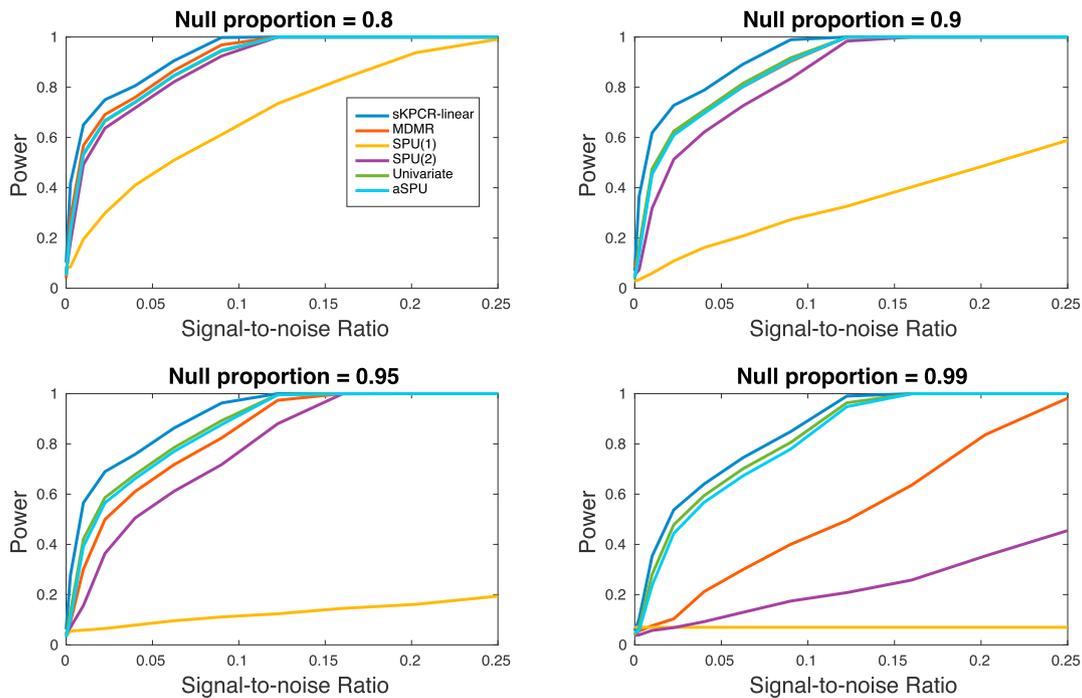


Fig. 3. Comparisons of the power of detecting linear signals with different methods using simulations in volume-based fMRI data. In this simulation, some functional connectivities were simulated to linearly correlated with a phenotype of interest. Each figure plots the power curves of 6 different methods with different signal-to-noise ratios (0–0.25) and with different null connectivity proportions (0.8–0.99). The results show that the proposed sKPCR approach with a linear kernel has the highest power.

#### 5.1.3. Comparing the statistical power of detecting nonlinear signals with other methods

Fig. 6 shows the results of comparing the power of different methods when the true signal is nonlinear, i.e. a subset of functional connectivities are nonlinearly correlated with a simulated phenotype of interest in volume-based fMRI data (see Section 3.4). Similarly, Fig. 7 reports the results of the same simulation using surface-based fMRI data.

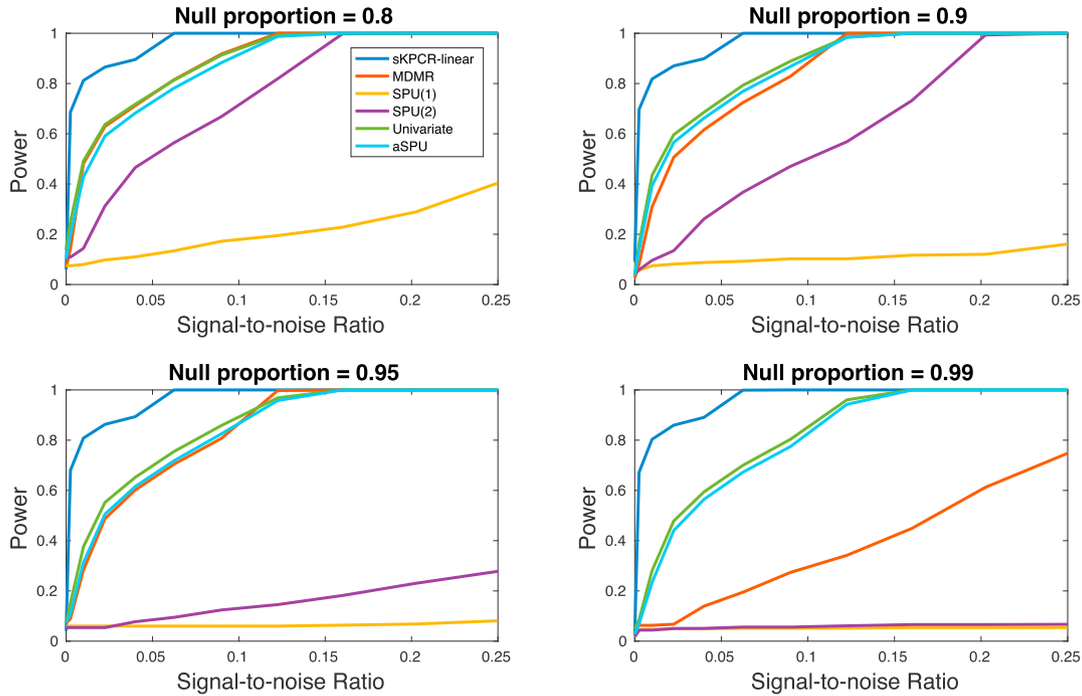
For both volume-based and surface-based data, it can be clearly seen that the proposed sKPCR method with a polynomial or linear kernel always has the highest power in different situations (different signal-to-

noise ratios and polynomial degrees). When the polynomial degree is an even number, sKPCR with a corresponding polynomial kernel outperforms all other approaches, but when the polynomial degree is an odd number, sKPCR with a corresponding polynomial kernel and linear kernel have the similar high power. This is because the polynomial signals with odd degree are quite similar to linear signals (e.g., consider  $y = x^3 + \epsilon$  and  $y = x + \epsilon$ ).

