

Computing Persistent Homology *

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ABSTRACT

We study the homology of a filtered d -dimensional simplicial complex K as a single algebraic entity and establish a correspondence that provides a simple description over fields. Our analysis enables us to derive a natural algorithm for computing persistent homology over an arbitrary field in any dimension. Our study also implies the lack of a simple classification over non-fields. Instead, we give an algorithm for computing individual persistent homology groups over an arbitrary PIDs in any dimension.

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General Terms: Algorithms, Theory

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1. INTRODUCTION

In this paper, we study the homology of a filtered d -dimensional simplicial complex K , allowing an arbitrary PID D as the ground ring of coefficients. A filtered complex is an increasing sequence of simplicial complexes, as shown in Figure 1. It determines an *inductive system* of homology groups, i.e. a family of Abelian groups $\{G_i\}_{i \geq 0}$ together with homomorphisms $G_i \rightarrow G_{i+1}$. If the homology is taken with field coefficients, we obtain an inductive system of vector spaces over the field. Each vector space is determined up to isomorphism by its dimension. In this paper, we obtain a simple classification of an inductive system of vector spaces. The classification is in terms of a graded module over the graded ring $F[t]$ of polynomials over the field F , which can be parametrized up to

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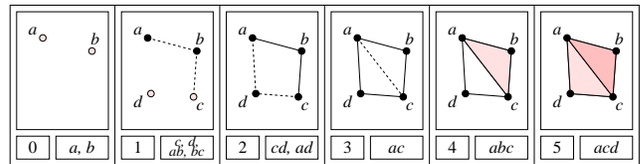


Figure 1: A filtered complex with newly added simplices highlighted.

isomorphism by finite families of intervals with integer endpoints. We also derive a natural algorithm for computing this family of intervals. Using this family, we may identify homological features that persist within the filtration, the *persistent homology* of the filtered complex.

Furthermore, our interpretation makes it clear that if the ground ring is not a field, there exists no similarly simple classification of persistent homology. Rather, the structures are very complicated, and although we may assign interesting invariants to them, no simple classification is, or is likely ever to be, available. In this case, we provide an algorithm for computing a single persistent group for the filtration.

In the rest of this section, we first motivate our study through three examples in which filtered complexes arise whose persistent homology is of interest. We then discuss prior work and its relationship to our work. We conclude this section with an outline of the paper.

1.1 Motivation

We call a filtered simplicial complex, along with its associated chain and boundary maps, a *persistence complex*. We will formalize this concept in section 3. Persistence complexes arise naturally whenever one is attempting to study topological invariants of a space computationally. Often, our knowledge of this space is limited and imprecise. Consequently, we must utilize a multi-scale approach to capture the connectivity of the space, giving us a persistence complex.

Example 1.1 (point cloud data) Suppose we are given a finite set of points X from a subspace $\mathbb{X} \in \mathbb{R}^n$. We call X *point cloud data* or *PCD* for short. It is reasonable to believe that if the sampling is dense enough, we should be able to compute the topological invariants of \mathbb{X} directly from the PCD. To do so, we may either compute the *Cech* complex, or approximate it via a *Rips* complex [11]. The latter complex $R_\epsilon(X)$ has X as its vertex set. We declare $\sigma = \{x_0, x_1, \dots, x_k\}$ to span a k -simplex of $R_\epsilon(X)$ iff

$d(x_i, x_j) \leq \epsilon$ for all pairs $x_i, x_j \in \sigma$. There is an obvious inclusion $R_\epsilon(X) \rightarrow R_{\epsilon'}(X)$ whenever $\epsilon < \epsilon'$. In other words, for any increasing sequence of non-negative real numbers, we obtain a persistence complex.

Example 1.2 (density) Often, our samples are not from a geometric object, but are heavily concentrated on it. It is important, therefore, to compute a measure of density of the data around each sample. For instance, we may count the number of samples $\rho(x)$ contained in a ball of size ϵ around each sample x . We may then define $R_\epsilon^r(X) \subseteq R_\epsilon$ to be the Rips subcomplex on the vertices for which $\rho(x) \leq r$. Again, for any increasing sequence of non-negative real numbers r , we obtain a persistence complex. We must analyze this complex to compute topological invariants attached to the geometric object around which our data is concentrated.

Example 1.3 (Morse functions) Given a manifold M equipped with a Morse function f , we may filter M via the excursion sets $M_r = \{m \in M \mid f(m) \leq r\}$. We again choose an increasing sequence of non-negative numbers to get a persistence complex. If the Morse function is a height function attached to some embedding of M in \mathbb{R}^n , persistent homology can now give information about the shape of the submanifolds, as well the homological invariants of the total manifold.

1.2 Prior work

We assume familiarity with basic group theory and refer the reader to Dummit and Foote [6] for an introduction. We make extensive use of Munkres [12] in our description of algebraic homology and recommend it as an accessible resource to non-specialists. There is a large body of work on the efficient computation of homology groups and their ranks [1, 4, 5, 9]. Persistent homology groups were initially defined only for three-dimensional simplicial complexes and coefficients in \mathbb{Z}_2 [7, 14]. The authors also provided an algorithm for generating a particular set of intervals for subcomplexes of S^3 over \mathbb{Z}_2 . Surprisingly, they showed that these intervals allowed the correct computation of the rank of persistent homology groups.

1.3 Our work

Viewing an inductive system of vector spaces as a single conceptual entity, we show that the algorithm in [7] extends to any inductive system of chain complexes and does not depend on simplicial complexes arising from coverings of sets in \mathbb{R}^3 . We extend and generalize this algorithm to arbitrary dimensions and ground fields by deriving it from the classic reduction scheme. In this manner, we illustrate how the algorithm derives its simple structure from the properties of the rich algebraic structures. We also show that if we consider integer coefficients or coefficients in some ring R , there is no similar simple classification of inductive systems of modules over R . This negative result suggests the possibility of interesting yet incomplete invariants of inductive systems. For now, we give an algorithm for classifying a single homology group over an arbitrary PID.

We finally observe that any filtered complex gives rise to a *spectral sequence*. In principle, we may use this method to compute the result of our algorithm. However, the method does not provide an *algorithm*, but a *scheme* that must be tailored for each problem independently. Our techniques, on the other hand, provide a complete and effective algorithm.

1.4 Outline

We review concepts from algebra and simplicial homology in Section 2. We also re-introduce persistent homology over integers and arbitrary dimensions. In Section 3, we define and study the persistence module, a structure that represents the homology of a filtered complex. In addition, we establish a relationship between our results and prior work. Using our analysis, we derive an algorithm for computation over fields in Section 4. For non-fields, we describe an algorithm in Section 5 that computes individual persistent groups. We conclude the paper in Section 7 with a discussion of current and future work.

2. BACKGROUND

In this section, we review the mathematical and algorithmic background necessary for our work. We begin by reviewing the structure of finitely generated modules over principal ideal domains. We then discuss simplicial complexes and their associated chain complexes. Putting these concepts together, we define simplicial homology and outline the standard algorithm for its computation. We conclude this section by describing persistent homology.

2.1 Algebra

Throughout this paper, we assume a ring R to be commutative with unity. A *polynomial* $f(t)$ with coefficients in R is a formal sum $\sum_{i=0}^{\infty} a_i t^i$, where $a_i \in R$ and t is the indeterminate. The set of all polynomials $f(t)$ over R forms a commutative ring $R[t]$ with unity. If R has no divisors of zero, and all its ideals are principal, it is a *principal ideal domain (PID)*, e.g. $\mathbb{R}, \mathbb{Q}, \mathbb{Z}, \mathbb{Z}_p$ for p prime, and $F[t]$ for F a field.

