Finding cosmic voids and filament loops using topological data analysis

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Abstract

We present a method called Significant Cosmic Holes in Universe (SCHU) for identifying cosmic voids and loops of filaments in cosmological datasets and assigning their statistical significance using techniques from topological data analysis. In particular, persistent homology is used to find different dimensional holes. For dark matter halo catalogs and galaxy surveys, the 0-, 1-, and 2-dimensional holes can be identified with connected components (i.e. clusters), loops of filaments, and voids. The procedure overlays dark matter halos/galaxies on a three-dimensional grid, and a distance-to-measure (DTM) function is calculated at each point of the grid. A nested set of simplicial complexes (a filtration) is generated over the lower-level sets of the DTM across increasing threshold values. The filtered simplicial complex can then be used to summarize the birth and death times of the different dimension homology group generators (i.e., the holes). Persistent homology summary diagrams, called persistence diagrams, are produced from the dimension, birth times, and death times of each homology group generator. Using the persistence diagrams and bootstrap sampling, we explain how \(p\)-values can be assigned to each homology group generator. The homology group generators on a persistence diagram are not, in general, uniquely located back in the original dataset volume so we propose a method for finding a representation of the homology group generators. This method provides a novel, statistically rigorous approach for locating informative generators in cosmological datasets, which may be useful for providing complementary cosmological constraints on the effects of, for example, the sum of the neutrino masses. The method is tested on a Voronoi foam simulation, and then subsequently applied to a subset of the SDSS galaxy survey and a cosmological simulation. Lastly, we calculate Betti functions for two of the MassiveNuS simulations and discuss implications for using the persistent homology of the density field to help break degeneracy in the cosmological parameters.

Keywords: cosmology: large-scale structure of universe, cosmology: cosmological parameters, methods: numerical, methods: statistical, methods: N-body simulations, methods: data analysis

1. Introduction

The large-scale distribution of matter in the Universe forms a connected network known as the cosmic web (Klypin and Shandarin 1993; Bond et al. 1996). Anisotropic gravitational collapse of matter has resulted in a picture where galaxy clusters form the nodes of this web and are interconnected by filaments, which form at the intersections of walls. The remaining majority of space is filled by cosmic voids: vast underdense regions that have experienced minimal non-linear growth of structure.

In the current era of multiband, high-resolution large-scale structure (LSS) surveys, there has been prolific investigation of the large dark matter halos at the nodes of the cosmic web, including cosmological analysis via the abundance of galaxy clusters (e.g., Vikhlinin et al. 2009; Mantz et al. 2015; Planck Collaboration et al. 2016a), and analysis of the large-scale matter distribution in dark matter halos via cosmic shear studies (e.g., Joudaki et al. 2018; Troxel et al. 2018) and the clustering of galaxies (e.g., Tinker et al. 2012; Cacciato et al. 2013). However, there has been a growing tension between cosmological constraints derived using cosmic microwave background (CMB) and LSS measurements (Planck Collaboration et al. 2016a; Joudaki et al. 2018; Troxel et al. 2018).

Recently, voids have begun to attract interest as a cosmological probe complementary to halos. Unlike halos, which are regions that have experienced high levels of growth and virialization that can partially destroy signatures of the primordial density field, voids have only evolved minimally. Information about the geometry of the initial density field present in the voids has the potential to help break degeneracies in the cosmological parameters and tighten their current constraints (Hamaus et al. 2016). Additionally, motivated by the growing tension between CMB and LSS measurements, void statistics may be able to provide a complementary probe of the growth of structure that is less sensitive to non-linear structure formation physics (Lavaux and Wandelt 2010, 2012). Furthermore, methods have been proposed for utilizing voids to constrain dark energy (Pisani et al. 2015) and the sum of the neutrino masses...
Various observables can be used for constraining cosmology through voids, some of which include weak gravitational lensing \( \sum m_n \) (Kreisch et al., 2018, Massara et al., 2015), both of which have large effects on large-scale, low-density regions.

Current datasets that are available and have been used to characterize some of the aforementioned void observables include galaxy redshift surveys such as the Sloan Digital Sky Survey (SDSS; Abolfathi et al., 2018) and the Dark Energy Survey (DES; Abbott et al., 2016) as well as CMB anisotropy maps from Planck (Planck Collaboration et al., 2016). In the near future, next-generation galaxy surveys will go online, including the Large Synoptic Survey Telescope (LSST; Ivezić et al. 2008) and the Dark Energy Spectroscopic Instrument (DESI; DESI Collaboration et al., 2016), and the secondary CMB anisotropies will be observed to greater precision than ever before by the Simons Observatory (SO; The Simons Observatory Collaboration et al., 2018) and the CMB-S4 experiment (Abazajian et al., 2016). In anticipation of these large upcoming datasets, many theoretical and computational approaches for identifying voids from an input simulation or galaxy survey have been and are currently being developed in order to characterize the potential future constraining power of void clustering and abundance statistics Libskind et al. 2018; Sutter et al. 2012; Neyrinck 2008; Platen et al. 2007; Pranav et al. 2016 and references therein. The majority of these methods allow for the identification of the physical location of the voids in the matter field, enabling one to study clustering statistics such as the void two-point correlation function and abundance statistics such as the void mass and volume functions.

In this paper, we propose a method called Significant Cosmic Holes in Universe (SCHU) for relating the cosmic matter distribution to topology using persistent homology. Persistent homology quantifies and summarizes the shape of a dataset by recording locations of the representations of these structures back into the data volume. This is because the homology group generators identified are not uniquely mapped in such a way that a loop forms, surrounding an empty or low density 3D volumes (2nd-dimensional homology groups, \( H_2 \)). The SCHU uses the output of the persistent homology algorithm constructed using alpha shapes; in particular, they demonstrated that the Betti numbers across the filtration can be used to distinguish the matter distribution resulting from different dark energy models. Additionally, Pranav et al. (2016) introduced a multiscale topological measurement of the cosmic matter distribution and explored the analysis of Betti numbers and topological persistence of different cosmological models. A scale-free and parameter-free method for identifying the cosmic environments (voids, walls, filaments, nodes) called Discrete Persistent Structures Extractor (DisPerSE) was proposed in Sousbie (2011). DisPerSE computes the discrete Morse-Smale complex of a spatial dataset using the Delaunay tessellation field estimator (DTFE) technique (Schaap and van de Weygaert 2000, van de Weygaert and Schaap 2009). The mathematical background and algorithm implementation is described in Sousbie (2011) and applications to 3D simulation datasets and observed galaxy surveys are found in Sousbie et al. (2011).

As noted previously, persistent homology is a tool within topological data analysis (TDA) that finds different dimensional holes in data (e.g. connected components, loops, voids) and summarizes the generators by their lifetime in a particular filtration. These persistence diagrams and their associated Betti numbers can then be used for various types of statistical inference or as inputs into machine learning algorithms (Reininghaus et al., 2015). Persistent homology has proven to be useful in a variety of applications, such as natural language processing (Zhu 2013), computational biology (Xia and Wei 2014), Lyman-alpha forest studies (Cisewski et al., 2014), angiography (Bendich et al., 2016), and dynamical systems (Emrani et al., 2014).