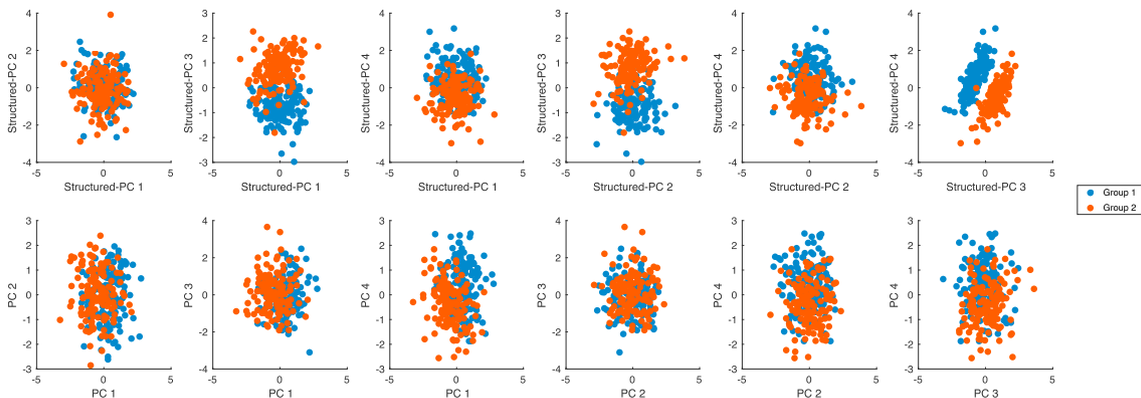
#### 5.1.4. The performance of sKPCR when the kernel is misspecified

Again, when the true signals are nonlinear, we can see from Figs. 6

### Surface-based fMRI with linear signal



**Fig. 4.** Comparisons of the power of detecting *linear* signals with different methods using simulations in *surface-based fMRI data*. In this simulation, some functional connectivities were simulated to linearly correlated with a phenotype of interest. Each figure plots the power curves of 6 different methods with different signal-to-noise ratio (0–0.25) and with different null connectivity proportions (0.8–0.99). The results show that the proposed sKPCR approach with a linear kernel has the highest power.



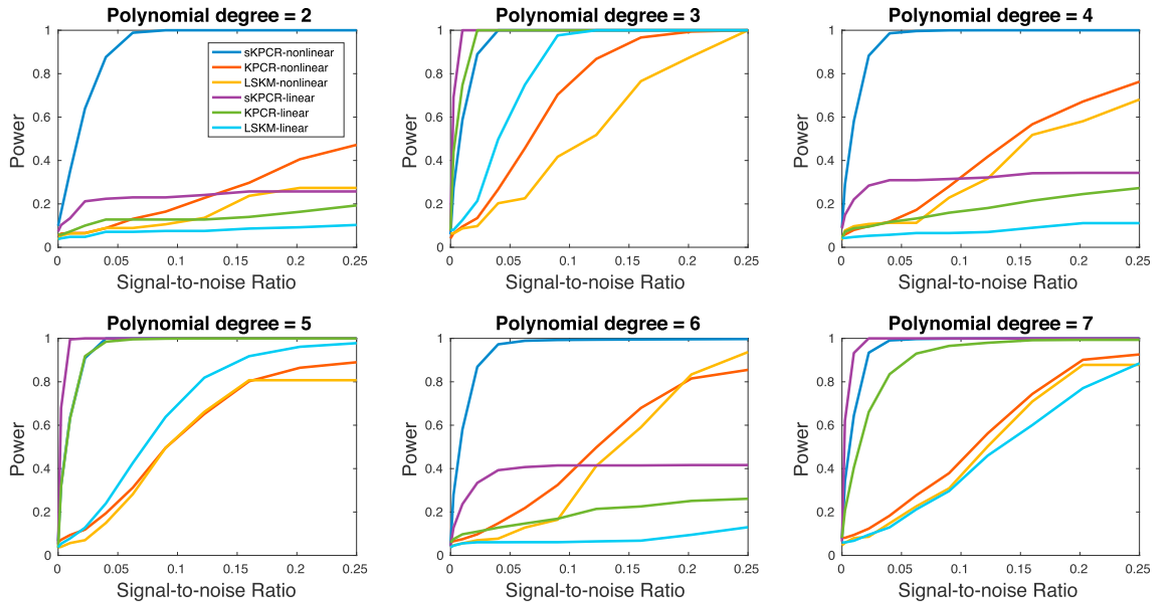
**Fig. 5.** An illustrative example of comparing structured kernel principal component analysis (sKPCA), which takes connectivity structure into account, with the original principal component regression (PCA) using simulated connectivity data. In this simulation, some functional connectivities connecting one voxel to other voxels in the brain were simulated to be higher in one group compared to another group. We applied these two methods to this dataset to learn the first 4 principal components. The top row shows the scatter plots of each pair of PCs in the two groups (blue and red) using sKPCA, while the bottom row shows the results for PCA.

and 7 that methods using a linear kernel usually have decreased power compared with the corresponding nonlinear kernel. This highlights the importance of specifying correct kernels in the analysis to achieve optimal power. However, this does not mean that our method is sensitive to the choice of kernels. In practice, we can run sKPCR with different kernels, and for each kernel, we will get a voxel-wise p-value map which reflects the strength of different types of association signals. In addition, although the power of sKPCR with a linear kernel decreases when the true signals are nonlinear, we can see that it is still higher than many other approaches, even with the correct nonlinear kernels. This may result from its effective modelling of the spatial structure of the data.

