

Spectral Bridges

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Abstract

In this paper, Spectral Bridges, a novel clustering algorithm, is introduced. This algorithm builds upon the traditional k-means and spectral clustering frameworks by subdividing data into small Voronoï regions, which are subsequently merged according to a connectivity measure. Drawing inspiration from Support Vector Machine's margin concept, a non-parametric clustering approach is proposed, building an affinity margin between each pair of Voronoï regions. This approach delineates intricate, non-convex cluster structures and is robust to hyperparameter choice. The numerical experiments underscore Spectral Bridges as a fast, robust, and versatile tool for clustering tasks spanning diverse domains. Its efficacy extends to large-scale scenarios encompassing both real-world and synthetic datasets. The Spectral Bridge algorithm is implemented both in Python (https://pypi.org/project/spectral-bridges) and R https://github.com/cambroise/spectral-bridges-Rpackage).

Keywords: spectral clustering, vector quantization, scalable, non-parametric

Contents

1	Intr	oduction	2
2	Rela	ated Work	3
3	Spec	ctral Bridges	3
	3.1	Bridge affinity	4
	3.2	Algorithm	7
	3.3	Hyperparameter settings	8

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4	Nur	nerical experiments	9
	4.1	Datasets	9
		4.1.1 Real-world data	9
		4.1.2 Synthetic data	9
		4.1.3 Datasets Summary & Class Balance	9
	4.2	Metrics	10
	4.3	Platform	10
	4.4	Sensitivity to hyperparameters	10
	4.5	Time complexity	10
	4.6	Accuracy	11
	4.7	Noise robustness	14
5	Con	aclusive remarks	1 4
6	App	pendix	17
	6.1	Derivation of the bridge affinity	17
	6.2	Code	18
		6.2.1 Implementation	18
		6.2.2 Affinity matrix computation	18
Re	efere	nces	19
Se	ssion	information	20

1 Introduction

Clustering is a fundamental technique for exploratory data analysis, organizing a set of objects into distinct homogeneous groups known as clusters. It is extensively utilized across various fields, such as biology for gene expression analysis (Eisen et al. 1998), social sciences for community detection in social networks (Latouche, Birmelé, and Ambroise 2011), and psychology for identifying behavioral patterns. Clustering is often employed alongside supervised learning as a pre-processing step, helping to structure and simplify data, thus enhancing the performance and interpretability of subsequent predictive models (Verhaak et al. 2010). Additionally, clustering can be integrated into supervised learning algorithms, such as mixture of experts (Jacobs et al. 1991), as part of a multi-objective strategy.

There are various approaches to clustering, and the quality of the results is largely determined by how the similarity between objects is defined, either through a similarity measure or a distance metric. Clustering techniques originate from diverse fields of research, such as genetics, psychometry, statistics, and computer science. Some methods are entirely heuristic, while others aim to optimize specific criteria and can be related to statistical models.

Density-based methods identify regions within the data with a high concentration of points, corresponding to the modes of the joint density. A notable non-parametric example of this approach is DBSCAN (Ester et al. 1996). In contrast, model-based clustering, such as Gaussian mixture models, represents a parametric approach to density-based methods. Model-based clustering assumes that the data is generated from a mixture of underlying probability distributions, typically Gaussian distributions. Each cluster is viewed as a component of this mixture model, and the Expectation-Maximization (EM) algorithm is often used to estimate the parameters. This approach provides a probabilistic framework for clustering, allowing for the incorporation of prior knowledge and the ability to handle more complex cluster shapes and distributions (McLachlan and Peel 2000).

Geometric approaches, such as k-means (MacQueen et al. 1967), are distance-based methods that aim to partition data by optimizing a criterion reflecting group homogeneity. The k-means++ algorithm (Arthur and Vassilvitskii 2006) enhances this approach by providing faster and more reliable results. However, a key limitation of these methods is the assumption of linear boundaries between clusters, implying that clusters are convex. To address non-convex clusters, the kernel trick can be applied, allowing for a more flexible k-means algorithm. This approach is comparable to spectral clustering in handling complex cluster boundaries (Dhillon, Guan, and Kulis 2004). The k-means algorithm can also be interpreted within the framework of model-based clustering under specific assumptions (Govaert and Nadif 2003), revealing that it is essentially a special case of the more general Gaussian mixture models, where clusters are assumed to be spherical Gaussian distributions with equal variance.

