Persistent Homology Based Characterization of the Breast Cancer Immune Microenvironment: A Feasibility Study

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- Abstract -

Persistent homology is a common tool of topological data analysis, whose main descriptor, the persistence diagram, aims at computing and encoding the geometry and topology of given datasets. In this article, we present a novel application of persistent homology to characterize the spatial arrangement of immune and epithelial (tumor) cells within the breast cancer immune microenvironment. More specifically, quantitative and robust characterizations are built by computing persistence diagrams out of a staining technique (quantitative multiplex immunofluorescence) which allows us to obtain spatial coordinates and stain intensities on individual cells. The resulting persistence diagrams are evaluated as characteristic biomarkers of cancer subtype and prognostic biomarker of overall survival. For a cohort of approximately 700 breast cancer patients with median 8.5-year clinical follow-up, we show that these persistence diagrams outperform and complement the usual descriptors which capture spatial relationships with nearest neighbor analysis. This provides new insights and possibilities on the general problem of building (topology-based) biomarkers that are characteristic and predictive of cancer subtype, overall survival and response to therapy.

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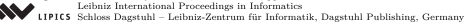
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1 Introduction

Descriptors computed with topological data analysis (TDA), such as persistence diagrams [21, 60] and Mapper [56], have shown strong analytical power in many real world biological data. Examples include (but are not limited to) neuronal structures [41, 33], cardiac trabeculae [24, 59], brain images [46, 39] and genomics data [44, 12, 51]. These methods capture multiscale geometric and structural patterns of the data with guaranteed robustness against potential noise introduced in measurement [17, 18] and in upstream preprocessing steps [7]. As such, they provide a systematic way to quantify complex biomedical systems. Furthermore, state-of-the-art discriminative models (i.e., classifiers) [11, 30, 34] and unsupervised models (i.e., clustering methods) [36] have been recently introduced, and are able to effectively connect topological features and clinical/biological outcomes of interest.

In this paper, we present a new application of topological data analysis, namely, the characterization of breast cancer immune microenvironment using persistence diagrams. Despite tremendous advancements in cancer screening, diagnostic methods and treatment, breast cancer remains the second leading cause of cancer death in women with projections of 270,000 new cases and approximately 42,000 deaths from invasive breast cancer in 2019 [53]. Therefore, identifying descriptors that indicate potential therapeutic targets and predict outcome is a critical yet unmet need in breast cancer [20]. The goal of this article is to show how persistence diagrams can help in fulfilling this task.

Cancer research and characterization of spatial cell arrangement. In the past decade, a major focus of cancer research has been on the interplay between the tumor and the immune environment, referred to as the tumor immune microenvironment [8]. By characterizing host-specific functional anti-tumor immune responses and their correlation to cancer subtype and overall survival, patient specific immunotherapeutic targets can be identified [49] with higher precision. To achieve the goal, it is necessary to characterize the complex spatial arrangement between cancer cells and a mixture of different immune cells, e.g., T-cells and macrophages, both of which play a versatile biological role and are believed to be crucially relevant to initiation and regulation of the immune response. This task involves two important steps: cell detection and characterization.

Thanks to the rapid development of imaging technology and deep learning methods, we are able to detect not only locations, but also types of different cells within a slide of tumor biopsy sample from a cancer patient. By staining the slide using immunohistochemical (IHC) markers, we are able to tag different types of cells with different stains, i.e., colors bounded with different protein biomarkers. Using a brightfield image scanner, we convert the stained slide into a whole slide image in which various cells can be identified by their respective stains [47, 32]. The identification of cells is referred to as *phenotyping*. Advanced deep learning methods [23, 1] have been developed to unmix the stains and to detect cells and their types. This approach, called multiplex IHC, is scalable but less precise as noise is introduced due to the additional deep learning cell detector. Alternatively, we may use quantitative multiplex immunofluorescence (qmIF), which stains different cells with different fluorescent stains and detect them using lenses with specific filters. The qmIF approach is highly reliable, albeit costly in material and in time.

Once cells of different types are detected, we need to quantitatively characterize their spatial arrangements in order to evaluate correlations with various outcomes of interest. There are two major challenges. First, the spatial arrangement is highly heterogeneous across different patients and even within a single tissue sample. Second, stain intensity is

relative, and phenotype thresholds must be manually determined. Discerning true signal from background isn't always clear, and currently is done in relation to other tissue samples. Nonetheless, qmIF imaging provides rich data for study; see Figure 1 for an example of the raw image data.

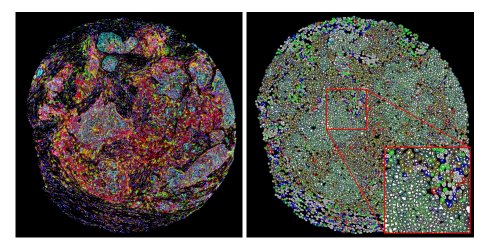


Figure 1 An example input data. Left: The raw microscopic image of a stained tissue sample. The sample is approximately 1x1 mm² large. The image is 2,000x2,000 pixels, 0.5x0.5 micron² per pixel. A sample usually contains 3,000 to 5,000 cells. Right: The processed results. Cells are identified by localizing their nuclei with a special stain (shown as white regions). The phenotype of each cell can be identified by the stain intensity of its cytoplasm and nucleus: T cells are tagged with CD8 (blue), macrophages are tagged with CD68 (green), tumor cells are tagged with pancytokeratin (cyan). Any cell may additionally be tagged with PD-L1 (red). The cells are abstracted into point clouds with different stain intensities, as shown in Figure 3.