A *graded ring* is a ring $\langle R, +, \otimes \rangle$ equipped with a direct sum decomposition of Abelian groups $R \cong \bigoplus_i R_i$, $i \in \mathbb{Z}$, so that multiplication is defined by bilinear pairings $R_n \otimes R_m \rightarrow R_{n+m}$. Elements in a single R_i are called *homogeneous* and have *degree* i , $\deg e = i$ for all $e \in R_i$. A *graded module* M over a graded ring R is a module equipped with a direct sum decomposition, $M \cong \bigoplus_i M_i$, $i \in \mathbb{Z}$, so that the action of R on M is defined by bilinear pairings $R_n \otimes M_m \rightarrow M_{n+m}$. A graded ring (module) is *non-negatively graded* if $R_i = 0$ ($M_i = 0$) for all $i < 0$. We may grade $R[t]$ non-negatively with the *standard grading* $(t^n) = t^n \cdot R[t]$, $n \geq 0$.

The standard structure theorem describes finitely generated modules and graded modules over PIDs.

Theorem 2.1 (structure) If D is a PID, then every finitely generated D -module is isomorphic to a direct sum of cyclic D -modules. That is, it decomposes uniquely into the form

$$D^\beta \oplus \left(\bigoplus_{i=1}^m D / d_i D \right), \quad (1)$$

for $d_i \in D, \beta \in \mathbb{Z}$, such that $d_i | d_{i+1}$. Similarly, every graded module M over a graded PID D decomposes uniquely into the form

$$\left(\bigoplus_{i=1}^n \Sigma^{\alpha_i} D \right) \oplus \left(\bigoplus_{j=1}^m \Sigma^{\gamma_j} D / d_j D \right), \quad (2)$$

where $d_j \in D$ are homogeneous elements so that $d_j | d_{j+1}$, $\alpha_i, \gamma_j \in \mathbb{Z}$, and Σ^α denotes an α -shift upward in grading.

In both cases, the theorem decomposes the structures into free (left) and torsion (right) parts. In the latter case, the torsional elements are also homogeneous.

2.2 Simplicial Complexes

A *simplicial complex* is a set K , together with a collection \mathcal{S} of subsets of K called *simplices* (singular *simplex*) such that for all $v \in K, \{v\} \in \mathcal{S}$, and if $\tau \subseteq \sigma \in \mathcal{S}$, then $\tau \in \mathcal{S}$. We call the sets $\{v\}$ the *vertices* of K . When it is clear from context what \mathcal{S} is, we refer to set K as a complex. We say $\sigma \in \mathcal{S}$ is a k -*simplex of dimension* k if $|\sigma| = k + 1$. If $\tau \subseteq \sigma, \tau$ is a *face* of σ , and σ is a *coface* of τ . An *orientation* of a k -simplex $\sigma, \sigma = \{v_0, \dots, v_k\}$, is an equivalence class of orderings of the vertices of σ , where $(v_0, \dots, v_k) \sim (v_{\tau(0)}, \dots, v_{\tau(k)})$ are equivalent if the sign of τ is 1. We denote an *oriented simplex* by $[\sigma]$. A simplex may be realized geometrically as the convex hull of $k + 1$ affinely independent points in $\mathbb{R}^d, d \geq k$. A realization gives us the familiar low-dimensional k -simplices: *vertices, edges, triangles, and tetrahedra*, for $0 \leq k \leq 3$. Within a realized complex, the simplices must meet along common faces. A *subcomplex* of a simplicial complex K is a simplicial complex $L \subseteq K$. A *filtration* of a complex K is a nested subsequence of complexes $\emptyset = K^0 \subseteq K^1 \subseteq \dots \subseteq K^m = K$. For generality, we let $K^i = K^m$ for all $i \geq m$. We call K a *filtered complex*. We show a small filtered complex in Figure 1.

2.3 Chain Complex

The k th *chain group* \mathbf{C}_k of K is the free Abelian group on its set of oriented k -simplices, where $[\sigma] = -[\tau]$ if $\sigma = \tau$ and σ and τ are differently oriented. An element $c \in \mathbf{C}_k$ is a k -chain, $c = \sum_i n_i [\sigma_i], \sigma_i \in K$ with *coefficients* $n_i \in \mathbb{Z}$. The *boundary operator* $\partial_k: \mathbf{C}_k \rightarrow \mathbf{C}_{k-1}$ is a homomorphism defined linearly on a chain c by its action on any simplex $\sigma = [v_0, v_1, \dots, v_k] \in c$,

$$\partial_k \sigma = \sum_i (-1)^i [v_0, v_1, \dots, \hat{v}_i, \dots, v_k],$$

where \hat{v}_i indicates that v_i is deleted from the sequence. The boundary operator connects the chain groups into a *chain complex* \mathbf{C}_* :

$$\dots \rightarrow \mathbf{C}_{k+1} \xrightarrow{\partial_{k+1}} \mathbf{C}_k \xrightarrow{\partial_k} \mathbf{C}_{k-1} \rightarrow \dots$$

We may also define subgroups of \mathbf{C}_k using the boundary operator: the *cycle group* $\mathbf{Z}_k = \ker \partial_k$ and the *boundary group* $\mathbf{B}_k = \text{im } \partial_{k+1}$. We show examples of cycles in Figure 2. An important property of the boundary operators is that the boundary of a boundary is always empty, $\partial_k \partial_{k+1} = \emptyset$. This fact, along with the definitions, implies that the defined subgroups are nested, $\mathbf{B}_k \subseteq \mathbf{Z}_k \subseteq \mathbf{C}_k$, as in Figure 3. For generality, we often define null boundary operators in dimensions where \mathbf{C}_k is empty.

2.4 Homology

The k th *homology group* is $\mathbf{H}_k = \mathbf{Z}_k / \mathbf{B}_k$. Its elements are classes of *homologous cycles*. To describe its structure, we view

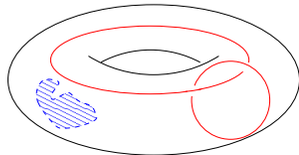


Figure 2: The dashed 1-boundary rests on the surface of a torus. The two solid 1-cycles form a basis for the first homology class of the torus. These cycles are non-bounding: neither is a boundary of a piece of surface.

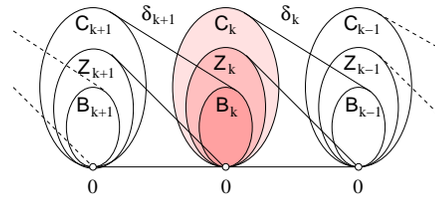


Figure 3: A chain complex with its internals: chain, cycle, and boundary groups, and their images under the boundary operators.

the Abelian groups we have defined so far as modules over the integers. This view allows alternate ground rings of coefficients, including fields. If the ring is a PID D, \mathbf{H}_k is a D -module and Theorem (2.1) applies: β , the rank of the free submodule, is the *Betti number* of the module, and d_i are its *torsion coefficients*. When the ground ring is \mathbb{Z} , the theorem above describes the structure of finitely generated Abelian groups. Over a field, such as \mathbb{R}, \mathbb{Q} or \mathbb{Z}_p for p a prime, the torsion submodule disappears. The module is a vector space that is fully described by a single integer, its rank β , which depends on the chosen field.

2.5 Reduction

The standard method for computing homology is the reduction algorithm. We describe this method for integer coefficients as it is the more familiar ring. The method extends to modules over arbitrary PIDs, however.