Graph-based methods represent data as a graph, with vertices symbolizing data points and edges weighted to indicate the affinity between these points. Spectral clustering can be seen as a relaxed version of the graph cut algorithm (Shi and Malik 2000). However, traditional spectral clustering faces significant limitations due to its high time and space complexity, greatly hindering its applicability to large-scale problems (Von Luxburg 2007).

The method we propose aims to find non-convex clusters in large datasets, without relying on a parametric model, by using spectral clustering based on an affinity that characterizes the local density of the data. The algorithm described in this paper draws from numerous clustering approaches. The initial intuition is to detect high-density areas. To this end, vector quantization is used to divide the space into a Voronoï tessellation. An original geometric criterion is then employed to detect pairs of Voronoï regions that are either distant from each other or separated by a low-density boundary. Finally, this affinity measure is considered as the weight of an edge in a complete graph connecting the centroids of the tessellation, and a spectral clustering algorithm is used to find a partition of this graph. The two main parameters of the algorithm are the number of Voronoï Cells and the number of clusters.

The paper begins with a section dedicated to presenting the context and related algorithms, followed by a detailed description of the proposed algorithm. Experiments and comparisons with reference algorithms are then conducted on both real and synthetic data.

2 Related Work

Spectral clustering is a graph-based approach that computes the eigenvectors of the graph's Laplacian matrix. This technique transforms the data into a lower-dimensional space, making the clusters more discernible. A standard algorithm like k-means is then applied to these transformed features to identify the clusters (Von Luxburg 2007). Spectral clustering enables capturing complex data structures and discerning clusters based on the connectivity of data points in a transformed space, effectively treating it as a relaxed graph cut problem.

Classical spectral clustering involves two phases: construction of the affinity matrix and eigendecomposition. Constructing the affinity matrix requires $O(n^2d)$ time and $O(n^2)$ memory, while eigendecomposition demands $O(n^3)$ time and $O(n^2)$ memory, where n is the data size and d is the dimension. As n increases, the computational load escalates significantly (Von Luxburg 2007).

To mitigate this computational burden, one common approach is to sparsify the affinity matrix and use sparse eigensolvers, reducing memory costs but still requiring computation of all original matrix entries (Von Luxburg 2007). Another strategy is sub-matrix construction. The Nyström method randomly selects m representatives from the dataset to form an $n \times m$ affinity sub-matrix (Chen et al. 2010). Cai et al. extended this with the landmark-based spectral clustering method, which uses k-means to determine m cluster centers as representatives (Cai and Chen 2014). Ultra-scalable spectral

clustering (U-SPEC) employs a hybrid representative selection strategy and a fast approximation method for constructing a sparse affinity sub-matrix (Huang et al. 2019).

Other approaches use the properties of the small initial clusters for the affinity computation. Clustering Based on Graph of Intensity Topology (GIT) estimates for example a global topological graph (topo-graph) between local clusters (Gao et al. 2021). It then uses the Wasserstein Distance between predicted and prior class proportions to automatically cut noisy edges in the topo-graph and merge connected local clusters into final clusters.

The issue of characterizing the affinity between two clusters to create an edge weight is central to the efficiency of a spectral clustering algorithm operating from a submatrix.

Notice that the clustering robustness of many Spectral clustering algorithms heavily relies on the proper selection of kernel parameter, which is difficult to find without prior knowledge (Ng, Jordan, and Weiss 2001).

3 Spectral Bridges

The proposed algorithm uses k-means centroids for vector quantization defining Voronoï region, and a strategy is proposed to link these regions, with an "affinity" gauged in terms of minimal margin between pairs of classes. These affinities are considered as weight of edges defining a completely connected graph whose vertices are the regions. Spectral clustering on the region provide a partition of the input space. The sole mandatory parameters of the algorithm are the number of Voronoï regions and the number of final clusters.