Related work. Previous methods [25, 54] focus on using nearest neighbor distances from cells of one type (obtained by thresholding the stain intensities) to cells of a second type. Unfortunately, this approach is sensitive to noise and lacks the ability to model stain concentration variations due to the thresholding. Moreover, it can only characterize fixed neighborhoods around the cells and is oblivious to larger cell arrangements.

Persistent homology has recently been used to characterize cellular architecture in pathology images in [37], where these descriptors were shown to successfully detect and quantify circular cell structures corresponding to glands. In contrast, our work operates on coordinates of phenotyped cells and deals with the global characterization of complex interactions between these cellular phenotypes.

Contributions. In this article, we propose the first topological analysis of tumor immune microenvironment. More specifically, we provide empirical evidence that persistence diagrams are suitable descriptors by experimentally demonstrating the following points:

First, stain concentration levels, or stain intensities, that are usually used by practitioners to filter cells, are natural candidates for defining *filtrations* (in the TDA vocabulary) from which persistence diagrams can be computed. This way, the whole range of stain intensities is taken into account instead of thresholding. We hypothesize that the stain intensity is biologically meaningful and the resulting persistence diagrams will be more predictive than just using cell coordinates from thresholding.

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Second, persistence diagrams are able to capture topological and structural features that are characteristic of the arrangement of the cells. This is because the structures encoded by persistence diagrams are robust to spatial deformation and other types of noise introduced in detection, which prevents the analysis from being biased by measurement errors, contrarily to other descriptors used in the literature.

Our study, although preliminary, demonstrates the potential of persistence homology being a novel tool to characterize the tumor immune microenvironment. With rich computation and learning tools available for persistence-derived features, we are confident that topological characterization will lead to powerful diagnostic and prognostic cancer biomarkers.

Plan of the article. We introduce our biological data, and briefly recall the basics of topological data analysis in Section 2. Then, we explain our methods for computing and running statistical tests on persistence diagrams in Section 3. Finally, we conclude and summarize future investigations and open questions in Section 4.

2 Data and Background

In this section, we introduce our biological data (Section 2.1), and briefly recall the rationale for nearest neighbor analysis (Section 2.2) and topological data analysis (Section 2.3).

2.1 Biological Data

We analyze a large cohort of patients with extensive 8.5 years of follow-up. For each tissue sample, qmIF imaging was obtained with a panel of immune markers for phenotyping the tumor immune microenvironment, including: CD8 (T-cells), CD68 (macrophages) and pancytokeratin (cancer cells). Then, a commercial software package (HALO, Indica Labs) was used to perform nuclear segmentation, cytoplasmic definition, and stain quantification. Cell phenotypes, based on a threshold applied to the stain intensity, were defined manually. See Figures 1 for the conventional threshold-based phenotype analysis. Let us now provide details on the important steps that were necessary to collect our data.

Patient Cohort. Our raw data is comprised of high-throughput tissue microarrays (TMA) consisting of 1mm × 1mm cores of tissue. The TMA were assembled with tissues from a cohort of 900 patients that underwent tumor resection following a diagnosis of breast cancer at Pitt County Memorial Hospital (now Vidant Hospital) in Greenville, North Carolina. Patient samples and clinicopathological data were collected under an IRB approved protocol at the Brody School of Medicine, East Carolina University [9]. The cohort is uniquely valuable for research as there is median 8.5 year follow-up data which allows for in depth evaluation for topological biomarkers with patient attributes and clinical outcomes.

Quantitative Multiplex Immunofluorescence. Unlike traditional immunohistochemistry, qmIF enables simultaneous staining of multiple markers in a single piece of tissue. We use the Ultivue UltiMapper I/O PD-L1 assay consisting of the following markers: CD8 (cytotoxic T-cells), CD68 (macrophages), PD-L1 (an immune suppressive protein), pancytokeratin (epithelial cells), and DAPI (DNA marker) for identification of cell nuclei. In our data, positively stained epithelial cells via pancytokeratin are considered to be tumor cells. Every cell in the tissue is designated with a PD-L1 status being either positive or negative corresponding to above or below threshold stain intensity. All staining thresholds are adaptively determined to

enhance signal (consistent with a positive staining pattern assessed visually) to background. The result of the phenotyping analysis is a text file for each tissue sample consisting of entries listing information about each cell location, including the manual phenotyping result and raw stain intensities. Each tissue sample consists of 3,000-5,000 cells.

