

# NIH Public Access

**Author Manuscript** 

Med Image Comput Comput Assist Interv. Author manuscript; available in PMC 2014 August

#### Published in final edited form as:

Med Image Comput Comput Assist Interv. 2013; 16(01): 300-307.

# Persistent Homological Sparse Network Approach to Detecting White Matter Abnormality in Maltreated Children: MRI and DTI Multimodal Study

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# Abstract

We present a novel persistent homological sparse network analysis framework for characterizing white matter abnormalities in tensor-based morphometry (TBM) in magnetic resonance imaging (MRI). Traditionally TBM is used in quantifying tissue volume change in each voxel in a massive univariate fashion. However, this obvious approach cannot be used in testing, for instance, if the change in one voxel is related to other voxels. To address this limitation of univariate-TBM, we propose a new persistent homological approach to testing more complex relational hypotheses across brain regions. The proposed methods are applied to characterize abnormal white matter in maltreated children. The results are further validated using fractional anisotropy (FA) values in diffusion tensor imaging (DTI).

# **1** Introduction

Traditionally tensor-based morphometry (TBM) in magnetic resonance imaging (MRI) has been massively univariate in that response variables are fitted using a linear model at each voxel producing massive number of test statistics (Figure 1). However, univariate approaches are ill-suited for testing more complex hypotheses about multiple anatomical regions. For example, the univariate-TBM cannot answer how the volume increase in one voxel is related to other voxels. To address this type of more complex relational hypothesis across different brain regions, we propose a new persistent homological approach.

The Jacobian determinant is the most often used volumetric mesurement in TBM. We propose to correlate the Jacobian determinant across different voxels and quantify how the volume change in one voxel is correlated to the volume changes in other voxels. However, existing multivariate statistical methods exhibit serious defects in applying to the whole brain regions due to the *small-n large-p problem* [2]. Specifically, the number of voxels *p* are substantially larger than the number of subjects *n* so the often used maximum likelihood estimation (MLE) of the covariance matrix shows the rank deficiency and it is no longer positive definite. In turn, the estimated correlation matrix is not considered as a good approximation to the true correlation matrix. The small-*n* large-*p* problem can be addressed

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by regularizing the ill-conditioned correlation or covariance matrices by sparse regularization terms.

Sparse model A is usually parameterized by a tuning parameter  $\lambda$  that controls the sparsity of the representation. Increasing the sparse parameter makes the representation more sparse. Instead of performing statistical inference at one fixed  $\lambda$  that may not be optimal, we propose to quantify how the topology of sparse solution changes over the increasing  $\lambda$  using the persistent homology. Then it is possible to obtain additional characterization of a population that cannot be obtained in the univariate-TBM.

The proposed framework is applied in characterizing abnormal white matter alterations in children who experienced maltreatment while living in post-institutional (PI) settings before being adopted by families in the US. The main contributions of the paper are (i) the introduction of a novel persistent homological approach to characterizing white matter abnormality and (ii) its application to MRI and DTI showing consistent results between the modalities.

# 2 Motivation

Let  $\mathbf{J}_{n \times p} = (J_{ij})$  be the matrix of Jacobian determinant for subject *i* at voxel position *j*. The subscripts denote the dimension of matrix. There are *p* voxels of interest and *n* subjects. The Jacobian determinants of all subjects at the *j*-th voxel is denoted as  $\mathbf{x}_j = (J_{1j}, \dots, J_{nj})'$ . The Jacobian determinants of all voxels for the *i*-th subject is denoted as  $\mathbf{y}_i = (J_{i1}, \dots, J_{ip})'$ .  $\mathbf{x}_j$  is the *j*-th column and  $\mathbf{y}_i$  is the *i*-th row of the data matrix  $\mathbf{J}$ . The covariance matrix of  $\mathbf{y}_i$  is given by Cov  $(\mathbf{y}_i) = \sum_{p \times p} = (\sigma_{kl})$  and estimated using the sample covariance matrix *S via* MLE. To remedy this small–*n* and large-*p* problem, the likelihood is regularized with  $L_1$ -penalty:

$$L(\sum^{-1}) = \text{logdet}\sum^{-1} - \text{tr}\left(\sum^{-1}S\right) - \lambda \|\sum^{-1}\|_{1^{1}}$$
(1)

where  $\|\cdot\|_1$  is the sum of the absolute values of the elements. *L* is maximized over all possible symmetric positive definite matrices. (1) is a convex problem and it is usually solved using the graphical-lasso (GLASSO) algorithm [4,6].

Since the different choice of parameter  $\lambda$  will produce different solutions, we propose to use the collection of  $\Sigma^{-1}(\lambda)$  for every possible value of  $\lambda$  for the subsequent statistical inference. This avoids the problem of identifying the optimal sparse parameter that may not be optimal in practice. The question is then how to use the collection of  $\Sigma^{-1}(\lambda)$  in a coherent mathematical fashion.

