Harmonic holes as the submodules of brain network and network dissimilarity

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Abstract. Persistent homology has been applied to brain network analysis for finding the shape of brain networks across multiple thresholds. In the persistent homology, the shape of networks is often quantified by the sequence of k-dimensional holes and Betti numbers. The Betti numbers are more widely used than holes themselves in topological brain network analysis. However, the holes show the local connectivity of networks, and they can be very informative features in analysis. In this study, we propose a new method of measuring network differences based on the dissimilarity measure of harmonic holes (HHs). The HHs, which represent the substructure of brain networks, are extracted by the Hodge Laplacian of brain networks. We also find the most contributed HHs to the network difference based on the HH dissimilarity. We applied our proposed method to clustering the networks of 4 groups, normal control (NC), stable and progressive mild cognitive impairment (sMCI and pMCI), and Alzheimer's disease (AD). The results showed that the clustering performance of the proposed method was better than that of network distances based on only the global change of topology.

Keywords: Topological data analysis · Brain network · Alzheimer's disease · Harmonic holes · Hodge Laplacian

1 Introduction

Persistent homology has been widely applied to brain network analysis for finding the topology of networks in multiscale [5,16,23,25] Since a 'simplicial complex' is not a familiar term in brain network analysis, we refer to it as a 'network' that is generally used. It quantifies the shape of brain networks by using k-dimensional holes and their cardinality, the kth Betti number [2,10]. A persistence diagram (PD) summarizes the change of Betti numbers during the filtration of networks by recording when and how holes appear and disappear during the filtration. The persistent homology also provides distances for distinguishing networks such as the bottleneck distance and kernel-based distances [10,20]. Such distances mostly find network differences in their PDs. The Betti numbers and PDs are more often used than holes themselves in network applications.

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Holes represent the submodule of brain networks. 0-dimensional holes, i.e., connected components, modules or clusters have been widely studied for finding functional or structural submodules in a brain [6,16,27]. On the other hand, 1-dimensional holes have been rarely used for brain network analysis [7,17,15,19,24,26] Most studies in brain network analysis do not use 2- and higher order simplexes in networks since networks. Therefore, all cycles in a network are considered as 1-dimensional holes. There are few network measures based on cycles in brain network analysis such as cycle probability and the change of the number of cycles during graph filtration [7,26]. These measures helped to compare the global property of networks but could not find the discriminative substructures of networks.

If higher order simplexes are introduced in a network, the number of 1dimensional holes is significantly reduced due to the removal of filled-in triangles. The previous brain network studies that studied higher order simplexes mostly found holes based on Zomorodian and Carlsson's (ZC) algorithm [19,24,28]. The ZC algorithm is very fast in linear-time, however, it finds the sparse representation of a hole that identifies only one path around the hole and ignores the other paths. This introduces an ambiguity in hole identification in practice. A better approach would be to localize the holes by the eigen-decomposition of Hodge Laplacian of a network. Such holes are called as the harmonic holes (HHs). The HH shows all possible paths around the hole with their weights [12,13,18]. The HHs have been applied to brain network analysis for localizing persistent holes [15,17]. The 1-dimensional holes in a network with higher order simplexes have at least one indirect path between every two nodes. Thus, the holes are related to the abnormality or inefficiency of the network. The previous studies found the persistent holes with long duration in a network as abnormal holes, and localized them by harmonic holes. Therefore, the duration of holes was used instead of HHs in network discrimination.

In this paper, we propose a new measure for estimating network dissimilarity based on persistent HHs (HH dissimilarity). The proposed HH dissimilarity is motivated from the bottleneck distance. The bottleneck distance first estimates the correspondence between holes between networks that are represented by points in PDs, and then chooses the maximum among all the distances between the estimated pairs of holes [8]. The HH dissimilarity also estimates the correspondence between HHs of two different networks that are represented by real-valued eigenvectors, and takes the averaged dissimilarities of the estimated pairs of HHs. The advantage of HH dissimilarity is not only to measure the network differences but also to quantify a HH's contributions to the network differences. We will call the amount of HH's of contribution the *citation* of HH. This allows us to identify the discriminative subnetworks of networks.