Though useful for summarizing information for complex data, one shortcoming of persistent homology is that the homology group generators identified are not uniquely mapped back into the data volume. This is because the homology group generators displayed on the summary diagrams each represent an equivalence class of representations of that particular hole. SCHU uses the output of the persistent homology algorithm in order to find a representation of the equivalence class back in the original data volume. SCHU detects and captures the locations of cosmic voids (\( H_2 \) generators) along with another cosmic structure that we call filament loops (\( H_1 \) generators). Filament loops are formed when filaments are connected together in such a way that a loop forms, surrounding an empty or low density region, as shown in Figure 1. Thus, SCHU, and the persistent homology underlying SCHU, enable analysis of the cosmological density field: the void and filament loop locations and sizes enable the standard clustering and abundance statistics, and the persistence diagrams and Betti numbers provide additional topological summary statistics of the density field that can be used to further discriminate between cosmological models.

This article is organized as follows. In §2, we provide an overview of the formalism of persistent homology, describing filtrations, persistence diagrams, and bootstrap confidence bands. In §3 we present SCHU for identifying statistically significant voids and filament loops in astronomical datasets. In §4 we test
the void identification capabilities of SCHU on Voronoi foam simulation data, which is generated such that the ground truth void locations are known. In §3, we apply SCHU to identify voids and filament loops in a subset of the SDSS galaxy survey dataset. Additionally, we identify voids and filament loops in the cosmological N-body simulation from Libeskind et al. (2018) and compare the void locations to those found by other methods. We then study the Betti numbers of two simulations from the MassiveNuS simulation suite (Liu et al. 2018). Finally, in §6 we summarize our results and provide concluding remarks.

2. Introduction to persistent homology

2.1. Background

Homology describes different dimensional holes of a manifold. To be specific, the generators of $H_0$ describes connected components, the generators of $H_1$ describes closed loops, and the generators of $H_2$ describes voids (i.e. low-density or empty regions). Put into the context of cosmic web environments, the $H_0$ generators represent clusters of galaxies, the $H_1$ generators represent filaments that form loops, and the $H_2$ generators represent cosmic voids. Figure 2 illustrates an example of $H_0$ and $H_1$: Figure 2a shows a circle, which forms one closed loop (one $H_1$ generator) and is one connected component (one $H_0$ generator). Figure 2b shows 15 points randomly sampled on a circle, which is a dataset that has 15 connected components (15 $H_0$ generators) and no closed loop (zero $H_1$ generators). While there is not closed loop present, it is clear that the data were sampled on a circle; persistent homology provides a framework for identifying such a loop. Our present interest is in the identification and analysis of voids and filament loops in cosmological datasets, and thus we will focus our attention in this work towards the $H_1$ and $H_2$ generators.

Persistent homology is a framework for computing the homology of real data across a particular filtration, and results are summarized in a persistence diagram. A persistence diagram summarizes the birth and death times of the homology group generators (i.e., the time when a homology group generator first appears in the filtration, and the time when it is no longer present due to merging with another homology group generator or the closing of the hole). For a detailed introduction of persistent homology, we refer the reader to Edelsbrunner and Harer (2010). Several software and packages are available to compute persistent homology (e.g., Maria et al. (2014); Morozov (2007)). In this work, we use Dionysus (Morozov 2007) via the R package TDA (Fasy et al. 2014) along with several of the other functions available in the R package.

2.2. Filtration construction

Given a point cloud of data, $S_n = \{X_1, X_2, \ldots, X_n\}$ in $\mathbb{R}^d$, sampled from some true distribution $P$, one may be interested in the homology of $P$. Persistent homology computes the homology of $S_n$ across different levels of a constructed filtration (Edelsbrunner and Harer 2010). We consider a function-based construction that relies on a robust version of a distance function.

A distance function, $d_{L,n}$, is a function that maps points from $\mathbb{R}^d$ to $\mathbb{R}$, defined as $d_{L,n}(x) = \inf_{y \in S_n} \|x - y\|_L$, where $\| \cdot \|_L$ is the $L^2$ norm. A filtration can then be constructed using the lower level sets of $d_{L,n}$, $L_{n,t} = \{x : d_{L,n}(x) \leq t\}$, where $t$ is the filtration parameter defining the threshold of the lower-level sets. Notice that $L_{n,0} = S_n$ and $L_{n,\infty} = \mathbb{R}^d$. Persistent homology describes how the homology of $L_{n,t}$ changes as $t$ increases from 0 to $\infty$. Let $p$ be the homology group dimension, and then $H_p(L_{n,t})$ is the $p$th-dimensional homology group of $L_{n,t}$. In practice, we use a grid over the point cloud data and evaluate $d_{L,n}$ for each point on the grid. Consider a sequence of $t$: $t_0, t_1, t_2, \ldots, t_k, t_{k+1}$. Let $t_0$ be 0 and $t_{k+1}$ be $\infty$, and then $L_{n,t_0} = S_n$ and $L_{n,t_{k+1}} = \mathbb{R}^d$. (In practice, this just goes to the size of the dataset box.) We can obtain a filtration of homology groups:

$$H_p(L_{n,t_0}) \rightarrow H_p(L_{n,t_1}) \rightarrow H_p(L_{n,t_2}) \rightarrow \cdots \rightarrow H_p(n, L_{n,t_{k+1}}). \quad (1)$$

This filtration of homology groups is constructed by computing the homology of the lower-level sets at each threshold value $t$. Along the filtration, topological structures gradually change: homology groups appear at some point and disappear later during the filtration. The threshold value $t_k$ where $H_p(L_{n,t_k})$ contains the first appearance of a homology group along the filtration is called its birth time and the threshold value $t_l$ where $H_p(L_{n,t_l})$ contains the last appearance of the homology group is called its death time.

A more robust version of the distance function is available called the distance-to-measure (DTM) function (Chazal et al. 2011), and is useful for TDA (Chazal et al. 2017). Given a probability measure $P$, a DTM function with respect to a set $X$
is defined for each \( y \in \mathbb{R}^d \) as

\[
d_{m_0}(y) = \left( \frac{1}{m_0} \int_0^{m_0} (G_t^{-1}(u)')^r \, du \right)^{1/r},
\]

where \( G_t(u) = P(\|X - y\| \leq u) \), and \( m_0 \in (0, 1) \) and \( r \in [1, \infty) \) are tuning parameters. The \( m_0 \) can be seen as a smoothing parameter and \( r \) is commonly set to 2 (using the \( L^2 \)-norm to measure distance). Given an observed \( S_n = \{x_1, \ldots, x_n\} \), the empirical version of the DTM function at some \( y \in \mathbb{R}^d \) is

\[
\hat{d}_{m_0}(y) = \left( \frac{1}{k} \sum_{i \in N_k(y)} \|x_i - y\|^r \right)^{1/r},
\]

where \( k = \lceil m_0 n \rceil \) (the ceiling of \( m_0 n \)) and \( N_k(y) \) is the set containing the \( k \) nearest neighbors of \( y \) among \( x_1, x_2, \ldots, x_n \). For more properties of the DTM function, see Chazal et al. (2017).