2.2 Nearest Neighbor Analysis

Nearest neighbor analysis is commonly performed with qmIF data [25]. We perform a nearest neighbor search between combinations of phenotypes for all possible phenotype pairs. More specifically, for a given pair of phenotypes P_1, P_2 , each composed of cells (detected with thresholds on their stain intensities) with two coordinates, we compute, for a given cell C_i belonging to P_1 , the Euclidean distances to all cells belonging to P_2 , excluding those whose distance is less than 0.05 microns to prevent cell-overlap. We keep the minimum distance value among those, which we call the nearest neighbor distance, and repeat this process for each cell in P_1 to form a distribution of nearest neighbor distances, d_i . The mean and standard deviation of d_i are then derived. We apply the same process for all pairs of phenotypes and used the corresponding means and deviations as features of biomarkers, potentially predictive of triple-negative status and prognostic of overall survival. This can also be written as a function of matrix operations involving the similarity matrix of cell coordinates between P_1 and P_2 :

$$\mathbf{q}_{ik} = \{C_{1k}, C_{2k}, ..., C_{ik}\}, \ \mathbf{t}_{kj} = \{C_{1k}, C_{2k}, ..., C_{jk}\}^{\mathrm{T}}, \ k = 1, 2$$
$$\mathbf{N}_{ij} = (\mathbf{q}_k^2)_i + (\mathbf{t}_k^2)_j - 2\mathbf{q}_{ik}\mathbf{t}_{kj}$$
$$\mathbf{d}_i = \sqrt{\min_j \mathbf{N}_{ij}} \ (\mathbf{N}_{ij} \ge 0.05)$$

2.3 Topological Data Analysis

In this article, we aim at characterizing the spatial arrangement of phenotypes using persistence diagrams, which are common descriptors of topological data analysis. Thus, we briefly recall, in this section, the basics of persistent homology and persistence diagrams. The interested reader can find a thorough treatment of persistence in several computational topology and algebraic topology textbooks such as [22, 14, 45].

Persistent homology. The aim of persistent homology is to encode the topological information contained in a dataset X through the lens of a filter function $f: X \to \mathbb{R}$. This is achieved by considering the sublevel sets of $f: F_{\alpha} = \{x \in X : f(x) \leq \alpha\}$. The family of sublevel sets $\mathcal{F} = \{F_{\alpha}\}_{\alpha \in \mathbb{R}}$ defines a filtration, i.e., a family of subsets of X that are nested with respect to the inclusion: $F_{\alpha} \subseteq F_{\beta}$ if $\alpha \leq \beta$. The idea of persistence is to track the topological changes occurring in the filtration as the sublevel set threshold α increases from $-\infty$ to $+\infty$. For instance, each time a topological structure such as a connected component, a handle or a void, appears in the sublevel set, we use the corresponding threshold as the so-called birth time for this structure. Similarly, each time a structure disappears in the sublevel set (think for instance of a handle being filled in after data points inside the handle were added to the sublevel set), we use the corresponding threshold as the death time. This tracking is eventually encoded in a persistence diagram, that we denote by D(f), which is a set of dots in the Euclidean plane \mathbb{R}^2 , each dot representing a topological structure whose birth and death times can be retrieved from the coordinates of the dot.

Persistence on images. In Figure 2, we provide an example of persistent homology computation performed on an image taken from the MNIST [38] dataset using the opposite of the pixel stain intensity as the filter function, so that it increases from white to black. Given a specific filter function value, the black pixels displayed in the top row of Figure 2 are those constituting the sublevel sets. One can see that at values b and d, handles are created in the union of black pixels, and they are eventually filled in at value e, for which the corresponding sublevel set includes all pixels. Other examples on our biological data are also displayed in Figures 5 and 6.

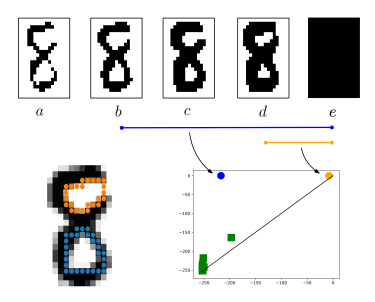


Figure 2 Example of a persistence diagram (lower right) computed on an image taken from the MNIST [38] dataset (lower left) using the opposite of the pixel stain intensity whose sublevel sets are displayed in the top row. Green squares represent connected components while the blue and orange circles represent handles, whose representative cycles are displayed on the original image.

Stability of persistence diagrams. One of the most useful properties of persistence diagrams is their *stability*: persistence diagrams computed from similar images must be similar themselves, w.r.t. the so-called *Wasserstein distances* between them.

▶ **Definition 1** ([14, 17]). The p-Wasserstein distance d_p between two persistence diagrams D, D' is defined as:

$$d_p(D,D')^p = \inf_{\gamma} \sum_{\mathrm{pt} \in D \cup \Delta} \|\mathrm{pt} - \gamma(\mathrm{pt})\|_{\infty}^p,$$

where Δ is made of an infinite number of copies of the diagonal $\{(x,x):x\in\mathbb{R}\}$ and γ ranges over all matchings between $D\cup\Delta$ and $D'\cup\Delta$.

When the sum in Definition 1 is replaced by a maximum, the Wasserstein distance becomes the so-called bottleneck distance d_{∞} . Using this distance, one can state the stability property of persistence diagrams, which shows that the Wasserstein distance between persistence diagrams is upper bounded by the distance (in the $\|\cdot\|_{\infty}$ norm) between filter functions.

▶ Theorem 2 ([13, 17]). Given a topological space X and two continuous functions $f, g: X \to \mathbb{R}$, the following inequality is true:

$$d_{\infty}(D(f), D(g)) \le ||f - g||_{\infty} \tag{1}$$

Note that similar stability results can be obtained with p-Wasserstein distances, with different upper bounds [45].

3 Methods and Results

In this section, we detail our methods to compute and analyze persistence diagrams from point clouds representing cells with different stain intensities. More specifically, we show how to discretize the cell domain into an image with stain intensity-valued pixels, from which we calculate the corresponding persistence diagrams (in homological dimension zero and one) in Section 3.1. Then, we show how to run statistical tests on persistence diagrams between different populations using Hilbert space embeddings with the *Sliced Wasserstein kernel* [11] in Section 3.2. Finally, we provide and discuss results for different patient groups (patients with different molecular subtypes, patients that survived after 8.5 years vs. deceased) in Section 3.3.