The proposed method is applied to metabolic brain networks obtained from the FDG PET dataset in Alzheimer's disease neuroimaging initiative (ADNI). The dataset consists of 4 groups: normal controls (NC), stable and progressive mild cognitive impairment (sMCI and pMCI), and Alzheimer's disease (AD). We generated 2400 networks by bootstrap, and compared the clustering performance with the existing network distances such as L_2 -norm (L2) of the difference between distance matrices, Gromov-Hausdorff (GH) distance, Kolmogorov-Smirnov (KS) distance of connected components and cycles (KS₀ and KS₁), and bottleneck distance of holes [3,6,7,8,16]. The results showed that the HH dissimilarity had the superior clustering performance than the other distance measures, and comparing local connectivities could be more helpful to discriminating the progression of Alzheimer's disease.

2 Materials and methods

2.1 Data sets, preprocessing, and the construction of metabolic connectivity

We used FDG PET images in ADNI data set (http://adni.loni.usc.edu). The ADNI FDG-PET dataset consists of 4 groups: 181 NC, 91 sMCI, 77 pMCI, and 135 AD (Age: 73.7 ± 5.9 , range $56.1 \sim 90.1$). FDG PET images were measured 30 to 60 minutes and they were averaged over all frames. The voxel size in the images were standardized in $1.5 \times 1.5 \times 1.5$ mm resolution. The images were spatially normalized to Montreal Neurological Institute (MNI) space using statistical parametric mapping (SPM8, www.fil.ion.ucl.ac.uk/spm). The details of data sets and preprocessing are given in [4]. The whole brain image was parcellated into 94 regions of interest (ROIs) based on automated anatomical labeling (AAL2) excluding cerebellum [21]. The 94 ROIs served as network nodes and their measurements were obtained by averaging FDG uptakes in the ROI. The averaged FDG uptake was globally normalized by the sum of 94 averaged FDG uptakes. The distance between 2 nodes was estimated by the diffusion distance on positive correlation between the measurements. The diffusion distance considers an average distance of all direct and indirect paths between 2 nodes via random walks [9]. The diffusion distance is known to be more robust to noise and outliers.

2.2 Harmonic holes

Simplicial complex The algebraic topology extends the concept of a graph further to a simplicial complex. Suppose that a non-empty node set V is given. If the set of all subsets of V is denoted by 2^V , an abstract simplicial complex Kis a subset of 2^V such that (1) $\emptyset \in K$, and (2) if $\sigma \in K$ and $\tau \in \sigma$, $\tau \in K$ [10,11]. Each $\sigma \in K$ is called a simplex. A *i*-dimensional simplex is an element with i + 1 nodes, $v_1, \ldots, v_{i+1} \in V$, denoted by $\sigma_i = [v_1, \ldots, v_{i+1}]$. The dimension of K, denoted as dimK, is the maximum dimension of a simplex $\sigma \in K$. The collection of σ_i 's in K is denoted by K_i ($-1 \leq i \leq \dim K$). The number of simplices in K_i is denoted as $|K_i|$. The *i*-skeleton of K is defined as $K^{(i)} = K_0 \cup \cdots \cup K_i$ ($0 \leq i \leq \dim K$). Thus, a graph with nodes and edges is 1-skeleton $K^{(1)}$. In this paper, we will only consider 2-skeleton $K^{(2)}$ of a simplicial complex that includes nodes, edges, and triangles. For convenience, we call it *a* (simplicial) *network* [14]. 4 H. Lee et al.

Incidence matrix We denote a $|K_i|$ -dimensional integer space as $\mathbb{Z}^{|K_i|}$. Given a finite simplicial complex K, a chain complex C_i is defined in $\mathbb{Z}^{|K_i|}$ [10,28]. The boundary operator ∂_i and coboundary operator ∂_i^{\top} for $i = 1, \ldots, N$ (N > 0) are functions such that $\partial_i : C_i \to C_{i-1}$ and $\partial_i^{\top} : C_{i-1} \to C_i$, respectively. We define $\partial_i = 0$ for i < 1 or i > N.