3.1 Bridge affinity

The basic idea involves calculating the difference in inertia achieved by projecting onto a segment connecting two centroids, rather than using the two centroids separately (see Figure 1). If the difference is small, it suggests a low density between the classes. Conversely, if this difference is large, it indicates that the two classes may reside within the same densely populated region.



Figure 1: Balls (left) versus Bridge (right). The inertia of each structure is the sum of the squared distances represented by grey lines.

Let us consider a sample $X=(\mathbf{x}_i)_{i\in\{1,\cdots,n\}}$ of vectors $\mathbf{x}_i\in\mathbb{R}^d$ and a set of m coding vectors $(\mathbf{\mu}_k)_{k\in\{1,\cdots,m\}}$ defining a partition $P=\{\mathcal{V}_1,\cdots,\mathcal{V}_m\}$ of \mathbb{R}^d into m Voronoï regions:

$$\mathcal{V}_k = \left\{ \mathbf{x} \in \mathbb{R}^d \mid \|\mathbf{x} - \boldsymbol{\mu}_k\| \le \|\mathbf{x} - \boldsymbol{\mu}_j\| \text{ for all } j \ne k \right\}.$$

In the following a ball denotes the subset of X in a Voronoï region. The inertia of two balls \mathcal{V}_k and \mathcal{V}_l is

$$I_{kl} = \sum_{\mathbf{x}_i \in \mathcal{V}_k} \|\mathbf{x}_i - \boldsymbol{\mu}_k\|^2 + \sum_{\mathbf{x}_i \in \mathcal{V}_l} \|\mathbf{x}_i - \boldsymbol{\mu}_l\|^2.$$

We define a bridge as a structure defined by a segment connecting two centroids μ_k and μ_l . The inertia of a bridge between \mathcal{V}_k and \mathcal{V}_l is defined as

$$B_{kl} = \sum_{\boldsymbol{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l} \|\boldsymbol{x}_i - \boldsymbol{p}_{kl}(\boldsymbol{x}_i)\|^2,$$

where

$$\boldsymbol{p}_{kl}(\boldsymbol{x}_i) = \boldsymbol{\mu}_k + t_i(\boldsymbol{\mu}_l - \boldsymbol{\mu}_k),$$

with

$$t_i = \min\left(1, \max\left(0, \frac{\langle \mathbf{x}_i - \mathbf{\mu}_k | \mathbf{\mu}_l - \mathbf{\mu}_k \rangle}{\|\mathbf{\mu}_l - \mathbf{\mu}_k\|^2}\right)\right).$$

Considering two centroïds, the normalized average of the difference between Bridge and balls inertia (see Appendix) constitutes the basis of our affinity measure between two regions:

$$\frac{B_{kl} - I_{kl}}{(n_k + n_l) \|\boldsymbol{\mu}_k - \boldsymbol{\mu}_l\|^2} = \frac{\sum_{\boldsymbol{x}_i \in \mathcal{V}_k} \langle \boldsymbol{x}_i - \boldsymbol{\mu}_k | \boldsymbol{\mu}_l - \boldsymbol{\mu}_k \rangle_+^2 + \sum_{\boldsymbol{x}_i \in \mathcal{V}_l} \langle \boldsymbol{x}_i - \boldsymbol{\mu}_l | \boldsymbol{\mu}_k - \boldsymbol{\mu}_l \rangle_+^2}{(n_k + n_l) \|\boldsymbol{\mu}_k - \boldsymbol{\mu}_l\|^4},$$

$$= \frac{\sum_{\boldsymbol{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l} \alpha_i^2}{n_k + n_l},$$

where

$$\alpha_i = \begin{cases} t_i, & \text{if } t_i \in [0, 1/2], \\ 1 - t_i, & \text{if } t_i \in]1/2, 1]. \end{cases}$$

The basic intuition behind this affinity is that t_i represents the relative position of the projection of \mathbf{x}_i on the segment $[\boldsymbol{\mu}_k, \boldsymbol{\mu}_l]$. α_i represents the relative position on the segment, with the centroid of the class to which \mathbf{x}_i belongs as the reference point.