3.1 Persistence Diagrams of Cells with Stain Intensity values

In this section, we explain how persistence diagrams were computed on our point clouds representing cells so as to make use of the associated stain intensities.

Point clouds. As mentioned above, the image data for each patient is summarized in a point cloud, where the points represent cells, and have four associated stain intensities, corresponding to the CD8, CD68, PD-L1, and pancytokeratin (tumor) stains (see Section 2.1). Each patient also has two binary labels corresponding to overall survival and whether the cancer subtype is triple-negative. After removing samples with bad quality or missing labels, our final dataset is comprised of 671 point clouds. See Figure 3 for an example of such point clouds, where we only kept the cells with stain intensities above a certain threshold to ease visualization. One can see from these point clouds that different topological structures seem to emerge depending on the stain being considered: structures can be either isolated connected components corresponding to the scattered spots of cells exhibiting large stain intensity values (such as pancytokeratin (tumor) in Figure 3) or small cycles corresponding to regions where there are no cells with large stain intensity (such as CD8 in Figure 3). The lack of any discernible structure is also a possible feature if the stain intensity is diffuse across the whole tissue (such as PD-L1 in Figure 3).

Persistence Diagrams. It is common in topological data analysis to use Rips, Cech or Alpha filtrations [13] when dealing with point clouds. However, this would leave the stain intensity values aside and only provide information about the shape of the whole point cloud, which might not be sufficient to successfully encode the spatial and geometrical relationships between phenotypes.

Hence, in order to take the stain intensities into account when computing topological descriptors, we first discretized the plane into a grid of 40×40 pixels. Next, we binned the stain intensity values on this grid, so as to obtain an image. Note that the choice of resolution (i.e. the number of pixels) has to be carefully done: if the number of pixels is too



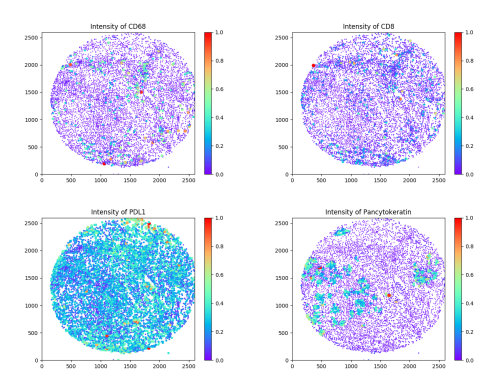


Figure 3 Illustration of the point clouds corresponding to the different stains (cell color and size is proportional to stain intensity to ease visualization). One can see that the different stain intensities induce different geometric patterns.

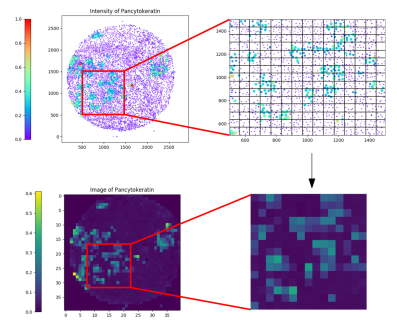


Figure 4 Discretization process turning a point cloud with stain intensity values into an image. We start with the full point cloud with the corresponding stain intensity values (upper left). Note that we only show cells above a certain stain intensity threshold to ease visualization. The cells are then placed into pixels of a grid drawn on top of the plane (upper right). These pixels with the corresponding stain intensity values are then turned into an image (down right and left).

small, one might not be able to see and compute the topological structures, but in the other hand, a resolution that is too large would induce artifacts, in the sense that all cells would be isolated, and no interesting topology could be computed. Our resolution of 40×40 pixels was manually chosen and seemed to be the best tradeoff on our data. See Figure 4 for an illustration of this process. Note also that it would be interesting to use Nadaraya-Watson kernel-based estimators (see Chapter 6 in [29]) to smooth the stain intensities of the pixels, but we left this possibility for future work.

Finally, we used persistent homology (see Section 2.3) to produce persistence diagrams out of our stain intensity-based images, by filtering the pixels with the opposite of the stain intensity (so that pixels with large stain intensity appear first). Note that points with ordinate 0 corresponds to topological structures that disappeared when adding the pixels with stain intensity 0, i.e., the pixels corresponding either to the cells not belonging to the corresponding phenotype or to pixels with no associated cells. These points should thus not be considered characteristic of the corresponding phenotype. See Figure 5 for examples of such persistence diagrams. One can see from these images that some patterns in the persistence diagrams, such as the distance-to-the-diagonal of points in homological dimension 0, or the number of points in homological dimension 1, seem to be correlated with how diffuse the cells with large stain intensity values are within the image.

Pairs of phenotypes. As mentioned in Section 1, characterizing the interactions, or colocalizations, between pairs of phenotypes might be as important, if not more, as characterizing them alone. Hence, we also computed persistence diagrams out of images with pixels colored by the average of pairs of phenotypes. This can be thought of as a similar but quite more general measure of co-localization than the one given by nearest neighbors (see Section 2.2). Indeed, the standard nearest neighbors analysis basically ranks the cells with respect to the distance to their closest neighbor. In terms of persistence, this ranking can be retrieved from the pixel filtration values: the lower they are, the more the corresponding pixels are likely to contain cells that co-localize from the two phenotypes. However, persistence diagrams also encode the interactions between the topological structures that are born from these co-localization spots. See Figure 6 for examples of such persistence diagrams. One can see from these images that the topological structures that are present in the image of a pair of phenotypes roughly include those of each phenotype alone, and that the structures that co-localize are emphasized.