As \mathbf{C}_k is free, the oriented k -simplices form the *standard basis* for it. We represent the boundary operator $\partial_k: \mathbf{C}_k \rightarrow \mathbf{C}_{k-1}$ relative to the standard bases of the chain groups as an integer matrix M_k with entries in $\{-1, 0, 1\}$. The matrix M_k is called the *standard matrix representation* of ∂_k . It has m_k columns and m_{k-1} rows (the number of k - and $(k-1)$ -simplices, respectively.) The null-space of M_k corresponds to \mathbf{Z}_k and its range-space to \mathbf{B}_{k-1} , as manifested in Figure 3. The *reduction algorithm* derives alternate bases for the chain groups, relative to which the matrix for ∂_k is diagonal. The algorithm utilizes the following *elementary row operations* on M_k :

1. exchange row i and row j ,
2. multiply row i by -1 ,
3. replace row i by $(\text{row } i) + q(\text{row } j)$, where q is an integer and $j \neq i$.

The algorithm also uses *elementary column operations* that are similarly defined. Each column (row) operation corresponds to a change in the basis for \mathbf{C}_k (\mathbf{C}_{k-1}). For example, if e_i and e_j are the i th and j th basis elements for \mathbf{C}_k , respectively, a column operation of type (3) amounts to replacing e_i with $e_i + qe_j$. A similar row operation on basis elements \hat{e}_i and \hat{e}_j for \mathbf{C}_{k-1} , however, replaces \hat{e}_j by $\hat{e}_j - q\hat{e}_i$. We shall make use of this fact in Section 4. The algorithm systematically modifies the bases of \mathbf{C}_k and \mathbf{C}_{k-1} using elementary operations to reduce M_k to its (*Smith*) *normal form*:

$$\tilde{M}_k = \left[\begin{array}{ccc|cc} b_1 & & 0 & & \\ & \ddots & & & 0 \\ & & b_{l_k} & & \\ \hline & & 0 & & 0 \end{array} \right],$$

where $l_k = \text{rank } M_k = \text{rank } \tilde{M}_k$, $b^i \geq 1$, and $b_i | b_{i+1}$ for all $1 \leq i < l_k$. The algorithm can also compute corresponding bases $\{e_j\}$ and $\{\hat{e}_i\}$ for \mathbf{C}_k and \mathbf{C}_{k-1} , respectively, although this is unnecessary if a decomposition is all that is needed. Computing the normal form in all dimensions, we get a full characterization of \mathbf{H}_k :

- (i) the torsion coefficients of \mathbf{H}_{k-1} (d_i in (1)) are precisely the diagonal entries b_i greater than one.
- (ii) $\{e_i \mid l_k + 1 \leq i \leq m_k\}$ is a basis for \mathbf{Z}_k . Therefore, $\text{rank } \mathbf{Z}_k = m_k - l_k$.
- (iii) $\{b_i \hat{e}_i \mid 1 \leq i \leq l_k\}$ is a basis for \mathbf{B}_{k-1} . Equivalently, $\text{rank } \mathbf{B}_k = \text{rank } M_{k+1} = l_{k+1}$.

Combining (ii) and (iii), we have

$$\beta_k = \text{rank } \mathbf{Z}_k - \text{rank } \mathbf{B}_k = m_k - l_k - l_{k+1}. \quad (3)$$

For the complex in Figure 1, the standard matrix representation of ∂_1 is

$$M_1 = \begin{bmatrix} & ab & bc & cd & ad & ac \\ a & -1 & 0 & 0 & -1 & -1 \\ b & 1 & -1 & 0 & 0 & 0 \\ c & 0 & 1 & -1 & 0 & 1 \\ d & 0 & 0 & 1 & 1 & 0 \end{bmatrix},$$

where we show the bases within the matrix. Reducing the matrix, we get the normal form

$$\tilde{M}_1 = \begin{bmatrix} & cd & bc & ab & z_1 & z_2 \\ d-c & 1 & 0 & 0 & 0 & 0 \\ c-b & 0 & 1 & 0 & 0 & 0 \\ b-a & 0 & 0 & 1 & 0 & 0 \\ a & 0 & 0 & 0 & 0 & 0 \end{bmatrix},$$

where $z_1 = ad - bc - cd - ab$ and $z_2 = ac - bc - ab$ form a basis for \mathbf{Z}_1 and $\{d-c, c-b, b-a\}$ is a basis for \mathbf{B}_0 .

We may use a similar procedure to compute homology over graded PIDs. A *homogeneous basis* is a basis of homogeneous elements. We begin by representing ∂_k relative to the standard basis of \mathbf{C}_k (which is homogeneous) and a homogeneous basis for \mathbf{Z}_{k-1} . Reducing to normal form, we read off the description provided by direct sum (2) using the new basis $\{\hat{e}_j\}$ for \mathbf{Z}_{k-1} :

- (i) zero row i contributes a free term with shift $\alpha_i = \text{deg } \hat{e}_i$,
- (ii) row with diagonal term b_i contributes a torsional term with homogeneous $d_j = b_j$ and shift $\gamma_j = \text{deg } \hat{e}_j$.

The reduction algorithm requires $O(m^3)$ elementary operations, where m is the number of simplices in K . The operations, however, must be performed in exact integer arithmetic. This is problematic in practice, as the entries of the intermediate matrices may become extremely large.

2.6 Persistence

We end this section with by re-introducing persistence. Given a filtered complex, the i th complex K^i has associated boundary operators ∂_k^i , matrices M_k^i , and groups $\mathbf{C}_k^i, \mathbf{Z}_k^i, \mathbf{B}_k^i$ and \mathbf{H}_k^i for all $i, k \geq 0$. Note that superscripts indicate the filtration index and are not related to cohomology.

The p -persistent k th homology group of K^i is

$$\mathbf{H}_k^{i,p} = \mathbf{Z}_k^i / (\mathbf{B}_k^{i+p} \cap \mathbf{Z}_k^i). \quad (4)$$

The definition is well-defined: both groups in the denominator are subgroups of \mathbf{C}_k^{i+p} , so their intersection is also a group, a subgroup

of the numerator. The p -persistent k th Betti number of K^i is $\beta_k^{i,p}$, the rank of the free subgroup of $\mathbf{H}_k^{i,p}$. We may also define persistent homology groups using the injection $\eta_k^{i,p}: \mathbf{H}_k^i \rightarrow \mathbf{H}_k^{i+p}$, that maps a homology class into the one that contains it. Then, $\text{im } \eta_k^{i,p} \simeq \mathbf{H}_k^{i,p}$ [7]. We extend this definition over arbitrary PIDs, as before. Persistent homology groups are modules and Theorem 2.1 describes their structure.

3. THE PERSISTENCE MODULE

In this section, we take a different view of persistent homology in order to understand its structure. Intuitively, the computation of persistence requires compatible bases for \mathbf{H}_k^i and \mathbf{H}_k^{i+p} . It is not clear when a succinct description is available for the compatible bases. We begin this section by combining the homology of all the complexes in the filtration into a single algebraic structure. We then establish a correspondence that reveals a simple description over fields. We end this section by illustrating the relationship of our view to the persistence equation (Equation (4)).