#### 5.1.5. The robustness of sKPCR when the number of principal components is misspecified

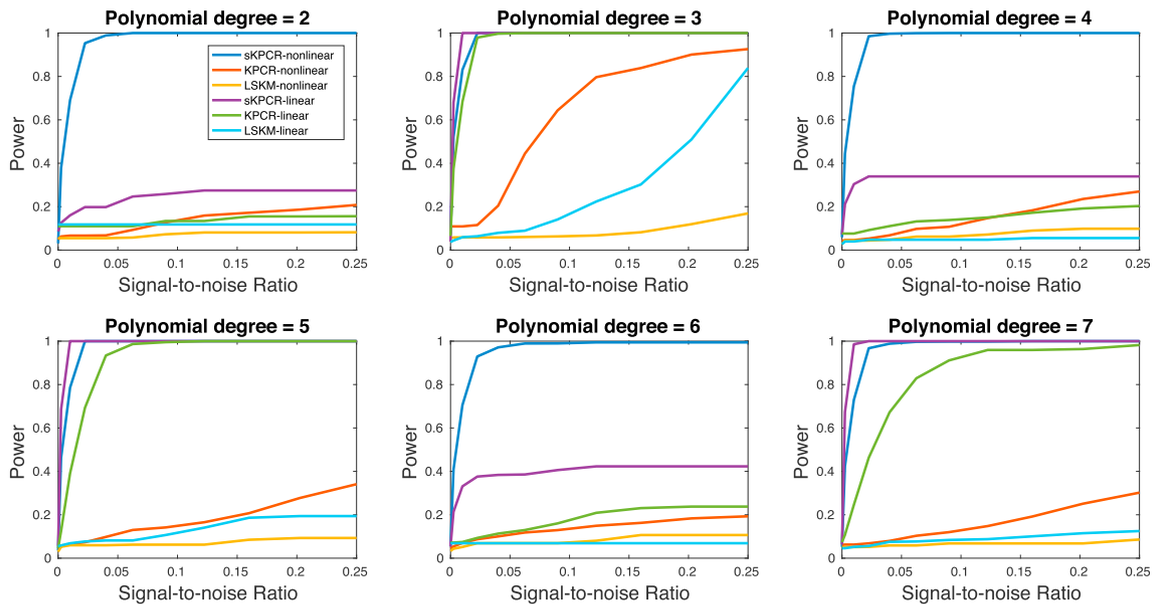
Fig. 8 shows the results of a power comparison between adaptive regression and the general linear model in association testing with different numbers of noise components and signal-to-noise ratios (see Section 3.6). It can be seen that adaptive regression is robust to the misspecification of components because its power does not change much, even when an increasing number of noise components are added to the model. However, the power of the general linear model decreased dramatically when an increasing number of noise components are included in the model.

### Volume-based fMRI with non-linear signal (polynomial relationship)



**Fig. 6.** Comparisons of the power of detecting *nonlinear* signals with different methods using simulations in *volume-based fMRI data*. In this simulation, polynomial (nonlinear) relationships were simulated to exist between some functional connectivities and a phenotype of interest with polynomial degree ranging from 2 to 7 in each figure (e.g.  $y = x^d$ , where  $y$  is a phenotype of interest,  $x$  is a functional connectivity and  $d = 2, \dots, 7$ ). Each figure plots the power curves of 3 different methods (sKPCR, KPCR and LSKM) with 2 different kernels (polynomial kernel and linear kernel) under different signal-to-noise ratios (0–0.25) and null connectivity proportion of 0.9.