Consider a sparse model  $\mathcal{A}(\lambda)$ , which gets more sparse as  $\lambda$  increases. Then under some condition, it is possible to have  $\mathcal{A}(\lambda_1) \supset \mathcal{A}(\lambda_2) \supset \mathcal{A}(\lambda_3) \supset \cdots$  for  $\lambda_1 \leq \lambda_2 \leq \cdots$ . Within the persistent homological framework [5],  $\mathcal{A}(\lambda)$  is said to be *persistent* if it has this type of nested subset structure. The collection of the nested subsets is called *filtration*.

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### 3 Persistent Structures for Sparse Network Models

#### Sparse Correlations

We assume the measurement vector  $\mathbf{x}_j$  at the *j*-th node is centered with zero mean and unit variance. These condition is achieved by centering and normalizing data. Let  $\mathbf{\Gamma} = (\gamma_{jk})$  be the correlation matrix, where  $\gamma_{jk}$  is the correlation between the nodes *j* and *k*. Sparse correlation  $\mathbf{\Gamma}$  is then estimated as

$$\hat{\boldsymbol{\Gamma}} = \arg\min_{\beta} \frac{1}{2} \sum_{j=1}^{p} \sum_{k\neq j} \|\mathbf{x}_j - \beta_{jk} \mathbf{x}_k\|_2^2 + \lambda \|\beta\|_1, \quad (2)$$

where  $\beta = (\beta_{jk})$ . When  $\lambda = 0$ , the sparse correlation is simply given by the sample correlation, i.e.  $\hat{\gamma}_{jk} = \mathbf{x}'_{j} \mathbf{x}_{k}$ . As  $\lambda$  increases, the correlation becomes more sparse. Using the sparse solution (2), we will explicitly construct a persistent structure on  $\Gamma(\lambda)$  over changing  $\lambda$ .

Let  $A = (a_{ik})$  be the adjacency matrix defined using the sparse correlation:

$$a_{jk}(\lambda) = \begin{cases} 1 & \text{if } \hat{\gamma}_{jk} \neq 0; \\ 0 & \text{otherwise.} \end{cases}$$

Let  $\mathcal{G}(\lambda)$  be the graph induced from the adjacency matrix *A*. It can be algebraically shown that the induced graph is persistent and from a filtration:

$$\mathscr{G}(\lambda_1) \supset \mathscr{G}(\lambda_2) \supset \mathscr{G}(\lambda_3) \supset \cdots$$
 (3)

for  $\lambda_1 \leq \lambda_2 \leq \lambda_3$ . The proof follows by simplifying the adjacency matrix A into a simpler but equivalent adjacency matrix  $B = (b_{ik})$ :

$$b_{jk}(\lambda) = \begin{cases} 1 & \text{if } |\mathbf{x}_j'\mathbf{x}_k| > \lambda; \\ 0 & \text{otherwise.} \end{cases}$$
(4)

Then it is not difficult to see the graph induced from the adjacency matrix *B* should be persistent. Hence, *the filtration on*  $\mathcal{G}(\lambda)$  *can be constructed by simply thresholding the* 

sample correlation  $\mathbf{x}'_{j}\mathbf{x}_{k}$  for each  $\lambda$  without solving the optimization problem (2). Figure 2 shows filtrations obtained from sparse correlations between Jacobian determinants on preselected 548 nodes in the two groups showing group difference. It is not necessary to perform filtrations for infinitely many possible filtration values. For an *n*-node network, it can be algebraically shown that at most n - 1 increments are sufficient to obtain a unique filtration.

#### Sparse Likelihood

The identification of a persistent homological structure out of the inverse covariance  $\Sigma^{-1}(\lambda)$ in (1) is similar. However, it is more involved than the sparse correlation case. Let  $A = (a_{ij})$  be the adjacency matrix

$$a_{ij}(\lambda) = \begin{cases} 1 & \text{if } \hat{\sigma}^{ij} \neq 0; \\ 0 & \text{otherwise.} \end{cases}$$
(5)

The adjacency matrix A induces a graph  $\mathcal{G}(\lambda)$  consisting of  $\kappa(\lambda)$  number of partitioned subgraphs

$$\mathscr{G}(\lambda) {=} \underset{l=1}{\overset{\kappa(\lambda)}{\cup}} G_l(\lambda) \quad \text{with} \quad G_l {=} \{V_l(\lambda), E_l(\lambda)\},$$

where  $V_l$  and  $E_l$  are vertex and edge sets of the subgraph  $G_l$  respectively. Unlike the sparse correlation case, we do not have full persistency on the induced graph  $\mathcal{G}$ . The partitioned graphs can be proven to be partially nested in a sense that only the partitioned node sets are persistent [4,6], i.e.