Robustness. From a theoretical point of view, the stability property that persistence diagrams enjoy (see Section 2.3 and Proposition 2) is very advantageous. Indeed, it is well-known that any nearest neighbors analysis is sensitive to measurement errors: even a slight mistake in the measurement of stain intensity can induce different phenotype assignments for the cells, and thus different outputs from a nearest neighbor analysis. Since we do not depend on thresholding to compute persistence diagrams, we avoid this issue. On the other hand, the stability theorem for persistence diagrams ensures that any measurement error only has a small effect, provided that the error is small itself.

3.2 Statistics on Persistence Diagrams

In this section, we provide details about the statistical methods we used to assess the efficiency of persistence diagrams as characteristic and predictive biological descriptors.

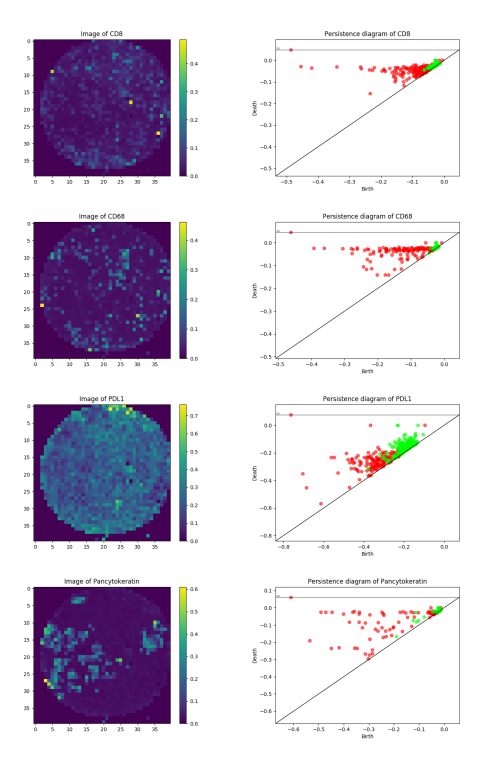


Figure 5 Examples of images with stain intensity-based pixels computed from point clouds (left) and their corresponding persistence diagrams (right). Points in homological dimension 0 are displayed in red and points in homological dimension 1 are displayed in green. From top to bottom: stains of CD8, CD68, PD-L1 and pancytokeratin.

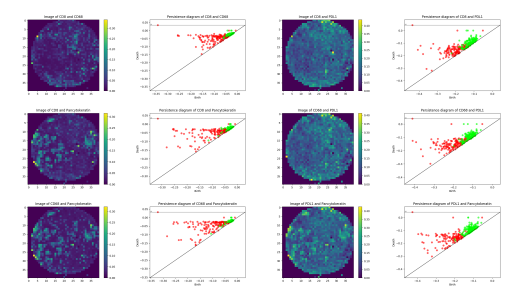


Figure 6 Examples of images and associated persistence diagrams computed from pairs of phenotypes/stains. Points in homological dimension 0 are displayed in red and points in homological dimension 1 are displayed in green.

Kernel-based Statistical Tests. In order to formally assess the statistical power of persistence diagrams with respect to the groups of interest, such as survived vs. not-survived, or triple-negative cancer subtype vs. other subtype, we need to be able to run statistical tests on distributions of persistence diagrams. Several recent works have looked at this question from a theoretical point of view [35, 52, 58]. In this article, we focus on Kernel Mean Embeddings [26], that is, we characterize a sample of a distribution \mathcal{D} of persistence diagrams $\hat{\mathcal{D}}_n = \{D_1, \dots, D_n\}$ by embedding the diagrams in a Hilbert space \mathcal{H} with a continuous map Φ , and by taking the mean (in the Hilbert space) of this sample: $\Phi(\hat{\mathcal{D}}_n) := \frac{1}{n} \sum_{i=1}^n \Phi(D_i)$. Now, given two samples $\hat{\mathcal{D}}_n$ and $\hat{\mathcal{D}}'_n$, one can compute the statistic:

$$\mathrm{MMD}(\hat{\mathcal{D}}_n, \hat{\mathcal{D}}'_n) := \|\Phi(\hat{\mathcal{D}}_n) - \Phi(\hat{\mathcal{D}}'_n)\|_{\mathcal{H}},$$

also called the maximum mean discrepancy, and use it to perform statistical tests in order to check whether \mathcal{D} and \mathcal{D}' are the same. This statistic has been shown to be a good proxy, with quantified approximation bounds, to its continuous version $\|\Phi(\mathcal{D}) - \Phi(\mathcal{D}')\|_{\mathcal{H}}$ in [26], where $\Phi(\mathcal{D})$ is defined as $\mathbb{E}_{\mathcal{D}\sim\mathcal{D}}[\Phi(\mathcal{D})]$.