Lower-level sets of \( \hat{d}_{m_0} \) from Eq. (3) can be used to construct the sequence of homology groups in \( \mathbb{R}^d \), where each homology generator has a birth time \( b_i \) and a death time \( d_i \). Then a persistence diagram is generated from the set of birth and death times \((b_1, d_1), (b_2, d_2), \ldots, (b_m, d_m)\). For example, the minima of the \( \hat{d}_{m_0}(y) \) (if present) are born early in the filtration as \( H_0 \) generators. As the filtration parameter \( t \) increases, homology group generators merge together leading to the death of some of the \( H_0 \) generators, and eventually all merge into a single connected component. Additionally, some \( H_0 \) generators can merge in such a way that an \( H_1 \) generator, a closed loop, is born. As the filtration parameter, \( t \), increases, the loop is filled in, indicating the death of the loop. Figure 3 shows an example of a DTM function and its corresponding persistence diagram produced. Figure 3a presents the point cloud we used to generate a persistence diagram, which contains three disconnected loops. Figure 3b and Figure 3c show the DTM function in 3D and as colored contours over a grid of points, respectively. The green plane in Figure 3b shows a threshold at the DTM value 3. The persistence diagram in Figure 3d has two types of points: the black dots are \( H_0 \) generators, and the red triangles are \( H_1 \) generators. The x-axis and y-axis represent birth time and death time, respectively. The farther a point is from the diagonal, the longer the homology group generator is present in the filtration. There are three distinct \( H_1 \) generators far from the diagonal, which is consistent with the three loops in the point cloud data.

2.3. Confidence band for persistence diagrams

The lifetime of a homology group generator \( i \) is called its persistence, defined as \( d_i - b_i \), where \( b_i \) and \( d_i \) are its birth time and death time, respectively. It is common to interpret more persistent generators (longer lifetimes) as topological signal and less persistent generators (shorter lifetimes) as topological noise. A method for defining confidence sets on a persistence diagram and this type of topological signal from topological noise was developed in Fasy et al. (2014). To define these confidence sets, or to carry out other types of statistical inference, about a persistence diagram, an appropriate metric on the space of persistence diagrams is needed. The bottleneck distance, \( W_\infty(\cdot, \cdot) \), is a popular metric for quantifying the distance between two persistence diagrams. It is defined as \( W_\infty(U, V) = \inf_{\eta: U \to V} \sup_{u \in U} \|u - \eta(u)\|_\infty \), where \( \eta \) is a bijection between two

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1 The DTM function can be understood as a smoothed version of the distance function.
Figure 3: (a) A simulated dataset, $S_n$. (b) The DTM function of $S_n$ in 3D over a grid of points. The green plane shows a threshold corresponding to a DTM value of 3. (c) The DTM function in colored contour over a grid of points. (d) The persistence diagram of $S_n$ using the lower level sets of the DTM. There are several black points and three red points obviously apart from the diagonal, which is consistent with the point cloud: three loops. The pink band is 90% confidence band and is used as a method for distinguishing topological signal from noise, which is introduced in §2.3.
persistence diagrams that matches each point on persistence diagram $U$ to a point on persistence diagram $V$. The diagonal (i.e., where $b_i = d_i$) is considered to have an infinite number of points so if there is an imbalance in the number of topological generators between the two diagrams, any excess of points can be mapped to the diagonal. The $||a - v||_\infty = \max|b_i - h_i|, |d_i - d_i|$, with $u = (b_i, d_i)$ and $v = (b_i, d_i)$, where $v = \eta(a)$.

Let $f$ be the DTM function of the true data distribution $M$ and $\hat{f}_n$ be the empirical DTM function of the point cloud data $S_n$, with $Dgm_p(f)$ as the persistence diagram of homology dimension $p$ for $f$ and $Dgm_p(\hat{f}_n)$ be the persistence diagram of dimension $p$ for $\hat{f}_n$. Following the notation of Fasy et al. (2014), for a given statistical significance level $\alpha \in (0, 1)$, we will find a constant $c_\alpha = c_\alpha(X_1, X_2, \ldots, X_n)$ such that

$$\limsup_{n \to \infty} P(W_n(Dgm_p(\hat{f}_n), Dgm_p(f)) > c_\alpha) \leq \alpha. \quad (4)$$

Hence $C_\alpha = [0, c_\alpha]$ is an asymptotic $1 - \alpha$ confidence set for $W_n(Dgm_p(\hat{f}_n), Dgm_p(f))$ (Fasy et al. 2014).

This result implies that a square $1 - \alpha$ confidence set can be centered on each point of $Dgm_p(\hat{f}_n)$ with a side of length $2c_\alpha$. If this box for some generator $i$ intersects the diagonal line on the persistence diagram (where the birth time = death time), then generator $i$ can be interpreted as topological noise at the significance level $\alpha$ (i.e., the $p$-value of generator $i$ would be greater than $\alpha$); if this box for some generator $i$ does not intersect the diagonal line, then generator $i$ can be interpreted as topological signal at the significance level $\alpha$. Instead of considering the separate confidence sets around each generator, a confidence band can be added to a persistence diagram using a band of width $\sqrt{2c_\alpha}$ on the diagonal of $Dgm_p(\hat{f}_n)$ where points in the band are considered indistinguishable from noise and points outside the band are considered significant topological generators. See Figure 3d for an example of a confidence band for the $H_1$ group generators.

The following bootstrap procedure, as described in Fasy et al. (2014), can be used as an option for computing the confidence band. First, randomly sample a new point cloud sample, $S_n^*$, from the original data set $S_n$. Next, compute a persistence diagram $Dgm_p(\hat{f}_n^*)$ for $S_n^*$, and calculate

$$w = W_n(Dgm_p(\hat{f}_n), Dgm_p(\hat{f}_n^*)).$$

Repeat the bootstrap sampling $N_{\text{boot}}$ times such that $N_{\text{boot}}$ persistence diagrams and bottleneck distances are obtained.

The empirical distribution of $w$ can be used to approximate the distribution of $W_n(Dgm_p(\hat{f}_n), Dgm_p(f))$. Extracting the $1 - \alpha$ quantile from this distribution, we get the value $c_\alpha$ for $1 - \alpha$ confidence band. For example, Figure 4 displays the bootstrapped bottleneck distances from the dataset in Figure 3a and the $c_\alpha$ for a 90% confidence interval is displayed as a vertical, dashed, red line.

The persistence diagram in Figure 3d displays the confidence band for $H_1$ corresponding to $\alpha = 10\%$. On the diagram, the red triangles outside the pink band can be considered a statistically significant topological generator of $H_1$ at a significance level of $10\%$.

This framework will be used to identify the statistically significant filament loops ($H_1$ generators) and cosmic voids ($H_2$ generators) on a persistence diagram. Rather than using the confidence band construction, we will assign individual $p$-values to each topological generator on a persistence diagram. This will be discussed in more detail in the next section.

3. Method

SCHU involves (i) computing $p$-values of the persistent homology group generators of a dataset by adapting the framework from §2.3 and (ii) addressing how to find a representation (i.e., physical locations and boundaries) of the $H_1$ and $H_2$ homology group generators from the persistence diagram back in the physical volume. As mentioned previously, one challenge in finding a representation of a homology group generator from the persistence diagram back in the data is that there are many possible representations of each homology group generator. In order to find a representation of the filament and cosmic voids in the data we exploit information available during the persistent homology computation. In this section, we present the details of SCHU for locating statistically significant filament and cosmic voids in the data.