Given $\sigma_i = [v_1, ..., v_{i+1}] \in C_i$, the boundary of σ_i is algebraically defined as

$$\partial_i \sigma_i = \sum_{j=1}^{i+1} (-1)^{j-1} [v_1, \dots, v_{j-1}, v_{j+1}, \dots, v_{i+1}].$$

If the sign of σ_{i-1} in $\partial_i \sigma_i$ is positive/negative, it is called positively/negatively oriented with respect to σ_i . We denote the positive/negative orientation by $\sigma_{i-1} \in_{+/-} \sigma_i$. The boundary of the boundary is always zero, i.e., $\partial_{i-1}\partial_i = 0$.

If the simplicial complex K has

$$K_{i} = \left\{ \sigma_{i}^{1}, \cdots, \sigma_{i}^{|K_{i}|} \right\}, \quad K_{i-1} = \left\{ \sigma_{i-1}^{1}, \cdots, \sigma_{i-1}^{|K_{i-1}|} \right\}$$

the boundary operator ∂_i is represented by the *i*th incidence matrix $M_i \in \mathbb{Z}^{|K_{i-1}| \times |K_i|}$ such that [12,13,18]

$$[\boldsymbol{M}_{i}]_{mn} = \begin{cases} 1 \text{ if } \sigma_{i-1}^{m} \in_{+} \sigma_{j}^{n}, \\ -1 \text{ if } \sigma_{i-1}^{m} \in_{-} \sigma_{j}^{n}, \\ 0 \text{ otherwise.} \end{cases}$$
(1)

The coboundary operator ∂_i^{\top} is represented by \boldsymbol{M}_i^{\top} . σ_i^n in K_i is represented by a vector in $\mathbb{Z}^{|K_i|}$ in which the *n*th entry is 1 and the rest is 0. The linear combination of σ_i 's can be represented by the linear combination of $|K_i|$ -dimensional vectors.

Combinatorial Hodge Laplacian A combinatorial Hodge Laplacian L_i : $C_i \rightarrow C_i$ is defined by

$$\boldsymbol{L}_{i} = \boldsymbol{L}_{i}^{up} + \boldsymbol{L}_{i}^{down} = \boldsymbol{M}_{i+1} \boldsymbol{M}_{i+1}^{\top} + \boldsymbol{M}_{i}^{\top} \boldsymbol{M}_{i}, \qquad (2)$$

where $\boldsymbol{L}_{i}^{up} \in \mathbb{Z}^{|K_{i}| \times |K_{i}|}$ and $\boldsymbol{L}_{i}^{down} \in \mathbb{Z}^{|K_{i}| \times |K_{i}|}$ are composite functions $\partial_{i+1}\partial_{i+1}^{\top}$: $C_{i} \to C_{i+1} \to C_{i}$ and $\partial_{i}^{\top}\partial_{i}: C_{i} \to C_{i-1} \to C_{i}$, respectively [12,13,14,18] The kernel and image of \boldsymbol{L}_{i} are denoted by ker \boldsymbol{L}_{i} and img \boldsymbol{L}_{i} , respectively. The ker \boldsymbol{L}_{i} is called harmonic classes H_{i} [14].

The *i*th homology and cohomology groups of $C = \{C_i, \partial_i\}$ are defined respectively by

$$\tilde{H}_i(C) = \ker \partial_i / \operatorname{img} \partial_{i+1}$$
 and $\tilde{H}^i(C) = \ker \partial_{i+1}^\top / \operatorname{img} \partial_i^\top$.

Theorem 1 (Combinatorial Hodge Theory [12,14,18]). Suppose that a chain complex $\{C_i(X; \mathbb{R}), \partial_i\}$ is given for i = 0, ..., N, and $C_i(X; \mathbb{R})$ is considered as an \mathbb{R} -vector space. Harmonic classes H_i obtained by the combinatorial Laplacian L_i are congruent to the *i*th homology and cohomology groups, \tilde{H}_i and \tilde{H}^i of C, *i.e.*,

$$H_i \cong H_i(C; \mathbb{R}) \cong H^i(C; \mathbb{R}).$$

Proof. rank H_i = rank C_i -rank L_i = rank C_i -(rank ∂_i +rank ∂_{i+1}) = rank $\tilde{H}_i(C; \mathbb{R})$.