### Surface-based fMRI with non-linear signal (polynomial relationship)

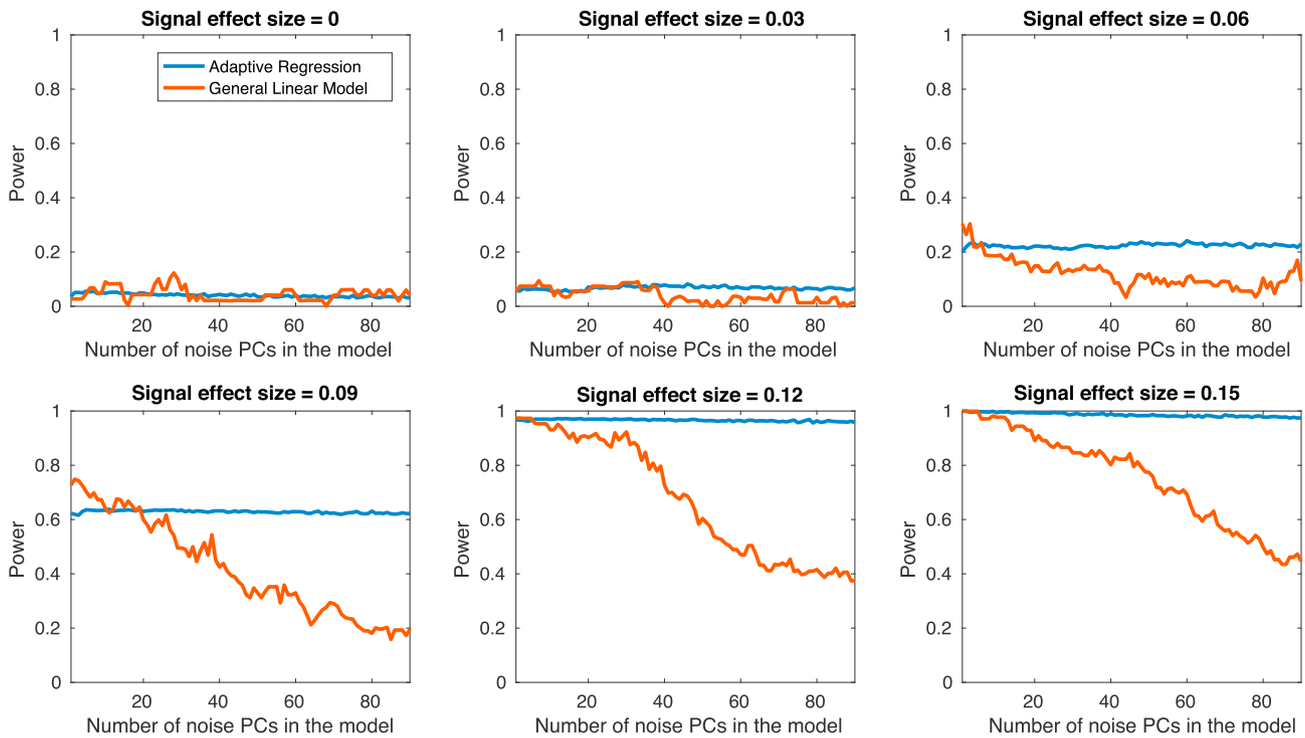


**Fig. 7.** Comparisons of the power of detecting *nonlinear* signals with different methods using simulations in *surface-based fMRI data*. In this simulation, polynomial (nonlinear) relationships were simulated to exist between some functional connectivities and a phenotype of interest, with polynomial degree ranging from 2 to 7 in each figure (e.g.  $y = x^d$ , where  $y$  is a phenotype of interest,  $x$  is a functional connectivity and  $d = 2, \dots, 7$ ). Each figure plots the power curves of 3 different methods (sKPCR, KPCR and LSKM) with 2 different kernels (a polynomial kernel and a linear kernel) under different signal-to-noise ratio (0–0.25) and a null connectivity proportion of 0.9.

#### 5.1.6. The stability of sKPCR under different covariance operators

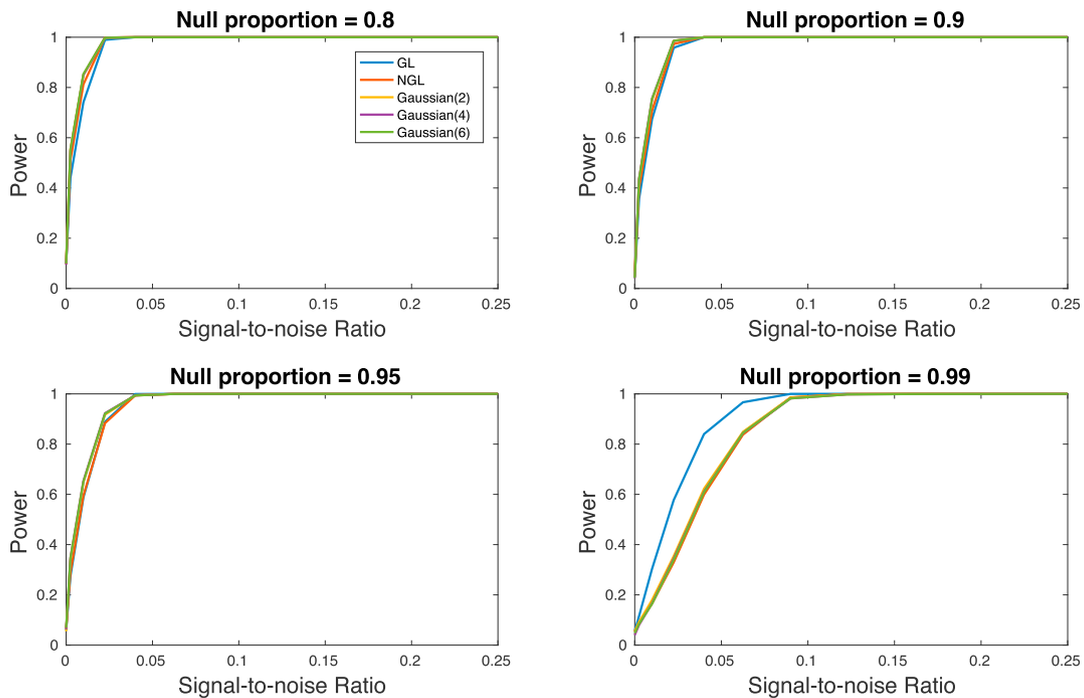
Fig. 9 shows that the power of sKPCR using different covariance operators under different signal-to-noise ratios and different proportions of null functional connectivities (see Section 3.7). We find that its power is

similar across three different covariance operators and operator parameters. The GL operator has a slightly higher power, so we used it throughout our simulations and real data analysis.



**Fig. 8.** A simulated example to demonstrate the power benefits of adaptive regression compared to the traditional general linear model  $F$  test. In the absence of a signal (no PCs correlate with phenotypes of interest, top left figure), both approaches can control the type one error rate accurately, i.e., power =  $\alpha \approx 5\%$ . When signals exist and the number of noise PCs increases, however, the power of the adaptive regression approach is stable, which means that it is sufficiently robust with respect to the selection of the number of PCs, while the power of the traditional general linear model decreases dramatically. When the number of noise PCs is small, it should be noted that adaptive regression has a somewhat lower power than the traditional general linear model.

### sKPCR power using different covariance operators



**Fig. 9.** The power of linear sKPCR with different covariance operators under different signal-to-noise ratio (0–0.25) and proportions of null functional connectivities (0.8–0.99). The covariance operators evaluated here are: Graph Laplacian (GL), Normalized Graph Laplacian (NGL), and Gaussian with  $\sigma^2 = 2, 4, 6$  (Gaussian (2), Gaussian (4), Gaussian (6)).

**Table 2**

Comparing the between-sites reproducibility of different approaches in 4 schizophrenia datasets using the Dice Coefficient (DC), with voxel-wise p-value threshold of 0.001 and permutation-based cluster-size FWER correction. In bold: the best performance among all methods.