3.1. $p$-values for filament loops and cosmic voids

Rather than computing the confidence bands from §2.3, $p$-values can be assigned for the individual homology group generators appearing on a persistence diagram. By changing the value of $c_\alpha$ from Eq. (4), a corresponding probability $1 - \hat{F}_{n,p}(w)$ can be obtained from the bootstrap empirical distribution function of $w$, denoted as $\hat{F}_{n,p}(w)$. For a generator $i$ on $Dgm_p(\hat{f}_n)$ with persistence $x_i$ (i.e., $x_i = d_i - b_i$), the shortest distance between the generator and the diagonal is $\frac{x_i}{\sqrt{n}}$. Noting that the bottleneck distance only considers vertical and horizontal distances (i.e. distances in the death time and birth time directions, respectively), the lowest significance level where generator $i$ would not be considered topological noise (i.e. the generator’s $p$-value) corresponds to a particular quantile of $\hat{F}_{n,p}(w)$ based on a bottleneck distance of $\frac{x_i}{\sqrt{n}}$. For example, in Figure 5, generator $i$ is the solid red triangle with persistence $x_i$. The $L^\infty$ norm distance from $i$ to the diagonal is $\frac{x_i}{\sqrt{n}}$. Any point with $L^\infty$ norm distance less than $\frac{x_i}{\sqrt{n}}$ to $i$ will be located within the pink box as shown in Figure 5. Thus, the lowest significance level where generator $i$ would not be rejected as noise corresponds to a bottleneck distance of $\frac{x_i}{\sqrt{n}}$.

Instead of using a specific significance level $\alpha$ as a cutoff as in §2.3, the individual $p$-values can be assigned as follows. Let $x_i$ be the persistence of generator $i$ on the observed persistence diagram $Dgm_p(\hat{f}_n)$. Then the $p$-value assigned to generator $i$ is

$$p\text{-value}(i) = 1 - \hat{F}_{n,p}\left(\frac{x_i}{\sqrt{n}}\right), \quad (5)$$

where a longer lifetime, $x_i$, results in a smaller $p$-value. Generators within the $1 - \alpha$ confidence band discussed in §2.3 suggests that the $p$-values are greater than $\alpha$ (i.e., the generators are considered indistinguishable from topological noise), and generators on or outside the confidence band have $p$-values less than
Bottleneck Distance
Frequency
0.0 0.5 1.0 1.5
0 500 1000 1500 2000

(a) Distribution of \( W_\infty(Dgm(\hat{f}_n), Dgm(\hat{f}_n^*)) \)

Bottleneck Distance
Frequency
0.1 0.2 0.3 0.4 0.5 0.6
0 500 1500 2500

(b) Distribution of \( W_\infty(Dgm(\hat{f}_n), Dgm(\hat{f}_n^*)) \)

Figure 4: Empirical distribution of the bottleneck distances between the persistence diagram of the data DTM, and the persistence diagrams of the bootstrap realization DTM, \( W_\infty(Dgm(\hat{f}_n), Dgm(\hat{f}_n^*)) \). Red dashed lines are the 90% quantiles. The number of bootstrap samples, \( N_{\text{boot}} \), is \( 10^4 \). The dataset used here is the same dataset shown in Figure 3a.

3.2. Locating filament loops and cosmic voids in the cosmological volume

Once a persistence diagram is obtained from a data set with \( p \)-values assigned to the topological generators, a natural question is where these generators (i.e., the holes) are located in the original data. For example, if statistically significant filament loops and cosmic voids exist in a given dataset, locating a representation of these voids back into the data set could be helpful for studying properties of the voids (Bos et al., 2012).

The points on a persistence diagram represent homology group generators so there are potentially many representations of the homology groups in the original dataset. In this section, we present a way of obtaining a representation of the homology groups using the quantities calculated during the persistent homology computations.

Additionally, the DTM affects the representations of the homology group generators through the resolution of the grid and the selected \( m_0 \). In Figure 6a, Figure 6c, and Figure 6e, there are three different representations of the \( H_1 \) generator shown in red circles. The three different representations correspond to contours of three lower level sets of three different DTM functions. In Figure 6b, Figure 6d, and Figure 6f, the contour maps of DTM are shown and the three representations are marked in black circles. The representation reported from persistent homology is the inner contour of the smallest lower level set that forms a \( H_1 \) generator (which will be explained later). Thus, the representation is determined by the DTM function constructed over the dataset.

or equal to \( \alpha \) (i.e., it is considered topological signal). Points on a persistence diagram outside the confidence band with confidence level 90% have \( p \)-values less than or equal to 10%.
In order to compute a DTM function, a grid needs to be defined with a specified grid size: for example, see the grid displayed in Figure 7a. Then the DTM is computed for each point on the grid using Eq. (3). The grid size and \( m_0 \) in Eq. (3) are two key parameters that influence resulting representations. The grid size determines the resolution of DTM and \( m_0 \) determines the smoothness of DTM. Figure 6b and Figure 6d have the same \( m_0 \) (smoothness) but different grid sizes. Figure 6b and Figure 6f have the same grid size but different \( m_0 \).

In the persistent homology algorithm used in this work, there is a way to record homology group generator locations during the computation of a persistence diagram (Edelsbrunner et al., 2002; Zomorodian and Carlsson, 2005; De Silva et al., 2011). The method for finding generators from persistence diagrams back in the original data uses the output of the computation of the persistence diagram. To understand the persistent homology computing, we first provide more details on the filtration construction on a grid.

As mentioned above, a grid needs to be defined to compute a DTM function, as displayed in Figure 7a. After the DTM is computed using Eq. (3), a sequence of lower level sets \( L_t = \{ x : d_{m_0}(x) \leq t \} \) are determined (see §2.2). See Figure 7b for an example of a lower level set on a grid using an arbitrary threshold \( t \), which consists of a subset of grid vertices drawn as red points.

A simplicial complex (Edelsbrunner and Harer, 2010; Zhu, 2013) is constructed using the lower level sets at threshold \( t \) (i.e., the vertices of the grid points with corresponding DTM values lower than the threshold, such as the red points displayed in Figure 7b). A simplicial complex \( K \) is a set of (possibly different order) simplices such that (i) any face of a simplex of \( K \) is also a simplex in \( K \), and (ii) the intersection of any two simplices in \( K \) is a face of both simplices or empty. In Figure 7b, we show the simplicial complex from the arbitrary lower level set displayed. The simplicial complex includes the red points (0-simplices), the blue segments (1-simplices), and the cyan triangles (2-simplices). Note that for the arbitrary threshold selected in the figure, the red points connected with blue segments represented a single connected component (i.e., an \( H_0 \) generator). As the lower level set threshold increases, eventually the red points would be connected in such a way that a loop forms (i.e. an \( H_1 \) generator).

The boundary of a p-simplex is the set of (p-1)-simplex faces (Zhu, 2013). A boundary matrix \( D \) records the boundary information of the simplicial complex across the filtration. As the DTM threshold increases, simplices get added to \( D \). Each row and each column represents a simplex: \( D_{i,j} = 1 \) if the i-th simplex is a boundary of the j-th simplex and \( D_{i,j} = 0 \) if not. The details of the persistent homology computations (which take \( D \) to a reduced form of \( D, R \)) are beyond the scope of this paper, but an interested reader can see Edelsbrunner et al. (2002), Zomorodian and Carlsson (2005), De Silva et al. (2011), David Cohen-Steiner (2006) for details. The reduced boundary matrix \( R \) contains the information about the various generators of a persistence diagram. If the j-th column of \( R \) represents an \( H_p \) generator (since the j-th simplex is a p-simplex) contains at least one non-zero element, the non-zero element(s) indicate which (p-1)-simplices formed the generator. The generator disappears at the time in the filtration when the j-th simplex appears (De Silva et al. 2011). The non-zero columns of matrix \( R \) are used as representations of the homology group generators. For example, suppose column \( j \) of \( R \) represents an \( H_1 \) generator with only three non-zero elements at positions \( i, i+1, \) and \( i+2 \). Then simplex \( i, i+1, \) and \( i+2 \) are 1-simplices (i.e., segments) that can be used as a representation for the \( H_1 \) generator (i.e., the loop) of column \( j \) of \( R \). In practice, the vertices (i.e., the grid points) of the p-simplices can be used as the representation of an \( H_p \) generator.