This quantity can also be understood in relation to a local form of PCA. Unlike the standard PCA inertia criterion, this approach is directional and limited to the union of two Voronoï cells.

Let $X_{|k|}$ denote the data matrix restricted to the region \mathcal{V}_k . The projection operator onto the μ_k -centered segment $[0, \mu_l - \mu_k]$ is given by:

$$\pi_{kl}(\cdot) = p_{kl}(\cdot) - \mu_{kl}$$

It then follows that:

$$\frac{\sum_{\mathbf{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l} \alpha_i^2}{n_k + n_l} = \frac{\|\mathbf{\pi}_{kl}(X_{|k})\|_F^2 + \|\mathbf{\pi}_{lk}(X_{|l})\|_F^2}{(n_k + n_l)\|\mathbf{\mu}_k - \mathbf{\mu}_l\|^2}$$

where $\|\cdot\|_F$ is the Frobenius norm.

This formulation is particularly useful for numerical implementation.

Moreover, the boundary that separates the two clusters defined by centroids μ_k and μ_l is a hyperplane \mathcal{P}_{kl} . This hyperplane is orthogonal to the line segment connecting the centroids and intersects this segment at its midpoint.

If we consider all points $\mathbf{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l$ which are not projected on centroids but somewhere on the segment, the distance from a point to the hyperplane is

$$d(\mathbf{x}_i, \mathscr{T}_{kl}) = (1/2 - \alpha_i) \|\mathbf{\mu}_k - \mathbf{\mu}_l\|.$$

This distance is similar to the concept of margin in Support Vector Machine (Cortes and Vapnik 1995). When the α_i values are small (close to zero since $\alpha_i \in [0, 1/2]$), the margins to the hyperplane are large, indicating a low density between the classes. Conversely, if the margins are small, it suggests that the two classes may reside within the same densely populated region. Consequently, the sum of the α_i or α_i^2 increases with the density of the region between the classes (See Figure Figure 2).



Figure 2: Spectral Bridge affinity illustration involving two centroids. The bold black dots mark the centroids of each cluster, while the colored cells represent the final partition of data points. In subfigures (a) and (c), the length of each dotted grey segment is proportional to $1/2 - \alpha_i$, whereas the thin black segments are proportional to α_i . Subfigures (b) and (d) depict the distribution of α_i , showing the behavior when clusters are either closely positioned (a, b) or well-separated (c, d).

Note that the criterion is local and indicates the relative difference in densities between the balls and the bridge, rather than evaluating a global score for the densities of the structures.

Eventually, we define the bridge affinity between centroids k and l as the square root of the variance gain:

$$a_{kl} = \begin{cases} 0, & \text{if } k = l, \\ \sqrt{\frac{\sum_{\mathbf{x_l} \in \mathcal{V}_k \cup \mathcal{V}_l} \alpha_i^2}{n_k + n_l}}, & \text{otherwise.} \end{cases}$$

The inclusion of the square root redefines the variance affinity measure. Rather than using the squared Euclidean norm, the affinity is interpreted as a quadratic mean, representing the ratio of the standard deviation to the length of the segment connecting two centroids.

This concept can be generalized by introducing the p-bridge affinity for any p>0 using the Minkowski mean:

$$a_{p,kl} = \begin{cases} 0, & \text{if } k = l, \\ \left(\frac{\sum_{\mathbf{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l} \alpha_i^p}{n_k + n_l}\right)^{1/p}, & \text{otherwise.} \end{cases}$$

Both definitions are equivalent when p = 2. For p = 1, the affinity aligns directly with the SVM model previously discussed. Note that this yields a bounded metric in [0, 1/2].

To allow points with large margin to dominate and make the algorithm more robust to noise and outliers we consider the following exponential transformation:

$$\tilde{a}_{kl} = g(a_{kl}) = \exp(\gamma a_{kl}). \tag{1}$$

where γ is a scaling factor. This factor is set to ensure a large enough separation between the final coefficients. This factor is determined by the equation:

$$\gamma = \frac{\log(M)}{q_{90} - q_{10}}$$

where q_{10} and q_{90} are respectively the 10th and 90th percentiles of the original affinity matrix and M > 1. Thus, since the transformation is order-preserving, the 90th percentile of the newly constructed matrix is M times greater than the 10th percentile. By default, M is arbitrarily set to a large value of 10^4 .