Sites	sKPCR	MDMR	SPU(1)	SPU(2)	Univariate	aSPU
COBRE & Taiwan	<b>42%</b>	24%	9%	19%	13%	22%
COBRE & NMorphCH	12%	14%	3%	4%	<b>20%</b>	14%
COBRE & BrainGluSchi	<b>16%</b>	9%	2%	0.3%	0%	1%
Taiwan & NMorphCH	<b>10%</b>	8%	2%	2%	9%	7%
Taiwan & BrainGluSchi	<b>13%</b>	7%	2%	0.2%	1%	1%
NMorphCH & BrainGluSchi	<b>7%</b>	4%	0%	0%	0.7%	0.5%

**Table 3**

Comparing the schizophrenia findings with different approaches with existing meta-analysis results (Neurosynth ‘schizophrenia’ term) in 4 datasets. The Dice Coefficient (DC) is used to quantify the proportion of overlap. The schizophrenia findings are thresholded using a voxel-wise p-value of 0.001 and permutation-based cluster-size FWER correction. The Neurosynth ‘schizophrenia’ forward inference map is thresholded using FDR = 0.01 (the default setting in the database (<http://neurosynth.org/analyses/terms/schizophrenia/>)). In bold: the best performance among all methods.

Sites	sKPCR	MDMR	SPU(1)	SPU(2)	Univariate	aSPU
COBRE	<b>39%</b>	30%	10%	12%	5%	14%
Taiwan	<b>69%</b>	56%	5%	34%	26%	37%
NMorphCH	<b>8%</b>	7%	0.3%	1%	3%	3%
BrainGluSchi	<b>11%</b>	1%	2%	0.1%	0.4%	1%

## 5.2. Evaluating sKPCR in real data: brain-wide associations of the schizophrenic connectome

### 5.2.1. Between-sites reproducibility

Table 2 shows the results of comparing the between-sites reproducibility of different approaches (see Section 4.2). In five of the six comparisons, our sKPCR approach with a linear kernel achieved the highest between-site reproducibility values, as measured by DC. Only the univariate approach outperformed our method in one comparison. The voxel-wise p-value map of sKPCR in four datasets is shown in Figs. S1–S4, and the corresponding results with global signal regression are shown in Figs. S6–S9, which were very similar to the results without global signal regression.

### 5.2.2. Literature evidence: overlaps with existing meta-analysis findings

Table 3 shows the results of comparing the overlap of the findings with different methods with existing meta-analysis findings in the Neurosynth database (see Section 4.3). Our sKPCR method with a linear kernel has the highest overlap with existing findings in the literature. The schizophrenia meta-analysis map is shown in Supplementary Fig. S5.

### 5.2.3. Computation time

Finally, we compare the computation time of sKPCR, MDMR and aSPU in the above analyses. All the analyses were implemented in MATLAB 2016b using 20 cores on a Linux workstation with Intel Xeon E5-2660 v3 (2.60 GHz) CPU and 128 GB memory. Table 4 shows that our method is the most efficient.

## 6. Discussion

The sKPCR described in this paper is a powerful and efficient multivariate approach for voxel-level connectome-wide association studies. It can identify voxels, the overall connectivity pattern of which, as summarised by sKPCA as low-dimensional features, correlates with the

**Table 4**

Computation time of different methods when analyzing different real datasets with an optimized software implementation in MATLAB (number of permutations = 2000, number of cores = 20).

Datasets	COBRE	Taiwan	NMorphCH	BrainGluSchi
sKPCR	<b>0.5h</b>	<b>1.2h</b>	<b>0.3h</b>	<b>0.5h</b>
MDMR	1.0h	4.3h	0.4h	1.0h
aSPU	7.8h	9.4h	7.7h	7.8h

phenotypes of interest. The idea behind sKPCR simply involves reducing the dimensionality of the connectivity features and then performing association studies. However, we went further and carefully refined these two steps, aiming to extract more information from the fMRI data. Specifically, sKPCR models the spatial noise structure in the dimension reduction step and automatically selects an optimal number of principal components in the association testing steps. In our simulation, we demonstrated that sKPCR usually had the highest power in both volume-based and surface-based fMRI data for detecting both linear and nonlinear signals. In real data analysis, we showed that sKPCR usually had better between-sites reproducibility, larger overlap with existing findings, and faster computation speed.

A voxel/vertex identified by this approach can be interpreted as ‘there may exist one or more functional connectivities which connect it that are associated with the phenotype of interest’. To know the associated connections, a subsequent seed-based analysis can be performed. That is, we can extract a seed time series by averaging the voxel/vertex's time series within a significant cluster and test the associations between the seed connectivity map and a phenotype of interest. However, no significant individual connections in the seed-based analysis may be found because our approach can detect more than simple linear association signals. For example, consider a scenario in which many of the connections only have small effect sizes. In addition, as our approach can produce a voxel-wise statistical map, but not a connectivity-wise result, it can be directly compared with results of other analyses, such as task-activation studies, voxel-based morphometry (VBM) analysis, and Neurosynth meta-analysis results, even though our results do not reflect the direction of the association.

Additional areas can be refined. First, the method currently can only analyse binary and continuous phenotype variables. However, it could be extended to analyse categorical and multivariate phenotypes. Second, a sparse version of sKPCA, which allows only a subset of functional connectivities related to a principal component, may further improve the performance of dimension reduction, just like sparse PCA (Witten et al., 2009) improves PCA. Third, with the larger size of the available datasets, such as HCP and the UK-Biobank, an online version of sKPCA should be an important extension because it is not currently possible to fit thousands of high-resolution fMRI data into memory. However, sKPCR could be equipped to analyse big datasets by borrowing ideas from the online/group PCA approach (Smith et al., 2014) and other related variants (Monti and Hyvärinen, 2018; Chen et al., 2015a). Fourth, it would also be very interesting to extend sKPCR to infer causal relationships (Tran and Blei, 2017). Finally, the current sKPCR method is designed for single-site studies. However, combining the sKPCR results from multiple imaging sites is an important extension for the future. Possible methods include conventional meta-analysis methods and the model-based site-effect adjustment methods, such as ComBat (Johnson et al., 2007).