Definition 3.1 (persistence complex) A *persistence complex* \mathcal{C} is a family of chain complexes $\{\mathbf{C}_*^i\}_{i \geq 0}$ over R , together with chain maps $f^i: \mathbf{C}_*^i \rightarrow \mathbf{C}_*^{i+1}$, so that we have the following diagram:

$$\mathbf{C}_*^0 \xrightarrow{f^0} \mathbf{C}_*^1 \xrightarrow{f^1} \mathbf{C}_*^2 \xrightarrow{f^2} \dots$$

Our filtered complex K with inclusion maps for the simplices becomes a persistence complex. Below, we show a portion of a persistence complex, with the chain complexes expanded. The filtration index increases horizontally to the right under the chain maps f^i , and the dimension decreases vertically to the bottom under the boundary operators ∂_k .

$$\begin{array}{ccccccc} \partial_3 \downarrow & & \partial_3 \downarrow & & \partial_3 \downarrow & & \\ \mathbf{C}_2^0 & \xrightarrow{f^0} & \mathbf{C}_2^1 & \xrightarrow{f^1} & \mathbf{C}_2^2 & \xrightarrow{f^2} & \dots \\ \partial_2 \downarrow & & \partial_2 \downarrow & & \partial_2 \downarrow & & \\ \mathbf{C}_1^0 & \xrightarrow{f^0} & \mathbf{C}_1^1 & \xrightarrow{f^1} & \mathbf{C}_1^2 & \xrightarrow{f^2} & \dots \\ \partial_1 \downarrow & & \partial_1 \downarrow & & \partial_1 \downarrow & & \\ \mathbf{C}_0^0 & \xrightarrow{f^0} & \mathbf{C}_0^1 & \xrightarrow{f^1} & \mathbf{C}_0^2 & \xrightarrow{f^2} & \dots \end{array}$$

Definition 3.2 (persistence module) A *persistence module* \mathcal{M} is a family of R -modules M^i , together with homomorphisms $\varphi^i: M^i \rightarrow M^{i+1}$.

For example, the homology of a persistence complex is a persistence module, where φ^i simply maps a homology class to the one that contains it.

Definition 3.3 (finite type) A persistence complex $\{\mathbf{C}_*^i, f^i\}$ (persistence module $\{M^i, \varphi^i\}$) is of *finite type* if each component complex (module) is a finitely generated R -module, and if the maps f^i (φ^i , respectively) are isomorphisms for $i \geq m$ for some integer m .

As our complex K is finite, it generates a persistence complex \mathcal{C} of finite type, whose homology is a persistence module \mathcal{M} of finite type. We showed in the introduction how such complexes arise in practice.

3.1 Correspondence

Suppose we have a persistence module $\mathcal{M} = \{M^i, \varphi^i\}_{i \geq 0}$ over ring R . We now equip $R[t]$ with the standard grading and define a graded module over $R[t]$ by

$$\alpha(\mathcal{M}) = \bigoplus_{i=0}^{\infty} M^i,$$

where the R -module structure is simply the sum of the structures on the individual components, and where the action of t is given by

$$t \cdot (m^0, m^1, m^2, \dots) = (0, \varphi^0(m^0), \varphi^1(m^1), \varphi^2(m^2), \dots).$$

That is, t simply shifts elements of the module up in the gradation.

Theorem 3.1 (correspondence) The correspondence α defines an equivalence of categories between the category of persistence modules of finite type over R and the category of finitely generated non-negatively graded modules over $R[t]$.

The proof is the Artin-Rees theory in commutative algebra [8].

The correspondence established by Theorem 3.1 shows that there exists no simple classification of persistence modules over a ground ring, such as \mathbb{Z} , that is not a field. It is well known in commutative algebra that the classification of modules over $\mathbb{Z}[t]$ is extremely complicated. While it is possible to assign interesting invariants to $\mathbb{Z}[t]$ -modules, a simple classification is not available, nor is it likely ever to be available.

On the other hand, the correspondence gives us a simple decomposition when the ground ring is a field F . Here, the graded ring $F[t]$ is a PID and its only graded ideals are homogeneous of form (t^n) , so the structure of the $F[t]$ -module is described by sum (2) in Theorem 2.1:

$$\left(\bigoplus_{i=1}^n \Sigma^{\alpha_i} F[t] \right) \oplus \left(\bigoplus_{j=1}^m \Sigma^{\gamma_j} F[t]/(t^{n_j}) \right). \quad (5)$$

We wish to parametrize the isomorphism classes of $F[t]$ -modules by suitable objects.

Definition 3.4 (\mathcal{P} -interval) A \mathcal{P} -interval is an ordered pair (i, j) with $0 \leq i < j \in \mathbb{Z}^\infty = \mathbb{Z} \cup \{+\infty\}$.

We associate a graded $F[t]$ -module to a set \mathcal{S} of \mathcal{P} -intervals via a bijection Q . We define $Q(i, j) = \Sigma^i F[t]/(t^{j-i})$ for \mathcal{P} -interval (i, j) . Of course, $Q(i, +\infty) = \Sigma^i F[t]$. For a set of \mathcal{P} -intervals $\mathcal{S} = \{(i_1, j_1), (i_2, j_2), \dots, (i_n, j_n)\}$, we define

$$Q(\mathcal{S}) = \bigoplus_{l=1}^n Q(i_l, j_l).$$

Our correspondence may now be restated as follows.

Corollary 3.1 The correspondence $\mathcal{S} \rightarrow Q(\mathcal{S})$ defines a bijection between the finite sets of \mathcal{P} -intervals and the finitely generated graded modules over the graded ring $F[t]$. Consequently, the isomorphism classes of persistence modules of finite type over F are in bijective correspondence with the finite sets of \mathcal{P} -intervals.

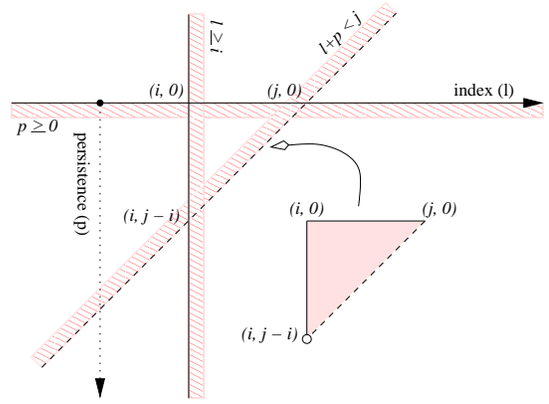


Figure 4: The inequalities $p \geq 0$, $l \geq i$, and $l + p < j$ define a triangular region in the index-persistence plane. This region defines when the cycle is a basis element for the homology vector space.

3.2 Interpretation

Before proceeding any further, we recap our work so far and relate it to prior results. Recall that our input is a filtered complex K and we are interested in its k th homology. In each dimension, the homology of complex K^i becomes a vector space over a field, described fully by its rank β_k^i . We need to choose compatible bases across the filtration in order to compute persistent homology for the entire filtration. So, we form the persistence module corresponding to K , a direct sum of these vector spaces. The structure theorem states that a basis exists for this module that provides compatible bases for all the vector spaces. In particular, each \mathcal{P} -interval (i, j) describes a basis element for the homology vector spaces starting at time i until time $j-1$. This element is a k -cycle e that is completed at time i , forming a new homology class. It also remains non-bounding until time j , at which time it joins the boundary group \mathbf{B}_k^j . Therefore, the \mathcal{P} -intervals discussed here are precisely the so-called k -intervals utilized in [7] to describe persistent \mathbb{Z}_2 -homology. That is, while component homology groups are torsion-less, persistence appears as torsional and free elements of the persistence module.

Our interpretation also allows us to ask when $e + \mathbf{B}_k^l$ is a basis element for the persistent groups $\mathbf{H}_k^{l,p}$. Recall Equation (4). As $e \notin \mathbf{B}_k^l$ for all $l < j$, we know that $e \notin \mathbf{B}_k^{l+p}$ for $l + p < j$. Along with $l \geq i$ and $p \geq 0$, the three inequalities define a triangular region in the index-persistence plane, as drawn in Figure 4. The region gives us the values for which the k -cycle e is a basis element for $\mathbf{H}_k^{l,p}$. In other words, we have just shown a direct proof of the k -triangle Lemma in [7], which we restate here in a different form.