This regularization is crucial: with a bounded affinity metric, exponentiation enhances the separation between low and high-density regions, controlled by a scaling parameter, as in traditional spectral clustering. Redefining the metric with a square root (or power 1/p for the generalized affinity) helps mitigate a converse issue. Omitting this step would entail $a_{p,kl} \leq 2^{-p}$. Machine error could cause numerical instability when solving the Laplacian eigenproblem, especially if values become too small or too large, since the range of affinity values can become wide when the initial ratio between the largest and smallest non-zero unscaled bridge affinities is high. This transformation reduces the maximum values in the affinity matrix while preserving the metric's interpretability and distance-like properties; importantly, this adjustment is not intended for outlier detection.

3.2 Algorithm

The Spectral Bridges algorithm first identifies local clusters to define Voronoï regions, computes edges with affinity weights between these regions, and ultimately cuts edges between regions with low inter-region density to determine the final clusters (see Algorithm 1 and Figure 3).

In spectral clustering, the time complexity is usually dominated by the eigendecomposition step, which is $O(n^3)$. However, in the case of Spectral Bridges, the k-means algorithm has a time complexity of $O(n \times m \times d)$. For datasets with large n, this can be more significant than the $O(m^3)$ time complexity of the Spectral Bridges eigendecomposition. As for the affinity matrix construction, there are m^2 coefficients to be calculated. Each a_{kl} coefficient requires the computation of $n_k + n_l$ dot products as well as the norm $\|\mu_k - \mu_l\|$, the latter often being negligeable. Assuming that the Voronoï regions are roughly balanced in cardinality, we have $n_k \approx \frac{n}{m}$. Since m should always be less than n, therefore $\frac{n}{m} > 1$ and the time complexity of the affinity matrix is $O(\frac{n}{m} \times m^2 \times d) = O(n \times m \times d)$ given the acceptable range of values for m. Nonetheless, this is rarely the bottleneck.

Algorithm 1 Spectral Bridges

- 1: **procedure** SpectralBridges $(X, k, m) \triangleright X$: input dataset, k: number of clusters, m: number of Voronoï regions
- 2: Step 1: Vector Quantization
- 3: centroids, voronoiRegions \leftarrow KMEANS(X, m) \triangleright Initial centroids and Voronoi regions using k-means++
- 4: Step 2: Affinity Computation
- 5: $A = \{g(a_{kl})\}_{kl} \leftarrow Affinity(X, centroids, voronoiRegions)$ \triangleright Compute affinity matrix A
- 6: **Step 3: Spectral Clustering** ▷ Assign each region to a cluster
- 7: labels \leftarrow SpectralClustering(A, k)
- 8: **Step 4: Propagate** > Assign each data point to the cluster of its region
- 9: clusters \leftarrow Propagate(X, labels, voronoiRegions)
- 10: **return** clusters \triangleright Return cluster labels for data points in X
- 11: end procedure



Figure 3: Illustration of the Spectral Bridges algorithm with the Iris dataset (first principal plane). The bold red dots represent the centroids of the clusters, while the colored cells indicate the final partition of the data points. Vector quantization (Step 1 of Algorithm 1), Affinity computation (Step 2 of Algorithm 1), Spectral clustering and spreading (Step 3-4 of Algorithm 1).

3.3 Hyperparameter settings

The proposed algorithm requires three input parameters: the number of clusters K, the number of Voronoï regions m, and a scaling parameter for the spectral clustering phase.

Model selection in non-parametric settings is challenging due to the absence of predefined model parameters. It relies heavily on data-driven approaches. Metrics like the Gap Statistic (Tibshirani,

Walther, and Hastie 2001) and the Laplacian eigengap (Von Luxburg 2007) are potential tools for hyperparameter selection.