Choice of the embedding function. It might not be totally clear how to choose such a map Φ for embedding persistence diagrams. This can actually be done quite easily with the use of kernels:

▶ **Definition 3.** Let $\mathcal{D}_{N,L}$ be the space of persistence diagrams with at most N points included in $[-L,L]^2$. A kernel is a pairwise function $k: \mathcal{D}_{N,L} \times \mathcal{D}_{N,L} \to \mathbb{R}$ such that the matrix $K = ((k(D_i,D_j)))_{1 \leq i,j \leq n}$ is positive semi-definite for any family of persistence diagrams $D_1, \dots, D_n \in \mathcal{D}_{N,L}$.

A useful result of kernel methods actually relates kernels to embeddings in Hilbert spaces:

▶ Proposition 4. Let k be a kernel on $\mathcal{D}_{N,L}$. Then, there exists a Hilbert space \mathcal{H}_k and a map Φ_k such that, for any $D, D' \in \mathcal{D}_{N,L}$, one has $k(D, D') = \langle \Phi(D), \Phi(D') \rangle_{\mathcal{H}_k}$.

In other words, any kernel matrix can be interpreted as a Gram matrix in an implicit (and potentially infinite-dimensional) Hilbert space. Moreover, the statistic MMD can be easily computed from k with:

$$MMD(\hat{D}_{n}, \hat{D}'_{n})^{2} = \left\langle \frac{1}{n} \sum_{i=1}^{n} \Phi(D_{i}) - \frac{1}{m} \sum_{j=1}^{m} \Phi(D'_{j}), \frac{1}{n} \sum_{i=1}^{n} \Phi(D_{i}) - \frac{1}{m} \sum_{j=1}^{m} \Phi(D'_{j}) \right\rangle_{\mathcal{H}_{k}}
= \frac{1}{n^{2}} \sum_{i=1}^{n} \sum_{u=1}^{n} \langle \Phi(D_{i}), \Phi(D_{u}) \rangle_{\mathcal{H}_{k}} + \frac{1}{m^{2}} \sum_{j=1}^{m} \sum_{v=1}^{m} \langle \Phi(D'_{i}), \Phi(D'_{v}) \rangle_{\mathcal{H}_{k}}
- \frac{2}{nm} \sum_{i=1}^{n} \sum_{j=1}^{m} \langle \Phi(D_{i}), \Phi(D'_{j}) \rangle_{\mathcal{H}_{k}}
= \frac{1}{n^{2}} ||K||_{1} + \frac{1}{m^{2}} ||K'||_{1} - \frac{2}{nm} ||\tilde{K}||_{1},$$

where K, K' and \tilde{K} are the kernel matrices computed on $\mathcal{D} \times \mathcal{D}$, $\mathcal{D}' \times \mathcal{D}'$, and $\mathcal{D} \times \mathcal{D}'$ respectively. Note however that it has been shown in [26] that MMD is a biased statistic – in practice, we compute the *unbiased* MMD, defined as:

$$\operatorname{MMD}_{u}(\hat{\mathcal{D}}_{n}, \hat{\mathcal{D}}'_{n})^{2} = \frac{1}{n(n-1)} \sum_{\substack{i=1\\u\neq i}}^{n} \langle \Phi(D_{i}), \Phi(D_{u}) \rangle_{\mathcal{H}_{k}} + \frac{1}{m(m-1)} \sum_{\substack{j=1\\v\neq j}}^{m} \langle \Phi(D'_{i}), \Phi(D'_{v}) \rangle_{\mathcal{H}_{k}} \\
- \frac{2}{nm} \sum_{i=1}^{n} \sum_{j=1}^{m} \langle \Phi(D_{i}), \Phi(D'_{j}) \rangle_{\mathcal{H}_{k}}$$

Now it only remains to pick a kernel for persistence diagrams. Several choices have been proposed in recent works [2, 6, 11, 34, 50], and we will focus on one called the *Sliced Wasserstein kernel* $k_{\rm SW}$ [11] in this work, since it has been shown to be one of the most efficient approach in different statistical tasks [11]. Its definition is based on the *Sliced Wasserstein distance* SW between persistence diagrams, which is defined (informally) as the integral over all possible lines of the 1-Wasserstein distance (see Section 2.3) computed between projections of these diagrams onto a line going through the origin. In practice, one does not compute this integral exactly but rather samples a fixed number of lines, finding the average Wasserstein distance between the corresponding projections. We refer the interested reader to [11] for a precise definition of this distance, and we merely recall the definition of the associated kernel:

▶ **Definition 5** ([11]). Let $D, D' \in \mathcal{D}_{N,L}$ and $\sigma > 0$. The Sliced Wasserstein kernel is:

$$k_{\text{SW}}(D, D') = e^{-\frac{\text{SW}(D, D')}{\sigma^2}}$$

where SW denotes the Sliced Wasserstein distance between persistence diagrams.

One can easily see that $k_{\rm SW}$ can be interpreted as a Gaussian kernel, with its only parameter σ being the corresponding bandwidth.

Characteristic kernels. There is a specific class of kernels in the literature that is of particular interest when it comes to statistical tests: the so-called *characteristic* kernels [57, 55].