Lemma 3.1 Let \mathcal{T} be the set of triangles defined by \mathcal{P} -intervals for the k -dimensional persistence module. The rank $\beta_k^{l,p}$ of $\mathbf{H}_k^{l,p}$ is the number of triangles in \mathcal{T} containing the point (l, p) .

Consequently, computing persistent homology over a field is equivalent to finding the corresponding set of \mathcal{P} -intervals.

4. ALGORITHM FOR FIELDS

In this section, we devise an algorithm for computing persistent homology over a field. Given the theoretical development of the last section, our approach is rather simple: we simplify the standard reduction algorithm using the properties of the persistence module.

Our arguments give an algorithm for computing the \mathcal{P} -intervals for a filtered complex directly over the field F , without the need for constructing the persistence module. This algorithm is a generalized version of the pairing algorithm shown in [7].

4.1 Derivation

We use the small filtration in Figure 1 as a running example and compute over \mathbb{Z}_2 , although any field will do. The persistence module corresponds to a $\mathbb{Z}_2[t]$ -module by the correspondence established in Theorem 2.1. Table 1 reviews the degrees of the simplices of our filtration as homogeneous elements of this module.

a	b	c	d	ab	bc	cd	ad	ac	abc	acd
0	0	1	1	1	1	2	2	3	4	5

Table 1: Degree of simplices of filtration in Figure 1

Throughout this section, we use $\{e_j\}$ and $\{\hat{e}_i\}$ to represent homogeneous bases for \mathbf{C}_k and \mathbf{C}_{k-1} , respectively. Relative to homogeneous bases, any representation M_k of ∂_k has the following basic property:

$$\deg \hat{e}_i + \deg M_k(i, j) = \deg e_j, \quad (6)$$

where $M_k(i, j)$ denotes the element at location (i, j) . We get

$$M_1 = \begin{bmatrix} & ab & bc & cd & ad & ac \\ d & 0 & 0 & t & t & 0 \\ c & 0 & 1 & t & 0 & t^2 \\ b & t & t & 0 & 0 & 0 \\ a & t & 0 & 0 & t^2 & t^3 \end{bmatrix}, \quad (7)$$

for ∂_1 in our example. The reader may verify Equation (6) using this example for intuition, e.g. $M_1(4, 4) = t^2$ as $\deg ad - \deg a = 2 - 0 = 2$, according to Table 1.

Clearly, the standard bases for chain groups are homogeneous. We need to represent $\partial_k: \mathbf{C}_k \rightarrow \mathbf{C}_{k-1}$ relative to the standard basis for \mathbf{C}_k and a homogeneous basis for \mathbf{Z}_{k-1} . We then reduce the matrix and read off the description of \mathbf{H}_k according to our discussion in Section 2.5. We compute these representations inductively in dimension. The base case is trivial. As $\partial_0 \equiv 0$, $\mathbf{Z}_0 = \mathbf{C}_0$ and the standard basis may be used for representing ∂_1 . Now, assume we have a matrix representation M_k of ∂_k relative to the standard basis $\{e_j\}$ for \mathbf{C}_k and a homogeneous basis $\{\hat{e}_i\}$ for \mathbf{Z}_{k-1} . For induction, we need to compute a homogeneous basis for \mathbf{Z}_k and represent ∂_{k+1} relative to \mathbf{C}_{k+1} and the computed basis. We begin by sorting basis \hat{e}_i in reverse degree order, as already done in the matrix in Equation (7). We next transform M_k into the *column-echelon form* \tilde{M}_k , a lower staircase form shown in Figure 5 [13]. The steps have variable height, all landings have width equal to one, and non-zero elements may only occur beneath the staircase. A boxed value in the figure is a *pivot* and a row (column) with a pivot is called a *pivot row (column)*. From linear algebra, we know that $\text{rank } M_k = \text{rank } \mathbf{B}_{k-1}$ is the number of pivots in an echelon form. The basis elements corresponding to non-pivot columns form the desired basis for \mathbf{Z}_k . In our example, we have

$$\tilde{M}_1 = \begin{bmatrix} & cd & bc & ab & z_1 & z_2 \\ d & \boxed{t} & 0 & 0 & 0 & 0 \\ c & t & \boxed{1} & 0 & 0 & 0 \\ b & 0 & t & \boxed{t} & 0 & 0 \\ a & 0 & 0 & t & 0 & 0 \end{bmatrix}, \quad (8)$$

$$\begin{bmatrix} \boxed{*} & 0 & & & & 0 \\ & \boxed{*} & 0 & \cdots & & \\ * & * & 0 & & & \vdots \\ & * & \boxed{*} & 0 & \cdots & \\ * & & & * & 0 & \cdots & 0 \end{bmatrix}$$

Figure 5: The column-echelon form. An * indicates a non-zero value and pivots are boxed.

where $z_1 = ad - cd - t \cdot bc - t \cdot ab$, and $z_2 = ac - t^2 \cdot bc - t^2 \cdot ab$ form a homogeneous basis for \mathbf{Z}_1 .

The procedure that arrives at the echelon form is Gaussian elimination on the columns, utilizing elementary column operations of types (1, 3) only. Starting with the left-most column, we eliminate non-zero entries occurring in pivot rows in order of increasing row. To eliminate an entry, we use an elementary column operation of type (3) that maintains the homogeneity of the basis and matrix elements. We continue until we either arrive at a zero column, or we find a new pivot. If needed, we then perform a column exchange (type (1)) to reorder the columns appropriately.

Lemma 4.1 (Echelon Form) The pivots in column-echelon form are the same as the diagonal elements in normal form. Moreover, the degree of the basis elements on pivot rows is the same in both forms.

PROOF. Because of our sort, the degree of row basis elements \hat{e}_i is monotonically decreasing from the top row down. Within each fixed column j , $\deg e_j$ is a constant c . By Equation (6), $\deg M_k(i, j) = c - \deg \hat{e}_i$. Therefore, the degree of the elements in each column is monotonically increasing with row. We may eliminate non-zero elements below pivots using row operations that do not change the pivot elements or the degrees of the row basis elements. We then place the matrix in diagonal form with row and column swaps. \square

The lemma states that if we are only interested in the degree of the basis elements, we may read them off from the echelon form directly. That is, we may use the following corollary of the standard structure theorem to obtain the description.

Corollary 4.1 Let \tilde{M}_k be the column-echelon form for ∂_k relative to bases $\{e_j\}$ and $\{\hat{e}_i\}$ for \mathbf{C}_k and \mathbf{Z}_{k-1} , respectively. If row i has pivot $\tilde{M}_k(i, j) = t^n$, it contributes $\Sigma^{\deg \hat{e}_i} F[t]/t^n$ to the description of \mathbf{H}_{k-1} . Otherwise, it contributes $\Sigma^{\deg \hat{e}_i} F[t]$. Equivalently, we get $(\deg \hat{e}_i, \deg \hat{e}_i + n)$ and $(\deg \hat{e}_i, \infty)$, respectively, as \mathcal{P} -intervals for \mathbf{H}_{k-1} .

In our example, $\tilde{M}_1(1, 1) = t$ in Equation (8). As $\deg d = 1$, the element contributes $\Sigma^1 \mathbb{Z}_2[t]/(t)$ or \mathcal{P} -interval (1,2) to the description of \mathbf{H}_0 .