During the computation, lower level sets are constructed using an increasing threshold \( t \). The lower level sets are in an increasing order, since a later set contains an earlier set. For a \( H_1 \) or \( H_2 \) generator, there exists a lower level set that first forms the \( H_1 \) or \( H_2 \) generator along the filtration, which is the smallest lower level set that forms the \( H_1 \) or \( H_2 \) generator. This smallest lower level set is generally what appears in the boundary matrix for representing that homology group generator. In general, the representation is the inner contour of the smallest lower level set that forms the \( H_1 \) or \( H_2 \) generator.

Remark 1: For \( H_1 \) generators, when there are points inside the loop, the representation returned by the algorithm can include an extra loop around the inner points. Figure 8 illustrates this special case. The representation is shown in red and includes two closed loops. However, only the larger loop is relevant as the inner points do not contribute to the \( H_1 \) generator and is an artifact of the algorithm. As discussed in §3.2, the homology group generator representation corresponds to a lower level set of the DTM function. Figure 8b shows the DTM function of the data in Figure 8a. The green surface is the threshold for the lower level set that corresponds to the \( H_1 \) generator represented in red in Figure 8a. Because of the inner points, the lower level set is empty around the inner points producing two closed loops. SCHU includes a step to remove the artificial inner loop from the representation.

Remark 2: In order to define a DTM for a given dataset, the grid resolution and \( m_0 \) need to be selected. The grid resolution should be high enough to pick out features at a meaningful physical scale; generally the higher the resolution the better. A statistically rigorous method for selecting \( m_0 \) for persistent homology is still an open question. However, Chazal et al. (2017) provide two suggestions for selecting \( m_0 \) using computationally intensive approaches (e.g. one approach selects the \( m_0 \) that maximizes the number of statistically significant homology group generators). Since there is not a statistically rigorous or computationally feasible approach for selecting \( m_0 \), we selected an \( m_0 \) so that the number of nearest neighbors was around 50. If \( m_0 \) is selected to be too small then the resulting DTM may be too noisy to isolate important homological signals; if \( m_0 \)
To compute a DTM function, a grid is generated with a specified grid size. The grid size and $m_0$ are two key parameters that influence resulting representations. (a), (c), and (e): Three representations of the same $H_1$ generator. (b), (d), and (f): Three contour maps of DTM functions with three representations marked in black circles.
selected to be too large, the resulting DTM may be too smooth and important homological signals may get washed out.

4. Application to Voronoi Foam Data

In order to demonstrate the performance of SCHU for finding statistically significant generators on a persistence diagram and then locating those generators in the original data, we consider a simulation study using data that mimic the large-scale structure of the Universe and focus on locating cosmic voids using $H_2$ generator representations. In the simulation study, we know the ground truth of where the voids are located and so can test its ability to find the true voids.

The generated data used in this study come from an approximation to the Voronoi foam model \cite{VoronoiFoamModel}, which can be considered an approximation of the large-scale structure of the Universe \cite{VoronoiFoamModel}. Our implementation of the Voronoi foam model begins with a Voronoi tessellation on a random set of points, or seeds, in the 3D simulation cube volume. These seeds are referred to as void points because the Voronoi cells become the cosmic voids. An illustration example of a 6-cell Voronoi foam simulation dataset is shown in Figure 9a where void points are drawn in red and the generated Voronoi foam data are drawn in black. A grid is generated over the 3D simulation cube at a specified resolution. Each grid point is assigned a void label indicating the nearest void point (i.e., indicating in which Voronoi cell the point is). For each grid point, its neighboring grid points are checked to see how many different void labels are assigned to its neighbors. If there are more than three different labels, the grid point is considered as a cluster point. This is because this grid point is equal-distance to more than three void points and so it is at the intersection of more than three Voronoi cells. Similarly, the grid point is considered as a filament point if there are exactly three different labels, and is considered as a wall point if there are exactly two different labels. If there is just one label, the grid point is considered within a void region. Then we have a set of cluster points, a set of filament points and a set of wall points. With specified proportions ($f_c$ for clusters, $f_f$ for filaments, and $f_w$ for walls), certain numbers of cluster points, filament points and wall points are sampled from these sets. In order for the points not to fall exactly on the grid, Gaussian noise with specified standard deviation, $\sigma_{\text{perturb}}$, is added to the sampled points. In Figure 9b, perturbed cluster points, filament points, and wall points are drawn as blue x’s, green pluses, and orange triangles, respectively. Also, a fraction, $f_n$, of clutter noise points are sampled uniformly over the simulation cube, shown in Figure 9b as purple diamonds. The perturbed cluster points, filament points and wall points, together with clutter noise points, form a Voronoi simulation dataset, with $f_c + f_f + f_w + f_n = 1$.

A persistence diagram is computed for a realization of the Voronoi foam simulation in order to find the cosmic voids (i.e., the $H_2$ generators). Since the positions of the seeds are known (i.e., the locations of cosmic voids), we are able to compare the locations of cosmic voids found using SCHU to the true void points to check how well we are able to locate a representation of the $H_2$ generators in the data. For $H_2$, the representation defines the surface of the void, and thus the average of the representation points is approximately the volume center.

The Voronoi foam simulated cosmological volume is represented by a cubic lattice with 100$h^{-1}$Mpc per dimension and a grid spacing of $1h^{-1}$Mpc. In order to initiate the foam sim-
Corresponding DTM with a threshold hyperplane.

Figure 8: (a) The red lines are the representation returned by the persistent homology algorithm. There are two separate loops in this representation, but there is only one meaningful $H_1$ generator representation (the larger red loop). (b) The red surface shows the DTM function for the dataset. The green surface is the threshold for the lower level set that corresponds to the representation in (a). The intersection of the red and green surfaces has two separate closed loops, which are returned as the representation in (a).