The harmonic classes $H_i = \ker L_k$ is also called a harmonic space [14]. The homology group \tilde{H}_i in persistent homology can be replaced with a harmonic space H_i , and the rank of H_i is the same as the *i*th Betti number. We call a hole in H_i a harmonic hole (HH), and a hole in \tilde{H}_i estimated by Smith normal form a binary hole [28].

Given a simplicial network with p nodes, q edges, and r filled-in triangles, we estimate $L_1 \in \mathbb{Z}^{q \times q}$ in (2), and $H_i = \{ \boldsymbol{x} \in \mathbb{R}^{q \times 1} | \boldsymbol{L}_1 \boldsymbol{x} = 0 \}$. The eigenvector of L_1 with zero eigenvalue, $\boldsymbol{x} \in \mathbb{R}^{q \times 1}$ represents a HH. The entry of \boldsymbol{x} can be positive or negative depending on the orientation of edges in the hole. The absolute value of the entry of \boldsymbol{x} represents the weight of the corresponding edge in the hole. Since \boldsymbol{x} and $-\boldsymbol{x}$ have zero eigenvalue, they represent the same hole, and $\parallel \boldsymbol{x} \parallel = 1$.

Computing persistent HHs In this study, we have the distances between pairs of nodes in a brain network. Given a set of nodes and their distances, Rips complex with threshold ϵ is the clique complex induced by a set of edges with their distances less than ϵ . Rips filtration is the nested sequence of Rips complexes obtained by increasing threshold ϵ . To compute persistent holes over threshold, we perform Rips filtration on brain network nodes [2,10].

Zomorodian and Carlsson developed an efficient algorithm for computing persistent holes based on the Smith normal form [28]. It is an incremental algorithm that updates the range and null spaces of incidence matrices during Rips filtration. The representation of a persistent binary hole is changed by adding simplexes during Rips filtration. The ZC algorithm chose the youngest binary hole at the birth of the persistent hole. The ZC algorithm is fast in practically linear-time, however, the obtained binary hole shows only one path around the hole and the other paths are ignored. On the other hand, a HH shows all possible paths around the persistent hole, and represents the contribution of a path to the generation of the hole by edge weights in the path. Thus, the HH is better in localizing a persistent hole than a binary hole when we want to extract local connectivity in a brain network. However, there is no algorithm for estimating persistent HHs during the filtration in literature.

In this study, we will estimate the youngest persistent HHs just like the ZC algorithm. First, we sort edges e_1, \ldots, e_q in the ascending order of an edge distance, and perform the Rips filtration by the fast ZC algorithm. To avoid having the same edge distance, we select the ordered index $1, \ldots, q$ as the filtration value, instead of the edge distance. The reason for performing the ZC algorithm first is that the computation of eigen-decomposition at every filtration value is too expensive. Then, we obtain a PD which is the set of the birth and death thresholds of persistent holes. If a persistent hole appears at i_X and disappears at i_Z , we perform the eigen-decomposition of Hodge Laplacian at i_X, i_Z , and $i_Y = i_Z - 1$ to estimate the corresponding HH. The i_Y is the threshold just before the death of the persistent hole.

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The harmonic spaces at i_X, i_Y , and i_Z are written by matrices

$$H_X = [\boldsymbol{x}_1, \cdots, \boldsymbol{x}_l] \in \mathbb{R}^{q \times l}, H_Y = [\boldsymbol{y}_1, \cdots, \boldsymbol{y}_m] \in \mathbb{R}^{q \times m}, H_Z = [\boldsymbol{z}_1, \cdots, \boldsymbol{z}_n] \in \mathbb{R}^{q \times n}$$

respectively. The HH appearing at i_X and disappearing at i_Z will be in H_X and H_Y , but not in H_Z . We find which $\boldsymbol{y} \in H_Y$ does not depend on \boldsymbol{z}_i 's in H_Z . If $\boldsymbol{y} \in H_Y$ depends on H_Z , the smallest singular value of the matrix $[H_Z, \boldsymbol{y}]$ is close to 0. It implies that \boldsymbol{y} still exists in H_Z . Therefore, we choose $\boldsymbol{y} \in H_Y$ such that

$$\boldsymbol{y} = \underset{\boldsymbol{y} \in H_Y}{\operatorname{arg\,max}} \{ \text{the smallest singular value of } [H_Z, \boldsymbol{y}] \}.$$
(3)