## Acknowledgements

Weikang Gong would like to thank Professor Stephen M. Smith for valuable discussions and suggestions. Jianfeng Feng is supported by the National High Technology Research and Development Program of China (No. 2015AA020507), the Key Program of National Natural Science Foundation of China (No. 91230201), the International (Regional) Collaborative and Exchange Program of National Natural Science (No.

71661167002), the Key Project of Shanghai Science and Technology Innovation Plan (No. 15JC1400101), the Shanghai Soft Science Research Program (No. 15692106604), and the National Centre for Mathematics and Interdisciplinary Sciences (NCMIS) of the Chinese Academy of Sciences.

## Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.neuroimage.2018.12.032>.

## References

- Allen, Genevera I., Grosebeck, Logan, Taylor, Jonathan, 2014. A generalized least-square matrix decomposition. *J. Am. Stat. Assoc.* 109 (505), 145–159.
- Bach, Francis R., Jordan, Michael I., 2005. *A Probabilistic Interpretation of Canonical Correlation Analysis*.
- Bai, Jushan, Ng, Serena, 2002. Determining the number of factors in approximate factor models. *Econometrica* 70 (1), 191–221.
- Bai, Jushan, Ng, Serena, 2007. Determining the number of primitive shocks in factor models. *J. Bus. Econ. Stat.* 25 (1), 52–60.
- Bartholomew, David J., Knott, Martin, 1999. *Latent Variable Models and Factor Analysis*, vol. 7. Arnold London.
- Beckmann, Christian F., Smith, Stephen M., 2004. Probabilistic independent component analysis for functional magnetic resonance imaging. *IEEE Trans. Med. Imag.* 23 (2), 137–152.
- Belilovsky, Eugene, Varoquaux, Gaël, Blaschko, Matthew B., 2016. Testing for differences in Gaussian graphical models: applications to brain connectivity. In: *Advances in Neural Information Processing Systems*, pp. 595–603.
- Bellec, Pierre, Benhajali, Yassine, Carbonell, Felix, Dansereau, Christian, Albouy, Geneviève, Pelland, Maxime, Craddock, Cameron, Collignon, Olivier, Julien, Doyon, Stip, Emmanuel, et al., 2015. Impact of the resolution of brain parcels on connectome-wide association studies in fmri. *Neuroimage* 123, 212–228.
- Benjamini, Yoav, Hochberg, Yosef, 1995. Controlling the false discovery rate: a practical and powerful approach to multiple testing. *J. Roy. Stat. Soc. B* 289–300.
- Bishop, Christopher M., 2006. Pattern recognition and machine learning. *J. Electron. Imag.* 16, 049901.
- Brown, Michael P.S., Grundy, William Noble, Lin, David, Cristianini, Nello, Walsh Sugnet, Charles, Furey, Terrence S., Ares, Manuel, Haussler, David, 2000. Knowledge-based analysis of microarray gene expression data by using support vector machines. *Proc. Natl. Acad. Sci. Unit. States Am.* 97 (1), 262–267.
- Chen, Po-Hsuan Cameron, Chen, Janice, Yeshurun, Yaara, Hasson, Uri, James, Haxby, Ramadge, Peter J., 2015a. A reduced-dimension fmri shared response model. In: *Advances in Neural Information Processing Systems*, pp. 460–468.
- Chen, Shuo, Kang, Jian, Xing, Yishi, Wang, Guoqing, 2015b. A parsimonious statistical method to detect groupwise differentially expressed functional connectivity networks. *Hum. Brain Mapp.* 36 (12), 5196–5206.
- Cheng, Wei, Rolls, Edmund T., Gu, Huaiguang, Zhang, Jie, Feng, Jianfeng, 2015. Autism: reduced connectivity between cortical areas involved in face expression, theory of mind, and the sense of self. *Brain* 138 (5), 1382–1393 page awv051.
- Cheng, Wei, Rolls, Edmund T., Jiang, Qiu, Liu, Wei, Tang, Yanqing, Huang, Chu-Chung, Wang, Xinfa, Zhang, Jie, Lin, Wei, Zheng, Lirong, et al., 2016. Medial reward and lateral non-reward orbitofrontal cortex circuits change in opposite directions in depression. *Brain* 139 (12), 3296–3309 page awv255.
- Choi, Yunjin, Taylor, Jonathan, Tibshirani, Robert, 2014. Selecting the Number of Principal Components: Estimation of the True Rank of a Noisy Matrix. 1410.8260.
- Eklund, Anders, Nichols, Thomas E., Knutsson, Hans, 2016. Cluster failure: why fmri inferences for spatial extent have inflated false-positive rates. *Proc. Natl. Acad. Sci. Unit. States Am.* 201602413.
- Escoufier, Y., 1977. Operators Related to a Data Matrix. *Recent Developments in Statistics*, pp. 125–131.
- Fiecas, Mark, Cribben, Ivor, Bahktari, Reyhaneh, Cummine, Jacqueline, 2017. A variance components model for statistical inference on functional connectivity networks. *Neuroimage* 149, 256–266.
- Finn, Emily S., Shen, Xilin, Scheinost, Dustin, Rosenberg, Monica D., Huang, Jessica, Chun, Marvin M., Papademetris, Xenophon, Constable, R Todd, 2015. Functional connectome fingerprinting: identifying individuals using patterns of brain connectivity. *Nat. Neurosci.* 18 (11), 1664–1671.
- Flandin, Guillaume, Penny, William D., 2007. Bayesian fmri data analysis with sparse spatial basis function priors. *Neuroimage* 34 (3), 1108–1125.
- Friston, Karl J., Worsley, Keith J., Frackowiak, R.S.J., Mazziotta, John C., Evans, Alan C., 1994. Assessing the significance of focal activations using their spatial extent. *Hum. Brain Mapp.* 1 (3), 210–220.
- Ge, Tian, Feng, Jianfeng, Hibar, Derrek P., Thompson, Paul M., Nichols, Thomas E., 2012. Increasing power for voxel-wise genome-wide association studies: the random field theory, least square kernel machines and fast permutation procedures. *Neuroimage* 63 (2), 858–873.
- Glasser, Matthew F., Sotiropoulos, Stamatios N., Wilson, J Anthony, Coalson, Timothy S., Fischl, Bruce, Andersson, Jesper L., Xu, Junqian, Jbabdi, Saad, Webster, Matthew, Polimeni, Jonathan R., et al., 2013. The minimal preprocessing pipelines for the human connectome project. *Neuroimage* 80, 105–124.
- Gong, Qiyong, He, Yong, 2015. Depression, neuroimaging and connectomics: a selective overview. *Biol. Psychiatry* 77 (3), 223–235.
- Gong, Weikang, Wan, Lin, Lu, Wenlian, Liang, Ma, Fan, Cheng, Cheng, Wei, Gruenewald, Stefan, Feng, Jianfeng, 2018. Statistical testing and power analysis for brain-wide association study. *Med. Image Anal.* 47, 15–30.