5 We tried various environment fractions, including values based on Table 2 of Libeskind et al. (2018), and found that the void/filament loop identification and significance levels are relatively insensitive to these ratios.
5.1. Cosmic voids and filament loops in SDSS

In this section, we identify filaments and voids in the SDSS main galaxy redshift survey from the Public Cosmic Void Catalog\(^6\) and compare our cosmic voids to the cosmic voids identified in Sutter et al. (2012). This catalog contains 63,639 galaxies within the redshift range 0.0 < z < 0.05. We only use the part of the catalog from the large contiguous region of the SDSS footprint in the northern part of the sky, which contains 57,795 galaxies, within 0.81 ≤ RA ≤ 261.25 and −3.71 ≤ Dec ≤ 70.13. Artificial boundary walls are added on the 3D boundary surface of the grid in order to detect generators that may extend beyond the boundary. In the celestial coordinate system (RA, DEC, z), the dataset is like a cylinder with the redshift direction as its height. We impose a grid on the side surface of the cylinder and randomly sample points from the top surface (the bottom surface is no longer a boundary after the transformation). A total of 11,130 grid points along the cylinder wall plus 3,000 sampled points at the top surface form the artificial walls. Then, the dataset is transformed into the Cartesian coordinate system and a DTM function is computed with an \(m_0\) of 0.001, which corresponds to using the 58 nearest neighbors when calculating the DTM function at each grid point. The resulting persistence diagram is displayed in Figure 11, and confidence bands are added as a reference (the 90% confidence band for \(H_1\) is drawn in pink, the 90% confidence band for \(H_2\) is drawn in blue, and the pink band shows the overlap). Using the bootstrap-derived empirical distribution of the bottleneck distances with \(N_{\text{boot}} = 1000\), the \(p\)-values were computed for all the \(H_1\) and \(H_2\) generators. In Figure 12, we highlight 10 of the most significant filament loops and 15 of the most significant cosmic voids found by SCHU. Figure 12a shows the 10 most significant filament loops in different colors (\(p\)-values < 0.001) and Figure 12b shows the 15 most significant cosmic voids (\(p\)-values < 0.09). As expected, the highlighted filament loops are surrounding sparse areas and the highlighted cosmic voids are located in areas with low galaxy density; this indicates that SCHU is indeed capable of identifying reasonable filament loops and cosmic voids based on our vi-

\(^6\)http://www.cosmicvoids.net/documents
Figure 10: (a) Voronoi foam data where the red points are the void points used to generate the Voronoi tessellation. (b) Volume centers of the 23 most significant $H_2$ generators with labels (c) Persistence diagram of (a), where the blue band is 90% confidence band for $H_2$. (d) Detailed shapes of the 23 voids found by SCHU. (e) For illustration, the red points show a representation of a $H_1$ generator with $p$-value 0.011.
sual comparison with the data. We note that while most cosmic voids are clustered around the bottom right corner of the Cartesian box (i.e. the area with lower redshift values), the identified filament loops span the entire redshift range.

We compare the results from SCHU with the identified cosmic voids from VIDE (Sutter et al., 2012), which is based on the ZOBOV void-finding method (Neyrinck, 2008). The VIDE algorithm begins by constructing a Delaunay tessellation based on the galaxy positions and assigns a density to each galaxy based on the volume of its corresponding Voronoi cell. Then, a watershed procedure (Platen et al., 2007) is carried out to combine some neighboring cells and ultimately assign certain regions as cosmic voids. To aid in visualizing the cosmic void locations and to easily compare the TDA-identified voids with those from VIDE, we use a similar rotation and subset as in Sutter et al. (2012) (i.e., the galaxy and void positions are rotated about the y-axis so that they lie on the x – y plane, and only those galaxies and voids within a 20 degree opening angle are plotted). The volume centers of the voids found by VIDE are plotted as red triangles in Figure 13.

For SCHU, the same DTM function as described above is used to generate the filtration with $m_0 = 0.001$. Since VIDE locates cosmic voids, we only compare cosmic voids ($H_2$ generators) here. After bootstrapping with $N_{\text{boot}} = 1000$, $p$-values for $H_2$ generators are obtained. The volume centers of the seven statistically significant voids found using SCHU ($p$-values < 0.1) are plotted as green pluses in Figure 13 with labels A through G; the corresponding $p$-value can be found in Table 2. Some of the voids found by VIDE are located in non-empty regions; however, these may be the smaller voids ($5 < 15h^{-1}$ Mpc) that Sutter et al. (2012) note are located at the edges of filaments and walls (non-empty regions). The cosmic voids found using the proposed TDA approach are concentrated more in empty regions. VIDE does not find the voids in the bottom region of Figure 13 while the proposed approach classifies those empty regions as cosmic voids. The orange pluses in Figure 13 are the artificial boundary points added to detect boundary voids; the side walls appear thick because of the projection.

Table 2: $p$-values for corresponding cosmic voids in Figure 13

| Label | $p$-value |
|-------|-----------|
| A     | <0.001    |
| B     | <0.001    |
| C     | <0.001    |
| D     | 0.064     |
| E     | <0.001    |
| F     | <0.001    |
| G     | 0.006     |

5.2. Cosmic web identification method comparison simulation

In this section, we compare SCHU with other void-finding methods by applying SCHU to the cosmological simulation used in the Libeskind et al. (2018) comparison study, created via the GADGET-2 dark-matter only N-body simulation code with $512^3$ particles (Springel, 2005). The GADGET-2 run uses the GADGET-2 dark-matter only N-body simulation code with $512^3$ particles (Springel, 2005). Libeskind et al. (2018) used this dataset in order to compare several cosmic environment classification methods; the data and the results for the void-finding methods considered are available online.

The simulation cube is $200h^{-1}$ Mpc on each side and was run using the 2015 Planck $\Lambda$CDM parameters (Planck Collaboration et al., 2016b). Dark matter halos were identified in the $z = 0$ snapshot using a friends-of-friends (FOF) algorithm (Huchra and Geller, 1982; Press and Davis, 1982) with a linking length of 0.2 and requiring a minimum of 20 particles per halo. There are a total of 281,465 halos in the catalog. Applying SCHU to a halo catalog is similarly well-motivated to that of a galaxy survey catalog, as dark matter halos (and the observable galaxies that lie within them) are both assumed to be biased tracers of the underlying matter field. Thus, underdensities in the halo field should correspond to physical voids in the matter distribution.

Libeskind et al. (2018) specified a $200 \times 200 \times 200$ grid for cosmic environment prediction in order to facilitate the comparison of the methods; we also use this same grid for our predictions. To detect boundary voids, we add artificial boundary walls by randomly sampling $1,000$ points on each of the six sides of the cube. The $m_0$ is set to 0.0002, which corresponds to using $56$ nearest neighbors when calculating the DTM function at each grid point. Figure 14 displays the resulting persistence diagram of the dataset along with the confidence bands. Figure 15a shows the 10 most significant filament loops with

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Figure 11: Persistence Diagrams with 90% confidence band for $H_1$ and $H_2$ of the SDSS dataset (with artificial boundary walls). A DTM function with $m_0 = 0.001$ is used to generate the persistence diagrams. The confidence band for $H_1$ is drawn in pink and the confidence band for $H_2$ is drawn in blue: the $H_1$ band is the overlap since it is shorter.

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http://data.aip.de/tracingthecosmicweb/

$h = 0.68$, $\Omega_m = 0.3$, $\Omega_{\Lambda} = 0.7$, $n_s = 0.96$ and $\sigma_8 = 0.82$
Figure 12: Filament loops (a) and voids (b) identified in the SDSS dataset using SCHU. The DTM function was constructed with $m_0 = 0.001$. The most significant 10 filament loops and the most significant 15 cosmic voids generators are shown in different colors. In the left figure, the 10 most significant filament loops are displayed with $p$-values < 0.001; in the right figure, the 15 most significant cosmic voids are displayed with $p$-values < 0.09.

Figure 13: Comparison between VIDE and SCHU. A slice of the SDSS dataset within a 20 degree opening angle after the rotation projection is shown. The red triangles are barycenters of voids found by VIDE; the green pluses are cosmic voids with $p < 0.1$ and the corresponding $p$-values can be found in Table 2.