The chosen \boldsymbol{y} by (3) is the oldest persistent HH. Next, we choose the youngest persistent HH $\boldsymbol{x} \in H_X$ such that

$$\boldsymbol{x} = \operatorname*{arg\,min}_{x \in H_X} \left\{ \text{the smallest singular value of } [\boldsymbol{x}, \boldsymbol{y}] \right\} = \operatorname*{arg\,min}_{x \in H_X} \left\{ 1 - |\boldsymbol{x}^\top \boldsymbol{y}| \right\}. (4)$$

This procedure is repeated for all persistent holes. The incidence matrices are already estimated during the ZC algorithm. Since the incidence matrices and their combinatorial Hodge Laplacian are very sparse, the computation of persistent HHs is not so hard in our experiments. In our experiments, the total number of persistent holes during the filtration is not more than 50, and the number of persistent holes at each filtration value is not more than 20.

2.3 HH dissimilarity

Bottleneck distance If K_a and K_b have m and n persistent holes. The PDs of K_a and K_b are denoted respectively by $PD_a = \{t_1^a, \dots, t_m^a\}$ and $PD_b = \{t_1^b, \dots, t_n^b\}$, where t_i is a point with the birth and death thresholds of the corresponding hole. Bottleneck distance between two simplicial complexes, K_a and K_b is defined by [8]

$$D_B(K_a, K_b) = d(PD_a, PD_b) = \inf_{\eta: PD_a \to PD_b} \sup_{t \in PD_a} \| t - \eta(t) \|_{\infty},$$

where η is a bijection from PD_a to PD_b and $||(x, y)||_{\infty} = \max\{|x|, |y|\}$ is the L_{∞} -norm. If there is no corresponding hole in the other PD because of $m \neq n$, the points on the diagonal line x = y that have the shortest distance from the point t are included. In this way, the bottleneck distance measures network distance by the difference of the birth and death thresholds of holes, not by the difference between holes themselves.

Dissimilarity between HHs If the eigenvectors with zero eigenvalues of two different combinatorial Laplacians are denoted by x and y, their dissimilarity is defined by one minus the absolute value of their inner product, i.e.,

$$d_h(\boldsymbol{x}, \boldsymbol{y}) = 1 - |\boldsymbol{x}^\top \boldsymbol{y}|.$$
(5)

This is the smallest singular value of the matrix [x, y] in (4) that shows the dependency between x and y. If x and y are similar, their dissimilarity is close to 0; otherwise, it is close to 1.

HH dissimilarity Suppose that two networks K_a and K_b have m and n persistent HHs, denoted by $\boldsymbol{H}_a = [\boldsymbol{x}_1^a, \cdots, \boldsymbol{x}_m^a]$ and $\boldsymbol{H}_b = [\boldsymbol{x}_1^b, \cdots, \boldsymbol{x}_n^b]$, respectively. The dissimilarity based on persistent HHs (HH dissimilarity) is defined by

$$D_H(K_a, K_b) = d(\boldsymbol{H}_a, \boldsymbol{H}_b) = \inf_{\zeta: H_a \to H_b} \frac{1}{\min(m, n)} \sum_{\boldsymbol{x} \in H_a} d_h(\boldsymbol{x}, \zeta(\boldsymbol{x})), \quad (6)$$

where ζ is a bijection from H_a to H_b .

The correspondence ζ between persistent HHs in two different networks is determined by minimizing the total distances between the pairs of HHs based on Munkres assignment algorithm, also known as Hungarian algorithm. Some of persistent HHs can not find their corresponding HHs in the other network because of $m \neq n$. In this study, we ignore them and average the dissimilarities of the obtained pairs of persistent HHs.