▶ **Definition 6.** A kernel k is called characteristic if its corresponding map Φ_k is injective on distributions, i.e., for any pair of distributions \mathcal{D} and \mathcal{D}' , one has:

$$\|\Phi(\mathcal{D}) - \Phi(\mathcal{D}')\|_{\mathcal{H}_k} = 0 \Longrightarrow \mathcal{D} = \mathcal{D}'$$

Obviously, any statistical test based on a kernel requires it to be characteristic in order to be theoretically backed-up. Even though it is not clear whether the Sliced Wasserstein kernel is characteristic or not, there exists a strategy to build a characteristic kernel out of another one, that was first presented in [35], and that we use again in this work:

▶ Theorem 7 ([35]). Let k be a kernel on $\mathcal{D}_{N,L}$ whose associated map Φ_k is continuous and injective and whose associated Hilbert space \mathcal{H}_k is separable. Then the kernel $\tilde{k} := e^k$ is a characteristic kernel.

Theorem 7 is actually a consequence of a more general theorem that is valid on any compact metric space (the fact that $\mathcal{D}_{N,L}$ is compact, with respect to the first Wasserstein distance between persistence diagrams, was proved in [35]). Moreover, it has been shown in [11] that the map $\Phi_{k_{\text{SW}}}$ associated to k_{SW} is continuous and injective. Finally, since it is also known that $\mathcal{D}_{N,L}$ is separable [43], it follows that the Hilbert space associated to k_{SW} is separable as well, as the completion of the span of a separable space. Hence the following result:

▶ Proposition 8. The kernel $\tilde{k}_{SW} := e^{k_{SW}}$ is characteristic.

All of the statistical analysis presented in the following section has been performed with the kernel $\tilde{k}_{\rm SW}$, that we call the *characteristic Sliced Wasserstein kernel*.

Comparison with NN features. Concerning the features given by nearest neighbors analysis, i.e., the means and variances of the distribution of Euclidean distances to the closest neighbors (see Section 2.2), we use kernel-based statistical tests based on the MMD computed with a standard Gaussian kernel (which is known to be characteristic). Moreover, we also test the independence between persistence diagrams and nearest neighbors features in order to check whether these two types of features are complementary or not. Again, kernel methods can be used to define the correlation between features living in different spaces. The so-called constrained covariance (COCO for short) [27] is defined as:

$$COCO(\hat{\mathcal{D}}_n^X, \hat{\mathcal{D}}_n^Y) = \frac{1}{n} \sqrt{\|\tilde{K}_X \tilde{K}_Y\|_2},$$

where $\hat{\mathcal{D}}_n^X$ (resp. $\hat{\mathcal{D}}_n^Y$) is a sample from a distribution in a space X (resp. Y), \tilde{K}_X (resp. \tilde{K}_Y) is the centered (i.e., multiplied with $\mathbf{I} - \frac{1}{n} \ \mathbf{1} \mathbf{1}^T$) version of the kernel matrix K_X (resp. K_Y) computed on $\hat{\mathcal{D}}_n^X$ (resp. $\hat{\mathcal{D}}_n^Y$), and $\|\cdot\|_2$ is the largest singular value. It has been shown in [27] that the COCO can be used as a general measure of correlation (for random variables that are not directly comparable), since having a null COCO is equivalent to being independent (see Theorem 6 in [27]) for characteristic kernels¹.

The cited result is actually proved for the so-called universal kernels but we leave this subtlety aside in the context of this work since it has no effect on our analysis

3.3 Results

We focus on statistical significance between populations of patients instead of building a classifier. This is due to the lack of tissue area and access to tissue heterogeneity typically available in whole-slide images which are typically used for diagnosis. In an ongoing analysis, we are expanding the analysis for the same patients to whole slide imaging, where the point clouds will be $\approx 400 \times$ larger. In this section, we provide the experimental results obtained on our data using the *characteristic Sliced Wasserstein kernel* $\tilde{k}_{\rm SW}$ presented in Section 3.2 for persistence diagrams and a standard Gaussian kernel for the NN features. For both types of descriptors, the kernel bandwidth was selected manually as the median of all pairwise distances (the distances used being the Sliced Wasserstein distance for persistence diagrams and the Euclidean distance for NN features). Moreover, the p-values were computed with $2 \cdot 10^3$ random permutations. The individual sample labels were shuffled and p-values were calculated from the rank of the true labels.

Triple-negative subtype. In this first experiment, we separate the patients with respect to their cancer subtypes. More specifically, we aim at distinguishing between patients diagnosed with triple-negative breast cancer and those with other subtypes. Triple-negative breast cancer is especially interesting due to its high ability to provoke an immune response, or immunogenecity, among subtypes. However, triple-negative breast cancer patients typically have poor prognosis due to the lack of response to hormonal or receptor-status therapy. By better understanding the immune profiles associated with triple-negative breast cancers and the association with treatment response (i.e. overall survival), it could be possible to design targeted immunotherapies [42].

We show in Figure 7 (left) the p-values obtained with persistence diagrams, and the ones computed with NN features, for each (pair of) phenotypes. It can be seen from this plot that the p-values obtained with persistence diagrams are most of the time comparable to those given by NN features, with the exception of CD8 and the CD8-pancytokeratin pair. We find that the NN metrics are not significant, and this was further verified with the full NN distribution shapes. On the other hand, persistence diagrams demonstrated consistency of the p-values including CD8-involved pairs, indicating they reveal topology beyond that quantified by the NN algorithm. Moreover, 1-dimensional persistence diagrams, which encode higher-order interactions between the phenotypes (that cannot be retrieved from NN analysis), also seem to be statistically more efficient than their 0-dimensional counterparts.