$p < 0.001$ in different colors and Figure 15b shows the 15 most significant cosmic voids with $p < 0.001$ in different colors. Cosmic voids are more straightforward to visualize and we can see that they are located at low density areas, as expected.

We compare SCHU with the nine different methods used in Libeskind et al. (2018), the Multiscale Morphology Filter-2 (MMF-2, Aragon-Calvo and Yang (2014)), Multi-Stream Web Analysis (MSWA, Ramachandra and Shandarin 2015).
CLASSIC (Kitaura and Angulo 2012), DisPerSE (Sousbie 2011), NEXUS+ (Cautun et al. 2012), Spinweb (Aragón-Calvo et al. 2010), ORIGAMI (Falck et al. 2012; Falck and Neyrinck 2015), the tidal shear tensor (T-web, Forero-Romero et al. 2009), and the velocity shear tensor (V-web, Hoffman et al. 2012). For a brief overview of these methods, we refer the reader to Table 1 of Libeskind et al. (2018). These methods are broadly classified as Hessian (CLASSIC, T-web, V-web), scale-space Hessian (MMF-2, NEXUS+), topological (DisPerSE, Spinweb), and phase-space (ORIGAMI, MSWA) methods. The Hessian methods use the Hessian of the gravitational potential (Hahn et al. 2007) or velocity shear tensor (Hoffman et al. 2012) in order to classify the cosmic environment type by counting the number of eigenvalues of the resulting matrix at each point in space. The scale-space Hessian methods are based on the Multiscale Morphology Filter (MMF) approach (Aragón-Calvo et al. 2007), which extends the Hessian method into a smoothing-scale-independent calculation by computing the optimum eigenvalue threshold after repeating the calculation across many different smoothing scales. The topological methods compute the topological structures of a reconstructed density field from the spatial datasets that can be constructed in various ways, such as using a Voronoi tessellation or a discrete Morse-Smale complex. SCHU falls into the topological category of cosmic environment classifiers. Lastly, the phase-space methods identify the cosmic environment based on the number of dimensions along which particles have shell-crossed.

A 2h−1 Mpc-thin slice of the simulation cube was selected to aid in the comparison of the methods, and is displayed in Figure 14. Persistence Diagrams with 90% confidence band for \( H_1 \) and \( H_2 \) from the Libeskind et al. (2018) dataset. A DTM function with \( m_0 = 0.0002 \) is used to generate the persistence diagrams. The confidence band for \( H_1 \) is drawn in pink and confidence band for \( H_2 \) is drawn in blue: they are overlapping so that the pink band, which is shorter, is the overlap.

Figure 14

The cosmic void regions found by the various methods span a range from visually smooth (larger contiguous regions defined as cosmic voids) to rough (many small regions defined as cosmic voids). Results of CLASSIC, MSWA, NEXUS+, ORIGAMI, and V-web in Figures 16d, 16g, 16h, 16i, and 16l, respectively, have similar appearances where the white regions appear tightly constrained to high density regions of matter and the remaining majority of space is defined as cosmic voids. Within those five methods, CLASSIC and V-web look smoother with larger contiguous high-density regions, while the other three methods look rougher with thinner, weblike, high-density regions. SCHU, DisPerSE, Spinweb, and T-web, Figures 16b and 16c, respectively, share visual similarities in that the white regions appear tightly constrained to low density regions of matter and the remaining majority of space is defined as matter structures. Persistent homology, DisPerSE, and T-web are smoother with large contiguous cosmic void regions, whereas Spinweb is rougher with many small cosmic void regions. MMF-2 is different from all the other methods as it identifies many disjoint small clusters of matter in regions where there are minimal halos within the slice.

5.3. Betti functions of the MassiveNuS simulations

Next we provide a demonstration of the potential usefulness of persistent homology by considering two simulations from the MassiveNuS suite (Liu et al. 2018). These simulations are dark matter-only, and trace the evolution of 10243 particles in a cubic volume of side length 512h−1 Mpc. The suite consists of 100 simulations with varied values of three cosmological parameters: the neutrino mass sum \( \sum m_\nu \), the total matter density \( \Omega_m \), and the primordial power spectrum amplitude \( A_s \). In the following analysis, we study the publicly-available Rockstar (Behroozi et al. 2013) halo catalogs generated from the \( z = 0 \) snapshots of two of these simulations. The two selected simulations both have \( \Omega_m = 0.3 \) and \( A_s = 2.1 \times 10^{-9} \), but different neutrino mass sums of \( \sum m_\nu = 0 \) eV and \( \sum m_\nu = 0.6 \) eV, which we denote \( S_1 \) and \( S_2 \), respectively. In this case, the goal is to use persistent homology to summarize the two simulation datasets and analyze whether the summaries can discriminate the different cosmologies used to produce the data.

When using persistent homology to analyze datasets, the typical summary statistic obtained is a persistence diagram. However, directly comparing persistence diagrams generated from...
two datasets in such a way that the differences are clearly captured is not straightforward. The bottleneck distance is an option for comparing two persistence diagrams, but it does not describe the differences across the filtration. Instead, functional summaries of persistence diagrams can be used. For example, the Betti number for homology dimension $p$, denoted $B_p$, is the rank of $H_p$, which can be interpreted as the number of $p$-dimensional holes. In persistent homology, $B_p$ can be computed across the varying threshold values $t$ in order to produce a Betti function, $B_p(t)$. Intuitively, $B_p(t)$ is the number of $H_p$ generators that were born before $t$ and are still alive at $t$. As noted in the introduction, Betti functions were used in van de Weygaert et al. (2011) and Pranav et al. (2016) to compare the persistent homology of different dark energy models and cosmic matter distributions, respectively. Betti functions, or other functional summaries of persistence diagrams (e.g., Berry et al. 2018), are straightforward to use in a comparison.

Figure 17 shows an example of Betti functions. We apply SCHU to an eighth of the simulation volume such that the halos are placed on a grid of size $256h^{-1}$ Mpc per side with grid size of $1h^{-1}$ Mpc. For computing the DTM function, $m_0 = 0.0001$ is used, which corresponds to the inclusion of roughly the 40 nearest neighbors. We denote the 0-, 1-, and 2-dimensional Betti functions of $S_1$ as $B_0(S_1(t))$, $B_1(S_1(t))$, and $B_2(S_1(t))$, respectively, for $i = 1, 2$. The resulting Betti functions of $S_1$ and $S_2$ are shown in Figure 17a and Figure 17b, respectively. The Betti functions of $S_1$ and $S_2$ have similar shapes. We calculate the ratio for each of the $p$-dimensional Betti functions between $S_1$ and $S_2$, but first carry out mild smoothing to reduce noise fluctuations so that trends in the Betti function ratios can be easily captured.

In particular, a local polynomial regression model (Cleveland 1979) with an adaptive bandwidth that uses 10% of the nearest observations is considered in order to smooth the Betti functions before computing the ratios displayed in Figures 17c and 17d. The black solid line in Figure 17c, which denotes the ratio of the $B_0(t)$, deviates substantially from unity when $1 < t < 3h^{-1}$ Mpc and then begins to fluctuate again at $6 < t < 8h^{-1}$ Mpc. Both ratios of $B_1(t)$ and $B_2(t)$ display similar behavior at different values of $t$. The threshold values where the ratio between the Betti functions of the two simulations is largest correspond to the phases where a particular dimension of homology group generators is just beginning to form. Thus, at these points, the ratios are between small counts of generators and can become large and noisy.