We propose a method for choosing the scaling parameter (see Equation Equation 1) that yields stable results. Selecting both m, the number of Voronoï regions, and K, the number of clusters, is difficult. We address this by adopting a heuristic: first, choose K, then determine m using a modified Laplacian eigengap.

If K represents the true number of clusters, the affinity matrix resembles a graph adjacency matrix with K connected components. This configuration is characterized by an eigengap at the Kth eigenvalue. In Self-Tuning Spectral Clustering (Zelnik-Manor and Perona 2004), the eigengap $\lambda_{K+1} - \lambda_K$ is used to evaluate clustering quality for K clusters. Following a similar strategy, and assuming K is known, the Laplacian eigengap at the Kth eigenvalue can select M, with the scaling parameter fixed.

Determining the optimal value of m using the eigengap is not straightforward. As the affinity matrix dimension increases, the number of eigenvalues grows, reducing gaps between them. This makes direct comparisons unreliable. To address this, we use the ratio $\rho = (\lambda_{K+1} - \lambda_K)/\lambda_{K+1}$. This metric is bounded between 0 and 1 and measures the relative difference between consecutive eigenvalues. It facilitates meaningful comparisons across different values of m. A value of R close to 1 suggests high clustering quality, whereas lower values indicate weaker performance.

Using this metric, we determine a near-optimal value for m by maximizing the average R across possible values of m. Additionally, the metric enhances robustness by running the algorithm with different random seeds and selecting the clustering result with the highest normalized eigengap.

4 Numerical experiments

In this section, the results obtained from testing the Spectral Bridges algorithm on various datasets, both small and large scale, including real-world and well-known synthetic datasets, are presented. These experiments assess the accuracy, time and space complexity, ease of use, robustness, and adaptability of our algorithm. We compare Spectral Bridges (SB) against several state-of-the-art methods, including k-means++ (KM) (MacQueen et al. 1967; Arthur and Vassilvitskii 2006), Expectation-Maximization (EM) (Dempster, Laird, and Rubin 1977), Ward Clustering (WC) (Ward Jr 1963), DB-SCAN (DB) (Ester et al. 1996) and GIT (Gao et al. 2021). This comparison establishes baselines across centroid-based clustering algorithms, hierarchical methods, and density-based methods.

The algorithms are evaluated on both raw and Principal Component Analysis processed (PCA-processed) data with varying dimensionality. For synthetic datasets, Gaussian and/or uniform noise is introduced to assess the robustness of the algorithm.

4.1 Datasets

4.1.1 Real-world data

- MNIST: A large dataset containing 60,000 handwritten digit images in ten balanced classes, commonly used for image processing benchmarks. Each image consists of $28 \times 28 = 784$ pixels.
- UCI ML Breast Cancer Wisconsin: A dataset featuring computed attributes from digitized images of fine needle aspirates (FNA) of breast masses, used to predict whether a tumor is malignant or benign.

4.1.2 Synthetic data

- **Impossible**: A synthetic dataset designed to challenge clustering algorithms with complex patterns.
- Moons: A two-dimensional dataset with two interleaving half-circles.
- Circles: A synthetic dataset of points arranged in two non-linearly separable circles.
- **Smile**: A synthetic dataset with points arranged in the shape of a smiling face, used to test the separation of non-linearly separable data.

4.1.3 Datasets Summary & Class Balance

Table 1: Datasets Summary & Class Balance

		#Sam-		
Dataset	#Dims ples		#Classes Class Proportions	
MNIST	784	60000	10	9.9%, 11.2%, 9.9%, 10.3%, 9.7%, 9%, 9.9%, 10.4%, 9.7%, 9.9%
UCI Breast	30	569	2	37.3%, 62.7%
Cancer				
Impossible	2	3594	7	24.8%, 18.8%, 11.3%, 7.5%, 12.5%, 12.5%, 12.5%
Moons	2	1000	2	50%, 50%
Circles	2	1000	2	50%, 50%
Smile	2	1000	4	25%, 25%, 25%, 25%

Class proportions are presented in ascending order starting from label 0.