- Greve, Douglas N., Fischl, Bruce, 2009. Accurate and robust brain image alignment using boundary-based registration. *Neuroimage* 48 (1), 63–72.
- Johnson, W Evan, Cheng, Li, Rabinovic, Ariel, 2007. Adjusting batch effects in microarray expression data using empirical bayes methods. *Biostatistics* 8 (1), 118–127.
- Jolliffe, Ian, 2002. *Principal Component Analysis*. Wiley Online Library.
- Kaczurkin, A.N., Moore, T.M., Calkins, M.E., Ciric, R., Detre, J.A., Elliott, M.A., Foa, E.B., Garcia de la Garza, A., Roalf, D.R., Rosen, A., et al., 2017. Common and dissociable regional cerebral blood flow differences associate with dimensions of psychopathology across categorical diagnoses. *Mol. Psychiatr.* 23 (10), 1981–1989.
- Kim, Junghi, Wozniak, Jeffrey R., Mueller, Bryon A., Shen, Xiaotong, Pan, Wei, 2014. Comparison of statistical tests for group differences in brain functional networks. *Neuroimage* 101, 681–694.
- Kim, Junghi, Pan, Wei, Alzheimer's Disease Neuroimaging Initiative, et al., 2015. Highly adaptive tests for group differences in brain functional connectivity. *Neuroimage: Clinical* 9, 625–639.
- Lam, Clifford, Yao, Qiwei, 2012. Factor modeling for high-dimensional time series: inference for the number of factors. *Ann. Stat.* 694–726.
- Lee, Seunggeun, Wu, Michael C., Lin, Xihong, 2012. Optimal tests for rare variant effects in sequencing association studies. *Biostatistics* 13 (4), 762–775.
- Li, Zeng, Wang, Qinwen, Yao, Jianfeng, et al., 2017. Identifying the number of factors from singular values of a large sample auto-covariance matrix. *Ann. Stat.* 45 (1), 257–288.
- Liu, Dawei, Lin, Xihong, Ghosh, Debashis, 2007. Semiparametric regression of multidimensional genetic pathway data: least-squares kernel machines and linear mixed models. *Biometrics* 63 (4), 1079–1088.
- Meskaldji, Djalel Eddine, Fischl-Gomez, Elda, Griffa, Alessandra, Hagmann, Patric, Morgenthaler, Stephan, Thiran, Jean-Philippe, 2013. Comparing connectomes across subjects and populations at different scales. *Neuroimage* 80, 416–425.
- Meskaldji, Djalel-Eddine, Vasung, Lana, Romascano, David, Thiran, Jean-Philippe, Hagmann, Patric, Morgenthaler, Stephan, Van De Ville, Dimitri, 2015. Improved statistical evaluation of group differences in connectomes by screening-filtering strategy with application to study maturation of brain connections between childhood and adolescence. *Neuroimage* 108, 251–264.
- Monti, Ricardo Pio, Hyvärinen, Aapo, 2018. A Unified Probabilistic Model for Learning Latent Factors and Their Connectivities from High-dimensional Data. 1805.09567.
- Nichols, Thomas E., Holmes, Andrew P., 2002. Nonparametric permutation tests for functional neuroimaging: a primer with examples. *Hum. Brain Mapp.* 15 (1), 1–25.
- Pan, Wei, Kim, Junghi, Zhang, Yiwei, Shen, Xiaotong, Peng, Wei, 2014. A powerful and adaptive association test for rare variants. *Genetics* 197 (4), 1081–1095.
- Patel, Ameera X., Kundu, Prantik, Rubinov, Mikail, Jones, P Simon, Vértes, Petra E., Ersche, Karen D., John Suckling, Bullmore, Edward T., 2014. A wavelet method for modeling and despiking motion artifacts from resting-state fmri time series. *Neuroimage* 95, 287–304.
- Penny, William D., Trujillo-Barreto, Nelson J., Friston, Karl J., 2005. Bayesian fmri time series analysis with spatial priors. *Neuroimage* 24 (2), 350–362.
- Ramsay, James O., 2006. *Functional Data Analysis*. Wiley Online Library.
- Robinson, Emma C., Jbabdi, Saad, Glasser, Matthew F., Andersson, Jesper, Burgess, Gregory C., Harms, Michael P., Smith, Stephen M., Van Essen, David C., Jenkinson, Mark, 2014. Msm: a new flexible framework for multimodal surface matching. *Neuroimage* 100, 414–426.
- Rolls, Edmund T., Cheng, Wei, Gong, Weikang, Jiang, Qiu, Zhou, Chanjuan, Zhang, Jie, Lv, Wujun, Ruan, Hongtao, Wei, Dongtao, Cheng, Ke, Meng, Jie, Xie, Peng, Feng, Jianfeng, 2018. Functional connectivity of the anterior cingulate cortex in depression and in health. *Cerebr. Cortex*. <https://doi.org/10.1093/cercor/bhy236> page bhy236.
- Romme, Ingrid AC., de Reus, Marcel A., Ophoff, Roel A., Kahn, René S., van den Heuvel, Martijn P., 2017. Connectome discontinuity and cortical gene expression in patients with schizophrenia. *Biol. Psychiatry* 81 (6), 495–502.
- Salimi-Khorshidi, Gholamreza, Douaud, Gwenaelle, Beckmann, Christian F., Glasser, Matthew F., Griffanti, Ludovica, Smith, Stephen M., 2014. Automatic denoising of functional mri data: combining independent component analysis and hierarchical fusion of classifiers. *Neuroimage* 90, 449–468.
- Satterthwaite, Theodore D., Vandekar, Simon N., Wolf, Daniel H., Danielle, S Bassett, Ruparel, Kosha, Shehzad, Zarrar, Cameron Craddock, R., Shinohara, Russell T., Moore, Tyler M., Gennatas, Efsthathios D., et al., 2015. Connectome-wide network analysis of youth with psychosis-spectrum symptoms. *Mol. Psychiatr.* 20 (12), 1508–1515.
- Schölkopf, Bernhard, Alexander, Smola, Klaus-Robert Müller, 1997. Kernel principal component analysis. In: *International Conference on Artificial Neural Networks*. Springer, pp. 583–588.
- Shehzad, Zarrar, Kelly, Clare, Reiss, Philip T., Craddock, R Cameron, Emerson, John W., McMahon, Katie, Copland, David A., Xavier Castellanos, F., Milham, Michael P., 2014. A multivariate distance-based analytic framework for connectome-wide association studies. *Neuroimage* 93, 74–94.
- Sidén, Per, Anders, Eklund, Bolin, David, Villani, Mattias, 2017. Fast bayesian whole-brain fmri analysis with spatial 3d priors. *Neuroimage* 146, 211–225.
- Simpson, Sean L., Laurienti, Paul J., 2015. A two-part mixed-effects modeling framework for analyzing whole-brain network data. *Neuroimage* 113, 310–319.
- Smith, Stephen M., Nichols, Thomas E., 2009. Threshold-free cluster enhancement: addressing problems of smoothing, threshold dependence and localisation in cluster inference. *Neuroimage* 44 (1), 83–98.
- Smith, Stephen M., Beckmann, Christian F., Andersson, Jesper, Auerbach, Edward J., Bijsterbosch, Janine, Douaud, Gwenaelle, Duff, Eugene, Feinberg, David A.,