Citation of HH The advantage of using HH dissimilarity is the ability to quantify how much a persistent HH contributes in differentiating networks. The degree of the contribution of HH is called the citation of HH. If a persistent HH \boldsymbol{x} in \boldsymbol{H}_a corresponds to a persistent HH $\boldsymbol{y} = \zeta(\boldsymbol{x})$ in \boldsymbol{H}_b in (6), their dissimilarity is $d_h(\boldsymbol{x}, \boldsymbol{y}) = 1 - |\boldsymbol{x}^\top \boldsymbol{y}|$, and their similarity is defined by $|\boldsymbol{x}^\top \boldsymbol{y}|$. If the persistent HHs of l networks are denoted by $\mathcal{H} = \{\boldsymbol{H}_1, \cdots, \boldsymbol{H}_l\}$ and they are compared with \boldsymbol{H}_a , the citation of \boldsymbol{x} is defined by

$$\sum_{\zeta(\boldsymbol{x})\in H, \forall H\in\mathcal{H}} |\boldsymbol{x}^{\top}\zeta(\boldsymbol{x})|.$$

If we find the most cited HHs by comparing networks within a group, we can determine which submodule makes two networks in a group close to each other. Furthermore, if we find the most cited HHs by comparing network between groups, we can determine which submodule makes differences.

3 Results

3.1 Brain network construction

We had 4 groups, NC, sMCI, pMCI, and AD which had 181, 91, 77 and 135 subjects, respectively. The subjects in a group could be heterogeneous. Thus, we obtained 600 bootstrap samples from each group by randomly selecting the subset of the number of subjects in each group with replacement [22]. The number of bootstrap samples was heuristically determined in comparison with previous study [22]. We constructed 600 bootstrapped networks from bootstrap samples in each group by diffusion distance in Sec. 2.1. The total number of generated brain networks was 2400.





Fig. 1. Distance of 2400 networks. (a) L2, (b) GH, (c) KS_0 , (d) KS_1 , (e) Bottleneck, and (f) HH. The 2400 networks were sorted in the order of NC, sMCI, pMCI, and AD. Each group had 600 networks. The clustering accuracy is shown in Table 1.

3.2 Network clustering

We clustered 2400 bootstrapped brain networks into 4 groups by Ward's hierarchical clustering method. The Ward's hierarchical clustering method found the group labels based on the distance between data points, which is a network in our application. The network distance was estimated by (a) L2, (b) GH distance, (c) KS₀, (d) KS₁, (e) bottleneck distance of holes, and (f) HH dissimilarity [3,6,7,8,16]. The obtained distance matrices of 2400 networks were shown in Fig. 1. After clustering networks, we matched the estimated group label with the true group label of networks and calculated the clustering accuracy of 8 distance matrices. The clustering accuracy of 8 distance matrices was shown in Table 1. We also clustered 1200 bootstrapped networks in sMCI and pMCI into 2 groups by the same way. The clustering accuracy was shown in Table 1.

3.3 The most cited HHs

We selected the 600 most cited HHs within NC, sMCI, pMCI, and AD, and divided them into 5 clusters based on the dissimilarity between HHs in (5). In Fig. 3 (a-d), because the dissimilarity of HHs in the cluster 5 was large, we considered HHs in the cluster 5 as outliers. We calculated the center of HHs in clusters 1, 2, 3, and 4, by selecting the HH with the minimum sum of dissimilarities with the other HHs in the cluster. The 4 representative HHs of 4 clusters were shown on the left of Fig. 3 (a-d). In each panel, the upper row showed the HHs in a brain, and the lower row showed the HHs in a 2-dimensional plane. The location

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of nodes in the 2-dimensional plane was estimated by Kamada-Kawai algorithm implemented in a network analysis/visualization toolbox, Pajek [1]. In Fig. 3 (a-d), the width of an edge was proportional to the edge weight in the HH. The larger the weight of an edge, the darker the color of an edge. The color of nodes represented the location of nodes in a brain. If a node was located in frontal, parietal, temporal, occipital, subcortical, and limbic regions, the color of the node was red, blue, green, purple, yellow, and orange, respectively.