However, in the intermediate values of the threshold where the homology group generators of a particular dimension are abundant, there are overall trends between the ratios for both $B_1(t)$ and $B_2(t)$. In Figure 17d, we focus on the normalized difference in the Betti functions between the simulations $(B_p(S_1(t)) - B_p(S_2(t))) / B_p(S_2(t))$ for the threshold values where there are more $p$-dimensional generators present in order to avoid large fluctuations due to low counts. The $H_1$ and $H_2$ generator abundances peak in the simulation with massive neutrinos ($S_2$) at a larger magnitude and at lower threshold values than the simulation without massive neutrinos ($S_1$). While these abundances by

The most significant 10 filament loops (a) and the most significant 15 cosmic voids generators (b) are shown in different colors. All of their $p$-values are less than 0.001.

Figure 15: Filament loops (a) and voids (b) identified in the Libeskind et al. (2018) dataset using SCHU. The DTM function was constructed with $m_0 = 0.0002$. The most significant 10 filament loops (a) and the most significant 15 cosmic voids generators (b) are shown in different colors. All of their $p$-values are less than 0.001.

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10The local polynomial regression model estimates a polynomial model using points in a local neighbourhood of the target point with weights defined based on distance from the target point. In this case, the neighborhood is specified to include the nearest 10% of observations.
Figure 16: Comparison of void-finding methods on the halo catalog of a $2h^{-1}$ Mpc slice of the cosmological simulation from Libeskind et al. (2018). (a) The halos in the slice of the simulation cube considered. (b) - (l) The various void-finding method results. The white regions denote voids and red regions denote walls, filaments, and nodes. (b) shows all $H_2$ generators in the slice from SCHU, whereas (c) shows only the $H_2$ generators with $p$-value < 0.1 from SCHU.
threshold are not directly an analog for physical size (the persistence of a generator would be a better proxy for size), the Betti functions do reveal an interesting difference between the simulations. The filament loops and voids form earlier in the filtration (i.e., at smaller distances) in the simulation that includes massive neutrinos ($S_2$), suggesting that the presence of massive neutrinos may act to change the density field in such a way that the filament loops and voids form at smaller distances; that is, the neutrinos affect the distribution of the dark matter halos along filaments and on the boundary of voids. Additional analysis of the dependence of these differences on redshift and the values of $(\sum m_\nu, \Omega_\Lambda, A_s)$ is needed to better understand the effect of neutrinos on the density field; this is a topic of future research. Overall, Figure 17d demonstrates that massive neutrinos have a distinct effect on the Betti functions, which could potentially be utilized in larger surveys and simulation datasets to help further constrain cosmological parameters and break degeneracies.

6. Conclusions and Discussions

In this work, we present a novel method, SCHU, for applying modern statistical methods from topological data analysis, specifically persistent homology, to identify filament loops and cosmic voids in astronomical datasets. While previous works used topological ideas to explore the underlying matter density field in order to study its Betti numbers and topological persistence [Pranav et al., 2016], SCHU strengthens and extends this by assigning p-values to individually-identified homology group generators as well as finding a representation of the statistically significant generators in the cosmological volume. The addition of this last capability enables simultaneous analysis of void clustering and abundance as well as analysis of the density field’s topology via persistence diagrams and Betti numbers. Furthermore, we emphasize the potentially complementary cosmological analysis that can be performed using filament loops, a subset of the cosmic filamentary distribution that SCHU can identify consisting of connected rings of filaments.

In this paper, we (i) introduce the necessary formalism that underlies persistent homology, (ii) detail SCHU for computing persistence diagrams and Betti numbers by embedding the cosmological dataset on a lattice and generating a discrete filtration using the lower-level sets of a DTM function, and (iii) apply SCHU to a controlled Voronoi foam simulation, a subset of the SDSS galaxy survey, a cosmological simulation from the Libeskind et al. (2018) cosmic environment identification survey, and a subset of the MassiveNuS simulation suite. We summarize our results as follows:

- By using an approximate Voronoi foam model of the cosmic web where the ground truth locations of the voids are known, we demonstrate that SCHU is capable of successfully identifying low-density void regions in cosmological datasets.
- Using a subset of SDSS survey data, we demonstrate that SCHU is able to locate statistically significant voids, and these voids are compared to those identified using VIDE (Sutter et al., 2012).
- We contrast the results of SCHU as applied to the Libeskind et al. (2018) comparison study, finding that while there is still no quantitative consensus between cosmic web environment classifiers, SCHU’s results do share similarities with those from various other tools, including Spinweb [Aragón-Calvo et al., 2010], DisPerSE [Sousbie, 2011], and T-web [Forero-Romero et al., 2009].
- The Betti functions of two simulations from the MassiveNuS suite [Liu et al., 2018] were analyzed with different values of $\sum m_\nu$, demonstrating that this statistic can potentially be used to help break cosmological parameter degeneracy when applied to larger datasets.

The methods presented in this paper lay the groundwork for a detailed follow-up analysis of the large-scale galaxy and matter density fields in state-of-the-art cosmological simulations. Data products that can be generated from SCHU have potential for constraining cosmological parameters and breaking degeneracy (e.g., $\Omega_\Lambda$ and $\sigma_8$) as well as discriminating between various models of dark energy, modified gravity, and massive neutrinos. With the imminent, massive influx of cosmological data that will soon be made available through next-generation surveys such as LSST and DESI, it is both important and timely that void-finding methods, such as the one presented here, are applied to realistic mock galaxy catalogs and lightcones in order to predict their model discrimination and parameter constraining power on these future datasets in a statistically rigorous way. Simulation suites that span the cosmological parameter space will provide a powerful base for applying SCHU to determine the dependence of void clustering, abundance, and Betti functions on the cosmological parameters. Summary statistics of these outputs could also be fed into a machine learning algorithm and trained to predict the underlying set of cosmological parameters. Based on our preliminary studies of the Betti functions of the MassiveNuS simulations, some example features of interest include the threshold values where homology group generators are first born and last appear, the threshold values of the peaks in the Betti functions, and the overall normalization of the Betti functions, for each homology group dimension and each simulation.

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Figure 17: Betti functions and ratios of two snapshots from the MassiveNuS simulation suite. The two datasets used are Rockstar halo catalogs generated from the $z = 0$ snapshots, both of which have $\Omega_m = 0.3$ and $A_s = 2.1 \times 10^{-9}$, but with different neutrino mass sums of $\sum m_{\nu} = 0$ eV (simulation $S_1$) and $\sum m_{\nu} = 0.6$ eV (simulation $S_2$). An eighth of the simulation volume is used such that the halos are placed on a grid of size $256h^{-1}$ Mpc per side with lattice spacing of $1h^{-1}$ Mpc. A DTM function with $m_0 = 0.0001$ is used to generate the Betti functions. (a) Betti functions of $S_1$. The black solid, red dashed, and blue dotted lines correspond to $H_0$, $H_1$, and $H_2$. (b) Betti functions of $S_2$. (c) Ratio of smoothed Betti functions of $S_1$ and $S_2$: $\frac{B^0_p(t)+1}{B^0_0(t)+1}$, where $p$ denotes the dimension of the homology group generators. (d) Normalized differences of smoothed Betti functions: $\frac{B^0_p(t)−B^0_0(t)}{B^0_0(t)+1}$. 

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