$$V_l(\lambda_1) \supset V_l(\lambda_2) \supset V_l(\lambda_3) \supset \cdots$$
 (6)

for  $\lambda_1 \leq \lambda_2 \leq \lambda_3$  and all *l*. Subsequently the collection of partitioned vertex set

 $\mathscr{V}(\lambda) = \bigcup_{l=1}^{\kappa(\lambda)} V_l(\lambda)$  is also persistent. On the other hand, the edge sets  $E_l$  may not be persistent. The identification of the vertex filtration can be fairly time consuming since it requires solving the convex optimization problem (1) for multiple  $\lambda$  values. However, it can be easily obtained by identifying a simpler adjacency matrix *B* that gives the identical vertex sets  $V_l$  (Figure 3).

Let  $B(\lambda) = (b_{ij})$  be another adjacency matrix given by

$$b_{ij}(\lambda) = \begin{cases} 1 & \text{if } |\hat{s}_{ij}| > \lambda; \\ 0 & \text{otherwise.} \end{cases}$$
(7)

where  $\hat{s}_{ij}$  is the sample covariance matrix. The adjacency matrix *B* similarly induces the graph  $\mathcal{H}$  with  $\tau(\lambda)$  disjoint subgraphs:

$$\mathscr{H}(\lambda) = \bigcup_{l=1}^{\tau(\lambda)} H_l(\lambda)$$

with  $H_l = \{W_l(\lambda), F_l(\lambda)\}$ .  $W_l$  and  $F_l$  are vertex and edge sets of the subgraph  $H_l$  respectively. Then trivially the node set  $W_l$  forms a filtration over the sparse parameter:

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$$W_l(\lambda_1) \supset W_l(\lambda_2) \supset W_l(\lambda_3) \supset \dots$$
 (8)

It can be further shown that  $\kappa(\lambda) = \tau(\lambda)$  and  $V_l(\lambda) = W_l(\lambda)$  for all  $\lambda > 0$  [6]. Hence, the filtration on the vertex set  $V_l(\lambda)$  is constructed by simply thresholding the sample covariance  $\hat{s}_{ij}(\lambda)$  for each  $\lambda$  without solving the time consuming optimization problem (2). Figure 3 shows the schematic of constructing the filtration on sparse likelihood by the covariance thresholding.

# 4 Application to Maltreated Children Study

#### MRI Data and Univariate-TBM

T1-weighted MRI were collected using a 3T GE SIGNA scanner for 23 children who experienced maltreatment while living in post-institutional (PI) settings before being adopted by families in the US, and age-matched 31 normal control subjects. The average age for PI is  $11.26 \pm 1.71$  years while that of controls is  $11.58 \pm 1.61$  years. A study specific template was constructed using the diffeomorphic shape and intensity averaging technique through Advanced Normalization Tools (ANTS) [1]. Image normalization of each individual image to the template was done using symmetric normalization with cross-correlation as the similarity metric. The 1mm deformation fields are then smoothed out with Guassian kernel with bandwidth  $\sigma = 4$ mm.

The computed Jacobian maps were feed into univariate-GLM at each voxel for testing the group effect while accounting for age and gender difference. Figure 1 shows the significant group difference between PI and controls. Any region above the *T*-statistic value of 4.86 or below –4.86 is considered significant at 0.05 (corrected). However, what the univariate-TBM can not determine is the dependency of Jacobian determinants at two different positions. It is possible that structural abnormality at one region of the brain might be related to the other regions due to interregional dependency. For this type of more complex hypothesis, we need the proposed persistent homological approach.

#### Inference on Barcodes

Since Jacobian determinants at neighboring voxels are highly correlated, we uniformly subsampled p = 548 number of nodes along the white matter template mesh vertices in order not to have spurious high correlation between two adjacent nodes (Figure 1). The proposed method is very robust under the change of node sizes. For the node sizes between 548 and 1856, the choice of node sizes did not affect the subsequent analysis. Following the proposed method, we constructed the filtrations on sparse correlations and inverse covariance without solving the optimizations (1) and (2). The filtrations are quantified using the barcode representation, which plots the change of Betti numbers over filtration values [5] (Figure 4). The first Betti number  $\beta_0(\lambda)$  counts the number of connected components of the graph  $\mathcal{G}(\lambda)$  at the filtration value  $\lambda$ .