4.2 Metrics

To evaluate the performance of the clustering algorithm, the Adjusted Rand Index (ARI) (Halkidi, Batistakis, and Vazirgiannis 2002) and Normalized Mutual Information (NMI) (Cover and Thomas 1991) are used. ARI measures the similarity between two clustering results, ranging from -0.5 to 1, with 1 indicating perfect agreement. NMI ranges from 0 to 1, with higher values indicating better clustering quality. In some tests, the variability of scores across multiple runs is also reported due to the random initialization in k-means, though k-means++ generally provides stable and reproducible results.

4.3 Platform

All experiments were conducted on an Archlinux machine with Linux 6.9.3 Kernel, 8GB of RAM, and an AMD Ryzen 3 7320U processor.

4.4 Sensitivity to hyperparameters

The hyperparameters of the Spectral Bridges algorithm were based on the size of each dataset, *n*, and the number of clusters, *K*.

To better grasp the sensitivity with respect to m, the number of Voronoï cells, Spectral Bridges was run on the PCA h=32 embedded MNIST dataset with varying values of $m \in \{10, 120, 230, 340, 450, 560, 670, 780, 890, 1000\}$. The case m=10 is equivalent to the k-means++ algorithm. ARI and NMI scores are recorded over 20 consecutive iterations and subsequently plotted. As shown by Figure 4, the accuracy seems to be consistently increasing with values of m, with the largest observed gap occurring between values of m=10 and m=120, and flattening thereafter,

indicating a tremendous improvement over the classical k-means++ framework even for empirically suboptimal hyperparameter values.



Figure 4: ARI and NMI scores of Spectral Bridges with varying values of m.

For other algorithms, such as DBSCAN, labels were used to determine the best hyperparameter values to compare our method against the "best case scenario", thus putting the Spectral Bridges algorithm at a voluntary disadvantage.

4.5 Time complexity

To assess the algorithm's time complexity, the average execution times over 50 runs were computed for varying numbers of Voronoï regions m as well as dataset sizes. With a constant number of clusters K=5 and an embedding dimension of d=10, the results (see Figure 5) highlight Spectral Bridges algorithm's efficacy. As discussed previously, we observe a linear relationship between m and the execution time because the matrix construction is highly optimized and the time taken is almost negligeable compared to that of the initial k-means++ centroids initalization.

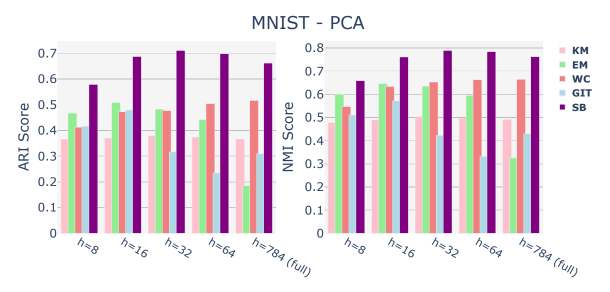


Figure 5: Average time taken per model fit.

4.6 Accuracy

The algorithm's accuracy was first evaluated on the MNIST dataset. Metrics were collected to compare our method with k-means++, EM, GIT, and Ward clustering. Metrics were estimated by computing the empirical average over 10 consecutive runs for each method. Due to limited computational resources, we randomly selected a sample of 20,000 data points (one-third of the total) for each run, on which all algorithms were trained and tested. To ensure reproducibility, a fixed random seed was set at the beginning of all scripts. Note, however, that this does not imply centroids were initialized identically for centroid based methods, as these may vary according to the implementation of each tested algorithm.

Let h denote the embedding dimension of the dataset. Spectral Bridges was tested both on the raw MNIST dataset without preprocessing (h = 784) and after reducing its dimension using PCA to $h \in \{8, 16, 32, 64\}$ (see Figure 6).



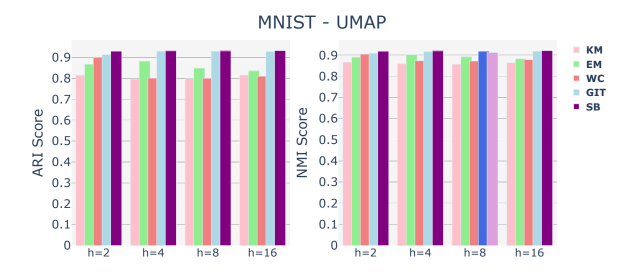
(a) ARI and NMI scores of k-means++ (pink), EM (green), Ward Clustering (red), GIT (blue), and Spectral Bridges (purple).