We now wish to represent ∂_{k+1} in terms of the basis we computed for \mathbf{Z}_k . We begin with the standard matrix representation M_{k+1} of ∂_{k+1} . As $\partial_k \partial_{k+1} = \emptyset$, $M_k M_{k+1} = 0$, as shown in Figure 6. Furthermore, this relationship is unchanged by elementary operations. Since the domain of ∂_k is the codomain of ∂_{k+1} , the elementary column operations we used to transform M_k into echelon form \tilde{M}_k give corresponding row operations on M_{k+1} . These row operations zero out rows in M_{k+1} that correspond to non-zero pivot columns in \tilde{M}_k , and give a representation of ∂_{k+1} relative to the basis we just computed for \mathbf{Z}_k . This is precisely what we are after. We can get it, however, with hardly any work.

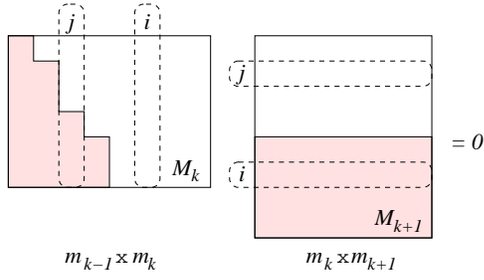


Figure 6: As $\partial_k \partial_{k+1} = \emptyset$, $M_k M_{k+1} = 0$ and this is unchanged by elementary operations. When M_k is reduced to echelon form \tilde{M}_k by column operations, the corresponding row operations zero out rows in M_{k+1} that correspond to pivot columns in \tilde{M}_k .

Lemma 4.2 (Basis Change) To represent ∂_{k+1} relative to the standard basis for \mathbf{C}_{k+1} and the basis computed for \mathbf{Z}_k , simply delete rows in M_{k+1} that correspond to pivot columns in \tilde{M}_k .

PROOF. We only used elementary column operations of types (1,3) in our variant of Gaussian elimination. Only the latter changes values in the matrix. Suppose we replace column i by (column i) + q (column j) in order to eliminate an element in a pivot row j , as shown in Figure 6. This operation amounts to replacing column basis element e_i by $e_i + qe_j$ in M_k . To effect the same replacement in the row basis for ∂_{k+1} , we need to replace row j with (row j) - q (row i). But row j is eventually zeroed-out, as shown in Figure 6, and rows i is never changed by any such operation. \square

Therefore, we have no need for row operations. We simply eliminate rows corresponding to pivot columns one dimension lower to get the desired representation for ∂_{k+1} in terms of the basis for \mathbf{Z}_k . This completes the induction. In our example, the standard matrix representation for ∂_2 is

$$M_2 = \left[\begin{array}{c|cc} & abc & acd \\ \hline ac & t & t^2 \\ ad & 0 & t^3 \\ cd & 0 & t^3 \\ bc & t^3 & 0 \\ ab & t^3 & 0 \end{array} \right].$$

To get a representation in terms of \mathbf{C}_2 and the basis (z_1, z_2) for \mathbf{Z}_1 we computed earlier, we simply eliminate the bottom three rows. These rows are associated with pivots in \tilde{M}_1 , according to Equation (8). We get

$$\tilde{M}_2 = \left[\begin{array}{c|cc} & abc & acd \\ \hline z_2 & t & t^2 \\ z_1 & 0 & t^3 \end{array} \right],$$

where we have also replaced ad and ac with the corresponding basis elements $z_1 = ad - bc - cd - ab$ and $z_2 = ac - bc - ab$.

4.2 Algorithm

Our discussion gives us an algorithm for computing \mathcal{P} -intervals of an $F[t]$ -module over field F . It turns out, however, that we can simulate the algorithm over the field itself, without the need for computing the $F[t]$ -module. Rather, we use two significant observations from the derivation of the algorithm. First, Lemma 4.1 guarantees that if we eliminate pivots in the order of decreasing degree, we may read off the entire description from the echelon form

and do not need to reduce to normal form. And second, Lemma 4.2 tells us that by simply noting the pivot columns in each dimension and eliminating the corresponding rows in the next dimension, we get the required basis change.

Therefore, we only need column operations throughout our procedure and there is no need for a matrix representation. We represent the boundary operators as a set of boundary chains corresponding to the columns of the matrix. Within this representation, column exchanges (type 1) have no meaning, and the only operation we need is of type 3. Our data structure is an array T with a slot for each simplex in the filtration, as shown in Figure 7 for our example. Each simplex gets a slot in the table. For indexing,

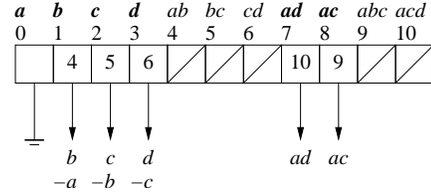


Figure 7: Data structure after running the algorithm on the filtration in Figure 1. Marked simplices are in bold italic.

we need a full ordering of the simplices, so we complete the partial order defined by the degree of a simplex by sorting simplices according to dimension, breaking all remaining ties arbitrarily (we did this implicitly in the matrix representation.) We also need the ability to mark simplices to indicate non-pivot columns.

Rather than computing homology in each dimension independently, we compute homology in all dimensions incrementally and concurrently. The algorithm, as shown in Figure 8, stores the list of \mathcal{P} -intervals for \mathbf{H}_k in L_k .

```

COMPUTEINTERVALS (K) {
  for k = 0 to dim(K) L_k = ∅;
  for j = 0 to m - 1 {
    d = REMOVEPIVOTROWS (σj);
    if (d = ∅) Mark σj;
    else {
      i = maxindex d; k = dim σj;
      Store j and d in T[i];
      L_k = L_k ∪ {(deg σi, deg σj)}
    }
  }
  for j = 0 to m - 1 {
    if σj is marked and T[j] is empty {
      k = dim σj; L_k = L_k ∪ {(deg σj, ∞)}
    }
  }
}

```

Figure 8: Algorithm COMPUTEINTERVALS processes a complex of m simplices. It stores the sets of \mathcal{P} -intervals in dimension k in L_k .

When simplex σ^j is added, we check via procedure REMOVEPIVOTROWS to see whether its boundary chain d corresponds to a zero or pivot column. If the chain is empty, it corresponds to a zero column and we mark σ^j : its column is a basis element for \mathbf{Z}_k , and the corresponding row should not be eliminated in the next dimension.

```

chain REMOVEPIVOTROWS ( $\sigma$ ) {
   $k = \dim \sigma$ ;  $d = \partial_k \sigma$ ;
  Remove unmarked terms in  $d$ ;
  while ( $d \neq \emptyset$ ) {
     $i = \text{maxindex } d$ ;
    if  $T[i]$  is empty, break;
    Let  $q$  be the coefficient of  $\sigma^i$  in  $T[i]$ ;
     $d = d - q^{-1}T[i]$ ;
  }
  return  $d$ ;
}

```

Figure 9: Algorithm REMOVEPIVOTROWS first eliminates rows not marked (not corresponding to the basis for Z_{k-1}), and then eliminates terms in pivot rows.

Otherwise, the chain corresponds to a pivot column and the term with the maximum index $i = \text{maxindex } d$ is the pivot, according to the procedure described for the $F[t]$ -module. We store index j and chain d representing the column in $T[i]$. Applying Corollary 4.1, we get \mathcal{P} -interval $(\deg \sigma^i, \deg \sigma^j)$. We continue until we exhaust the filtration. We then perform another pass through the filtration in search of infinite \mathcal{P} -intervals: marked simplices whose slot is empty.