Given the barcode  $\beta_0^i(\lambda)$  for group *i*, we tested if the barcodes were different between the groups, i.e.  $\beta_0^1(\lambda) \neq \beta_0^2(\lambda)$  for some  $\lambda \in [0, 1]$ . A Kolmogorov-Smirnov (KS) like test

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statistic  $T = \sup_{\lambda \in [0,1]} |\beta_0^1(\lambda) - \beta_0^2(\lambda)|$  is used. Since each group produces one barcode, we used the Jackknife resampling technique for inference. For a group with *n* subjects, one subject is removed and the remaining *n* – 1 subjects are used in constructing the barcode. This process is repeated for each subject to produce *n* barcodes (Figure 4). The Jackknife resampling produces 23 and 31 barcodes respectively for PI and controls. In order for the permutation test to converge for our data set, it requires tens of thousands permutations and it is really time consuming. So we used a much simpler Jackknife resampling. Then the test statistic *T* is constructed between 23 × 31 pairs of barcodes. Under the null, *T* is expected to be zero. One-sample T-test on the collection of *T* is then subsequently performed to show huge group discrimination for sparse correlations in Figure 4 (*p*-value < 0.001). The barcodes for normal controls show much higher Betti numbers at the given threshold. This suggests higher non-uniformity in Jacobian determinants across the brain that causes increased disconnections in correlations. The inverse covariance was not able to discriminate the groups. The MATLAB codes for constructing barcodes.

#### Validation Against DTI

For children who suffered early neglect and abuse, white matter microstructures are more diffusely organized [3]. So we expect less white matter variability not only in the Jacobian determinants but also in the fractional anisotropy (FA) values in DTI as well. The MRI data in this study has the corresponding DTI. The DTI acquisition are done in the same 3T GE SIGNA scanner and the acquisition parameters can be found in [3]. We applied the proposed persistent homological method in obtaining the filtrations for sparse correlations and inverse covariances in the same 548 nodes (Figure 1). The resulting filtration patterns also show similar pattern of rapid increase in disconnected components (Figure 2 and 4) for sparse correlations. The Jackknife-based one-sample T-test also shows significant group difference for correlations (p-value < 0.001). This results are due to consistent abnormality observed in both MRI and DTI modalities. PI exhibited stronger white matter homogeneity and less spatial variability compared to normal controls in both MRI and DTI measurements. The inverse covariance was not able to discriminate the groups.

# **5** Conclusions and Discussions

Using the persistent homological framework, we have shown that PI group shows less anatomic variation in MRI compared to the normal controls. This result is consistent with DTI, which shows similar patterns. The reason we did not detect the group difference in the inverse covariances might be that, as shown in Figure 4, the changes in the first Betti number are occurring in a really narrow window and losing the discrimination power. On the other hand the sparse correlations exhibit more slow changes in the Betti number over the wide window making it easier to discriminate the groups. The proposed method is general enough to run on any type of volumetric imaging data that is spatially normalized.

# Acknowledgments

This work was supported by NIH Research Grants MH61285 and MH68858 to S.D.P., NIMH Grant MH84051 to R.J.D. and US National Institute of Drug Abuse (Fellowship DA028087) to J.L.H. The authors like to thank

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Matthew Arnold of University of Bristol for the discussion on the KS-test procedure and pointing out the reference [6].

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# Fig. 1.

(a) *T*-statistic map of group differences (PI-controls) on Jacobian determinants. (b) 548 uniformly sampled nodes in MRI where the persistent homology is applied. The nodes are sparsely sampled in the template to guarantee there is no spurious high correlation due to proximity between nodes. (c) The same nodes are taken in DTI to check the consistency against the MRI results.

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#### Fig. 2.

Networks  $\mathcal{G}(\lambda)$  obtained by thresholding sparse correlations for the Jacobian determinant from MRI and fractional anisotropy from DTI at different  $\lambda$  values. The collection of the thresholded graphs forms a filtration. PI shows more dense network at a given  $\lambda$  value. Since PI is more homogenous, in the white matter region, there are more dense high correlations between nodes. The filtration is visualized using the equivalent dendrogram [5], which also shows more dense linkages for PI at high correlations. Chung et al.



# Fig. 3.

Schematic of graph filtrations obtained by sparse-likelihood (5) and sample covariance thresholding (7). The vertex set of  $\mathcal{G}(\lambda_1) = \mathcal{H}(\lambda_1)$  consists of gray nodes. For the next filtration value  $\lambda_2$ ,  $\mathcal{G}(\lambda_1) \neq \mathcal{H}(\lambda_1)$ . However, the partitioned vertex sets (yellow and red) of  $\mathcal{G}(\lambda_1)$  and  $\mathcal{H}(\lambda_1)$  match.

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# Fig. 4.

The barcodes on the sparse inverse covariance (top) and correlation (bottom) for Jacobian determinant (left) and FA (right). Unlike the inverse sparse covariance, the sparse correlation shows huge group separation between normal controls and post-institutionalized (PI) children (p-value < 0.001).