Survival. In this second experiment, we now aim at distinguishing between patients that were alive at the latest follow-up after diagnosis. Although this includes causes unrelated to the breast cancer morbidity and associated treatment, such as dying of natural causes or other disease, this is still a good measure of overall disease-free survival. The corresponding p-values are displayed in Figure 7 (right). It can be seen that the p-values corresponding to persistence diagrams are in general much lower than those corresponding of NN features, especially in PD-L1 involved pairs. PD-L1 combinations are relatively rare and, as explained at the end of Section 3.1, NN features are sensitive to noise and the counting statistics on the number of phenotype pairs. Characterizing the spatial interactions of PD-L1 expression, however, would provide valuable insight into the possible immuno-repressive patterns in the tumor immune microenvironment.

We see, on the other hand, stability of persistence diagrams providing statistically significant measures. This makes diagrams a more robust descriptor than NN alone at the same statistical power. Similarly, it is clear from the distribution of values that persistence diagrams are more stable descriptors than NN features, picking up topology relating to PD-L1.

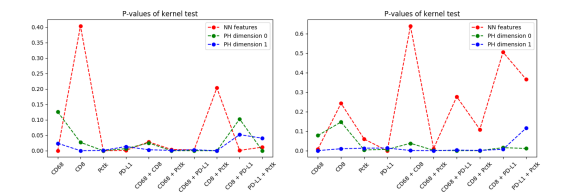


Figure 7 P-values computed by kernel-based statistical tests for NN features (red), 0-dimensional persistence diagrams (green) and 1-dimensional persistence diagrams (blue), for different (pairs of) phenotypes. "Pancytokeratin" has been abbreviated to "Pctk".

Correlation. Finally, we check the correlation values, as measured with COCO, between the NN features and persistence diagrams. We show the computed values in Figure 8. One can see that the correlation is always less than 0.1, which indicates that these features are almost statistically completely independent, and thus complementary. Moreover, these correlations seem to be oblivious to homological dimension since the shape of the curves for 0- and 1-dimensional persistence diagrams is roughly the same.

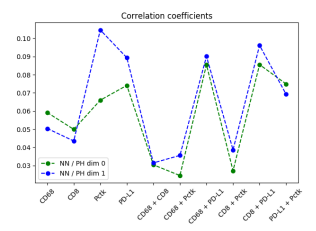


Figure 8 Correlation coefficients, measured with COCO, between NN features and 0-dimensional persistence diagrams (green), and 1-dimensional persistence diagrams (blue).

4 Open Questions and Future Work

We presented a novel approach for cancer research through the analysis of qmIF data using persistent homology, evaluated our method on a unique cohort of 671 patients using high-throughput tumor microarrays with a median 8.5 year follow-up. Our preliminary analyses show that features derived from persistent homology between groups of patients stratified by survival and triple negative status are statistically significant and are complementary to the state-of-the-art nearest neighbor approach. This indicates that the persistent homology features can be used as a complementary biomarker. Although the features do not separate the groups well enough to form a viable classifier, our results indicate feasibility of strong classification results in future work using more tissue area and larger associated point clouds. We are actively pursuing this by performing qmIF on tissue sections that contain $400\times$ more area. If a successful classifier can be built, it could be possible to characterize patient immune profiles to build specific treatments. It could also be possible to use features derived from persistent homology to study functional breast cancer dynamics. For example, the relationship between persistence diagrams and other biological data such as proteomics or genomic sequencing could reveal factors that play a role in cancer initiation or progression.

Open questions. Our preliminary study is by no means comprehensive, and many questions remain open. Here is a list of the future investigations that we plan to work on:

- We only considered single phenotypes and pairs of phenotypes. However, one might be interested in the interactions between more than two phenotypes, although this would greatly increase the number of persistence diagrams computed for each patient. Moreover, there is no single solution on how to combine the different stain intensities. In this work, we merely took the average between normalized stain intensities, even though it would be interesting to weight the filtrations given by stain intensities in order to take the range of stain intensity values into account. The weight coefficients could even be learned so as to avoid a brute force search, using for instance recent works on differentiability of persistence diagrams for learning [5, 15, 31, 48].
- More generally, the question of turning a point cloud with different stain intensity values into one (or more) persistence diagram has many different solutions, the most natural one being to use Alpha or Rips filtrations, even though it would not be satisfactory since stain intensity values would be left aside. In this work, we built images with fixed resolution, that is, number of pixels, on top of the point clouds and used these images to compute persistence. However, other choices of filtrations are possible. For instance, one could think of constructing a graph on top of the point cloud, such as a δ -neighborhood graph, and then filter this graph with the stain intensity values on the nodes. Note that the δ parameter actually plays the role of the resolution of the image.
- Multiple stain intensities actually fits into the multiparameter persistence framework [10, 28], where data is filtered by several filtrations at the same time. Our approach of taking linear combinations of stain intensities actually amounts to draw lines in this multiparameter space and compute usual persistence along this line, which is the approach that is also advocated in recent works [19, 40]. However, multiparameter persistence is a current area of research, and invariants have been obtained in recent works, at least for bifiltrations, that is, filtrations with two parameters [3, 4, 16]. Even though they are harder to encode than persistence diagrams, it might be interesting to apply these results in our context.

■ There are many different choices available when it comes to computing statistics on persistence diagrams. In this work, we restricted to kernel-based approaches, even though many other choices are available, see for instance [2, 6, 34, 50]. Overall, statistics and machine learning with persistence diagrams is also a current area of research, with new methods appearing regularly.

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