We give the function REMOVEPIVOTROWS in Figure 9. Initially, the function computes the boundary chain d for the simplex. It then applies Lemma 4.2, eliminating all terms involving unmarked simplices to get a representation in terms of the basis for Z_{k-1} . The rest of the procedure is Gaussian elimination in the order of decreasing degree, as dictated by our discussion for the $F[t]$ -module. The term with the maximum index $i = \max d$ is a potential pivot. If $T[i]$ is non-empty, a pivot already exists in that row, and we use the inverse of its coefficient to eliminate the row from our chain. Otherwise, we have found a pivot and our chain is a pivot column. For our example filtration in Figure 7, the marked 0-simplices $\{a, b, c, d\}$ and 1-simplices $\{ad, ac\}$ generate \mathcal{P} -intervals $L_0 = \{(0, \infty), (0, 1), (1, 1), (1, 2)\}$ and $L_1 = \{(2, 4), (3, 5)\}$, respectively.

4.3 Discussion

From our derivation, it is clear that the algorithm has the same running time as Gaussian elimination over fields. That is, it takes $O(m^3)$ in the worst-case, where m is the number of simplices in the filtration. The algorithm is very simple, however, and represents the matrices efficiently. In our preliminary experiments, we have seen a linear time behavior for the algorithm.

5. ALGORITHM FOR PIDS

The correspondence we established in Section 3 eliminated any hope for a simple classification of persistent groups over rings that are not fields. Nevertheless, we may still be interested in their computation. In this section, we give an algorithm to compute the persistent homology groups $H_k^{i,p}$ of a filtered complex K for a fixed i and p . The algorithm we provide computes persistent homology over any PID D of coefficients by utilizing a reduction algorithm over that ring.

To compute the persistent group, we need to obtain a description of the numerator and denominator of the quotient group in Equation (4). We already know how to characterize the numerator. We

simply reduce the standard matrix representation M_k^i of ∂_k^i using the reduction algorithm. The denominator, $\mathbf{B}_k^{i,p} = \mathbf{B}_k^{i+p} \cap \mathbf{Z}_k^i$, plays the role of the boundary group in Equation (4). Therefore, instead of reducing matrix M_{k+1}^i , we need to reduce an alternate matrix $M_{k+1}^{i,p}$ that describes this boundary group. We obtain this matrix as follows:

- (1) We reduce matrix M_k^i to its normal form and obtain a basis $\{z^j\}$ for \mathbf{Z}_k^i , using fact (ii) in Section 2.5. We may merge this computation with that of the numerator.
- (2) We reduce matrix M_{k+1}^{i+p} to its normal form and obtain a basis $\{b^l\}$ for \mathbf{B}_k^{i+p} using fact (iii) in Section 2.5.
- (3) Let $N = [\{b^l\} \{z^j\}] = [B \ Z]$, that is, the columns of matrix N consist of the basis elements from the bases we just computed, and B and Z are the respective submatrices defined by the bases. We next reduce N to normal form to find a basis $\{u^q\}$ for its null-space. As before, we obtain this basis using fact (ii). Each $u^q = [\alpha^q \ \zeta^q]$, where α^q, ζ^q are vectors of coefficients of $\{b^l\}, \{z^j\}$, respectively. Note that $Nu^q = B\alpha^q + Z\zeta^q = 0$ by definition. In other words, element $B\alpha^q = -Z\zeta^q$ is belongs to the span of both bases. Therefore, both $\{B\alpha^q\}$ and $\{Z\zeta^q\}$ are bases for $\mathbf{B}_k^{i,p} = \mathbf{B}_k^{i+p} \cap \mathbf{Z}_k^i$. We form a matrix $M_{k+1}^{i,p}$ from either.

We now reduce $M_{k+1}^{i,p}$ to normal form and read off the torsion coefficients and the rank of $\mathbf{B}_k^{i,p}$. It is clear from the procedure that we are computing the persistent groups correctly, giving us the following.

Theorem 5.1 For coefficients in any PID, persistent homology groups are computable in the order of time and space of computing homology groups.

6. EXPERIMENTS

In this section, we discuss experiments using an implementation of the persistence algorithm for arbitrary fields. Our aim is to further elucidate the contributions of this paper. We look at two scenarios where the previous algorithm would not be applicable, but where our algorithm succeeds in providing information about a topological space.

6.1 Implementation

We have implemented our field algorithm for \mathbb{Z}_p for p a prime, and \mathbb{Q} coefficients. Our implementation is in C and utilizes GNU MP, a multi-precision library, for exact computation [10]. We have a separate implementation for coefficients in \mathbb{Z}_2 as the computation is greatly simplified in this field. The coefficients are either 0, or 1, so there is no need for orienting simplices or maintaining coefficients. A k -chain is simply a list of simplices, those with coefficient 1. Each simplex is its own inverse, reducing the group operation to the *symmetric difference*, where the sum of two k -chains c, d is $c + d = (c \cup d) - (c \cap d)$. We use a 2.2 GHz Pentium 4 Dell PC with 1 GB RAM running Red Hat Linux 7.3 for computing the timings.

6.2 Data

Our algorithm requires a persistence complex as input. In the introduction, we discussed how persistence complexes arise naturally in practice. In Example 1.3, we discussed generating persistence complexes using excursion sets of Morse functions over

	number s_k of k -simplices					χ
	0	1	2	3	4	
K	2,000	6,000	4,000	0	0	0
E	3,095	52,285	177,067	212,327	84,451	1
J	17,862	297,372	1,010,203	1,217,319	486,627	1

Table 2: Datasets. K is the Klein bottle, E is potential around electrostatic charges. J is supersonic jet flow.

manifolds. We have implemented a general framework for computing complexes of this type. We must emphasize, however, that our persistence software processes persistence complexes of any origin.

Our framework takes a tuple (K, f) as input and produces a persistence complex $\mathcal{C}(K, f)$ as output. K is a d -dimensional simplicial complex that triangulates an underlying manifold. And $f: \text{vert } K \rightarrow \mathbb{R}$ is a discrete function over the vertices of K that we extend linearly over the remaining simplices of K . The function f acts as the Morse function over the manifold, but need not be Morse for our purposes. Frequently, our complex is augmented with a map $\varphi: K \rightarrow \mathbb{R}^d$ that immerses or embeds the manifold in Euclidean space. Our algorithm does not require φ for computation, but φ is often provided as a discrete map over the vertices of K and is extended linearly as before. For each dataset, Table 2 gives the number s_k of k -simplices, as well as the Euler characteristic $\chi = \sum_k (-1)^k s_k$. We use the Morse function to compute the excursion set filtration for each dataset. Table 3 gives information on the resulting filtrations.

	$ K $	len	filt (s)	pers (s)
K	12,000	1,020	0.03	< 0.01
E	529,225	3,013	3.17	5.00
J	3,029,383	256	24.13	50.23

Table 3: Filtrations. The number of simplices in the filtration $|K| = \sum_i s_i$, the length of the filtration (number of distinct values of function f), time to compute the filtration, and time to compute persistence over \mathbb{Z}_2 coefficients.

6.3 Field Coefficients

A contribution of this paper is the generalization of the persistence algorithm to arbitrary fields. This contribution is important when the manifold under study contains torsion. To make this clear, we compute the homology of the Klein bottle using the persistence algorithm. Here, we are interested only in the Betti numbers of the final complex in the filtration for illustrative purposes. In homology, the non-orientability of the Klein bottle manifests itself as a torsional 1-cycle c where $2c$ is a boundary (indeed, it bounds the surface itself.) The homology groups over \mathbb{Z} are: $H_0(K) = \mathbb{Z}$, $H_1(K) = \mathbb{Z} \times \mathbb{Z}_2$, and $H_2(K) = \{0\}$. Note that $\beta_1 = \text{rank } H_1 = 1$. We now use the “height function” as our Morse function, $f = z$, to generate the filtration in Table 3. We then compute the homology of dataset K with field coefficients using our algorithm, as shown in Table 4.