We also selected the 600 most cited HHs when we compared networks between sMCI and pMCI, and divided them into 5 clusters. In Fig. 2 (a), the cluster 5 contained the outliers. Thus, we estimated the center HHs in cluster 1-4. The representative HHs in sMCI and the corresponding holes in pMCI were shown in Fig. 2 (b).

4 Discussion and conclusions

In this study, we proposed a new network dissimilarity, called HH dissimilarity. Unlike a binary hole estimated by the ZC algorithm, a HH show all possible paths of edges around a hole, and the contribution of paths to forming the hole is represented by the weight of edges on the paths. If an edge belongs to a unique path that forms a hole, its edge weight will be large. If an edge belongs to one of many alternative paths as in a module, its edge weight will be small. In this way, HHs can extract the substructures of a brain network including holes and modules. Moreover, since the HHs can be represented as real-valued orthonormal vectors we can define the dissimilarity between HHs as well as HH dissimilarity between brain networks easily using vector product.

Brain networks of different groups may share common substructure as well as have different substructures that make individual and group differences. The proposed HH dissimilarity first finds candidates of common substructures between brain networks and estimates the over all dissimilarities between candidates. The clustering results showed that brain networks of different groups had similar substructures, however, the averaged similarities was much larger than that of brain networks within a group.

The goal of persistent homology may be to find persistent features that last for a long duration. However, in brain network analysis, it has been applied for

Distance	4 groups	2 groups
	(NC, sMCI, pMCI, and AD)	(sMCI and pMCI)
L2	66.09~%	98.50~%
GH	45.96~%	87.58~%
KS_0	52.54~%	74.00~%
KS_1	77.38~%	79.83~%
Bottleneck	45.71 %	76.58~%
HH	$100 \ \%$	$100 \ \%$
	Distance L2 GH KS ₀ KS ₁ Bottleneck HH	$\begin{array}{c c} & 4 \mbox{ groups} \\ \hline {\rm Distance} & ({\rm NC, \ sMCI, \ pMCI, \ and \ AD}) \\ \hline {\rm L2} & 66.09 \ \% \\ {\rm GH} & 45.96 \ \% \\ {\rm KS}_0 & 52.54 \ \% \\ {\rm KS}_1 & 77.38 \ \% \\ {\rm Bottleneck} & 45.71 \ \% \\ {\rm HH} & 100 \ \% \end{array}$

 Table 1. Clustering accuracy



Fig. 2. (a) Clustering of the 600 most cited HHs when sMCI and pMCI were compared. (b) Representative HHs in cluster 1, 2, 3 and 4. The left two columns showed HHs in sMCI and the right two columns showed the corresponding HHs in pMCI. Each HH was visualized in a brain and in a 2-dimensional plane. The shape of the HH was more clearly shown in the plane, and the location of the HH could be checked in the brain. The color of nodes was determined by the location of nodes in a brain: frontal (red), parietal (blue), temporal (green), occipital (purple), subcortical (yellow), and limbic (orange) regions. If the edge weight was larger in a HH, the color of edge was darker and the width of edge was larger.

finding the change of topology, especially the change of connected components, instead of the persistence of topology. This study suggested a more coherent framework to observe, capture, and quantify the change of holes in brain networks. Depending on imaging modality and study populations, brain networks may have different characteristics of shapes. Therefore, it is necessary to apply proper network measures to brain networks depending on modality and population. The results showed that when the Alzheimer's disease progresses, the hole structure was changed in metabolic brain networks, and HHs and HH dissimilarity could predict the disease progression.

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Fig. 3. Distance matrix of the 600 most cited HHs within (a) NC, (b) sMCI, (c) pMCI, and (d) AD. The most cited holes were clustered into 5 groups. The last cluster 5 had outliers with large dissimilarities between HHs. The representative HHs of the first 4 clusters were plotted on the right. The upper row showed the HHs in a brain and the lower row showed the HHs in a 2-dimensional plane.