- Griffanti, Ludovica, Harms, Michael P., et al., 2013. Resting-state fmri in the human connectome project. *Neuroimage* 80, 144–168.
- Smith, Stephen M., Hyvärinen, Aapo, Varoquaux, Gaël, Miller, Karla L., Beckmann, Christian F., 2014. Group-pca for very large fmri datasets. *Neuroimage* 101, 738–749.
- Smith, Stephen M., Nichols, Thomas E., Vidaurre, Diego, Winkler, Anderson M., Behrens, Timothy E.J., Glasser, Matthew F., Ugurbil, Kamil, Barch, Deanna M., Van Essen, David C., Miller, Karla L., 2015. A positive-negative mode of population covariation links brain connectivity, demographics and behavior. *Nat. Neurosci.* 18 (11), 1565–1567.
- Sur, Pragma, Chen, Yuxin, Candès, Emmanuel J., 2017. The Likelihood Ratio Test in High-dimensional Logistic Regression Is Asymptotically a Rescaled Chi-square. 1706.01191.
- Tipping, Michael E., Bishop, Christopher M., 1999. Probabilistic principal component analysis. *J. Roy. Stat. Soc. B* 61 (3), 611–622.
- Tran, Dustin, Blei, David M., 2017. Implicit Causal Models for Genome-wide Association Studies. 1710.10742.
- Witten, Daniela M., Tibshirani, Robert, Hastie, Trevor, 2009. A penalized matrix decomposition, with applications to sparse principal components and canonical correlation analysis. *Biostatistics* 10 (3), 515–534.
- Woo, Choong-Wan, Krishnan, Anjali, Tor, D Wager, 2014. Cluster-extent based thresholding in fmri analyses: pitfalls and recommendations. *Neuroimage* 91, 412–419.
- Worsley, Keith J., Marrett, Sean, Peter, Neelin, Vandal, Alain C., Friston, Karl J., Evans, Alan C., et al., 1996. A unified statistical approach for determining significant signals in images of cerebral activation. *Hum. Brain Mapp.* 4 (1), 58–73.
- Xia, Mingrui, He, Yong, 2017. Functional Connectomics from a Big Data Perspective. *NeuroImage*.
- Yarkoni, Tal, Poldrack, Russell A., Nichols, Thomas E., Van Essen, David C., Wager, Tor D., 2011. Large-scale automated synthesis of human functional neuroimaging data. *Nat. Methods* 8 (8), 665–670.
- Zalesky, Andrew, Fornito, Alex, Bullmore, Edward T., 2010. Network-based statistic: identifying differences in brain networks. *Neuroimage* 53 (4), 1197–1207.
- Zhang, Hui, Nichols, Thomas E., Johnson, Timothy D., 2009. Cluster mass inference via random field theory. *Neuroimage* 44 (1), 51–61.
- Zhu, Hongtu, Shen, Dan, Peng, Xuwei, Liu, Leo Yufeng, 2017. Mwpccr: multiscale weighted principal component regression for high-dimensional prediction. *J. Am. Stat. Assoc.* 112 (519), 1009–1021.