Over \mathbb{Z}_2 , we get $\beta_1 = 2$ as homology is unable to recognize the torsional boundary $2c$ with coefficients 0 and 1. Instead, it observes an additional class of homology 1-cycles. By the Euler-Poincaré relation, $\chi = \sum_i \beta_i$, so we also get a class of 2-cycles to compensate for the increase in β_1 [12]. Therefore, \mathbb{Z}_2 -homology misidentifies

F	β_0	β_1	β_2	time (s)
\mathbb{Z}_2	1	2	1	0.01
\mathbb{Z}_3	1	1	0	0.23
\mathbb{Z}_5	1	1	0	0.23
\mathbb{Z}_{3203}	1	1	0	0.23
\mathbb{Q}	1	1	0	0.50

Table 4: Field coefficients. The Betti numbers of K computed over field F and time for the persistence algorithm. We use a separate implementation for \mathbb{Z}_2 coefficients.

the Klein bottle as the torus. Over any other field, however, homology turns the torsional cycle into a boundary, as the inverse of 2 exists. In other words, while we cannot observe torsion in computing homology over fields, we can deduce its existence by comparing our results over different coefficient sets. Similarly, we can compare sets of \mathcal{P} -intervals from different computations to discover torsion in a persistence complex.

Note that our algorithm’s performance for this dataset is about the same over arbitrary finite fields, as the coefficients do not get large. The computation over \mathbb{Q} takes about twice as much time and space, since each rational is represented as two integers in GNU MP.

6.4 Higher Dimensions

A second contribution of this paper is the extension of the persistence algorithm from subcomplexes of \mathbb{S}^3 to complexes in arbitrary dimensions. We have already utilized this capability in computing the homology of the Klein bottle. We now examine the performance of this algorithm in higher dimensions. For practical motivation, we use large-scale time-varying volume data as input. Advances in data acquisition systems and computing technologies have resulted in the generation of massive sets of measured or simulated data. The datasets usually contain the time evolution of physical variables, such as temperature, pressure, or flow velocity at sample points in space. The goal is to identify and localize significant phenomena within the data. We propose using persistence as the significance measure.

The underlying space for our datasets is the four-dimensional space-time manifold. For each dataset, we triangulate the convex hull of the samples to get a triangulation. Each complex listed in Table 2 is homeomorphic to a four-dimensional ball and has $\chi = 1$. Dataset E contains the potential around electrostatic charges at each vertex. Dataset J records the supersonic flow velocity of a jet engine. We use these values as Morse functions to generate the filtrations. We then compute persistence over \mathbb{Z}_2 coefficients to get the Betti numbers. We give filtration sizes and timings in Table 3. Figure 10 displays β_2 for dataset J. We observe large number of two-dimensional cycles (voids), as the co-dimension is 2. Persistence allows us to do to decompose this graph into the set of \mathcal{P} -intervals. Although there are 730,692 \mathcal{P} -intervals in dimension two, most are empty as the topological attribute is created and destroyed at the same function level. We draw the 502 non-empty \mathcal{P} -intervals in Figure 11. We note that the \mathcal{P} -intervals represent a compact and general *shape descriptor* for arbitrary spaces.

For the large data sets, we do not compute persistence over alternate fields as the computation requires in excess of two gigabytes of memory. In the case of finite fields \mathbb{Z}_p , we may restrict the prime p to be so that the computation fits within an integer. This is a reasonable restriction, as on most modern machines with 32-bit integers, it implies $p < 2^{16} - 1$. Given this restriction, any coefficient will

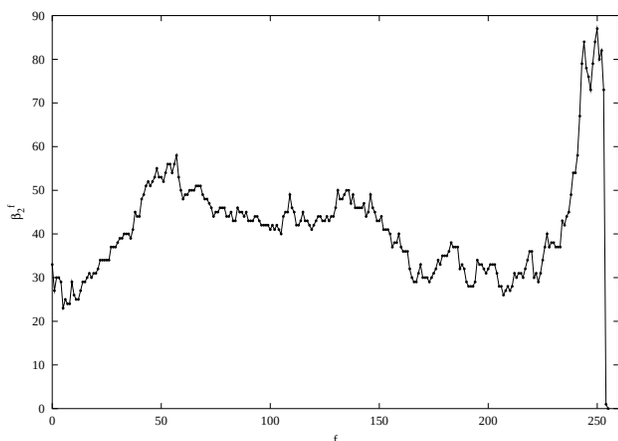


Figure 10: Graph of β_1^f for dataset J, where f is the flow velocity.

be less than p and representable as a 4-byte integer. The GNU MP exact integer format, on the other hand, requires at least 16 bytes for representing any integer.

7. CONCLUSION

We believe the most important contribution of this paper is a reinterpretation of persistent homology within the classical framework of algebraic topology. Our interpretation allows us to:

1. establish a correspondence that fully describes the structure of persistent homology over any field, not only over \mathbb{Z}_2 , as in the previous result,
2. and relate the previous algorithm to the classic reduction algorithm, thereby extending it to arbitrary fields and arbitrary dimensional complexes, not just subcomplexes of \mathbb{S}^3 as in the previous result.

We provide implementations of our algorithm for fields, and show that they perform quite well for large datasets. Finally, we give an algorithm for computing a persistent homology group with fixed parameters over arbitrary PIDs.

Our software for n -dimensional complexes enables us to analyze arbitrary-dimensional point cloud data and their derived spaces. One current project uses this implementation for feature recognition using a novel algebraic method [2]. Another project applies persistence to derived spaces to arrive at compact shape descriptors for geometric objects [3]. Future theoretical work include examining invariants for persistent homology over non-fields and defining *multivariate persistence*, where there is more than one persistence dimension. An example would be tracking a Morse function as well as density of sampling on a manifold.

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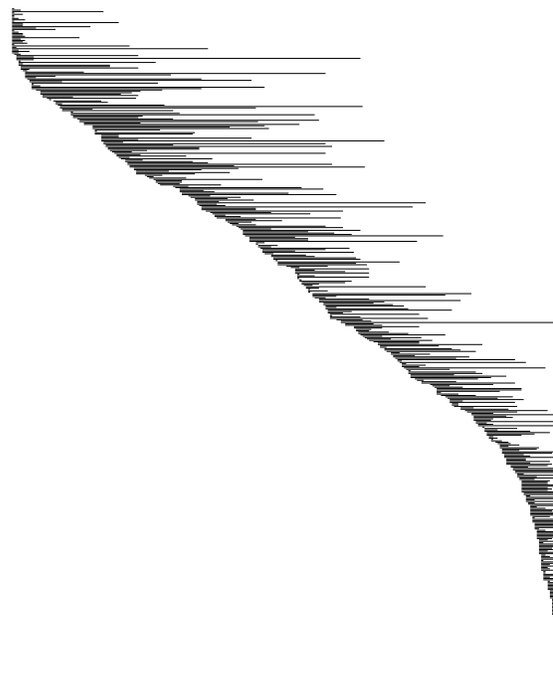


Figure 11: The 502 non-empty \mathcal{P} -intervals for dataset J in dimension two. The amalgamation of these intervals gives the graph in Figure 10.

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