	Algo-					
Metric	rithm	h = 8	h = 16	h = 32	h = 64	h = 784 (full)
ARI	KM	0.3661	0.3701	0.3794	0.3740	0.3665
	EM	0.4672	0.5081	0.4826	0.4422	0.1850
	WC	0.4123	0.4729	0.4771	0.5043	0.5164
	GIT	0.4162	0.4792	0.3166	0.2347	0.3094
	SB	0.5789	0.6875	0.7110	0.6983	0.6619
NMI	KM	0.4778	0.4899	0.5037	0.4969	0.4915
	EM	0.6007	0.6462	0.6355	0.5952	0.3252
	WC	0.5471	0.6340	0.6532	0.6634	0.6648
	GIT	0.5110	0.5720	0.4227	0.3314	0.4297
	SB	0.6592	0.7616	0.7895	0.7846	0.7628

Figure 6: Comparison of clustering algorithms on PCA embedded and full MNIST with ARI and NMI metrics. The highest mean scores among tested algorithms are in bold.

Additionally, the proposed algorithm was evaluated on the MNIST dataset after reducing its di-

mensionality to $h \in \{2, 4, 8, 16\}$ using UMAP (McInnes et al. 2018), a state-of-the-art nonlinear dimensionality reduction technique (see Figure 7). To enhance the clustering performance of Spectral Bridges, the normalized eigengap method was applied. This approach trains the algorithm with multiple initializations and determines the optimal number of Voronoï cells by selecting the configuration with the largest normalized eigengap (refer to Section 3.3).



(a) ARI and NMI scores of k-means++ (pink), EM (green), Ward Clustering (red), GIT (blue), Spectral Bridges (purple).

	Algo-				
Metric	rithm	h = 2	h = 4	h = 8	h = 16
ARI	KM	0.8154	0.7971	0.7985	0.8160
	EM	0.8670	0.8828	0.8494	0.8359
	WC	0.9008	0.8015	0.8004	0.8105
	GIT	0.9141	0.9298	0.9297	0.9289
	SB	0.9304	0.9318	0.9315	0.9332
NMI	KM	0.8664	0.8586	0.8574	0.8641
	EM	0.8908	0.9025	0.8916	0.8826
	WC	0.9047	0.8733	0.8721	0.8785
	GIT	0.9081	0.9172	0.9183	0.9184
	SB	0.9182	0.9198	0.9125	0.9214

Figure 7: Comparison of clustering algorithms on UMAP embedded MNIST with ARI and NMI metrics. The highest mean scores among tested algorithms are in bold.

Note the Spectral Bridges algorithm shines particularly even with quite simple dimension reduction algorithms.

For visualization purposes, the predicted clusters by Spectral Bridges and k-means++ were projected using UMAP to compare them against the ground truth labels and to better understand the cluster shapes (see Figure 8). Note that this projection was not used in the experiments as an embedding, and thus does not play any role in the clustering process itself. As a matter of fact, the embedding used was obtained with Principal Componant Analysis (PCA), h = 32 and 500 Voronoï regions. Note that the label colors match the legend only in the case of the ground truth data. Indeed, the ordering

of the labels have no significance on clustering quality. Importantly, Spectral Bridges demonstrates remarkable accuracy, with the exception of the classes representing digits "4" and "9", which appear to have been merged.

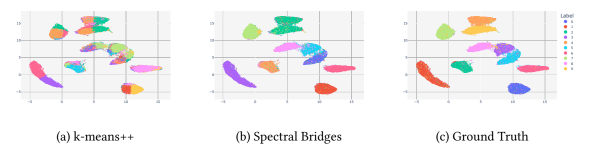


Figure 8: UMAP projection of predicted clusters against the ground truth labels.

The Spectral Bridges algorithm was also put to the test against the same competitors using scikit-learn's UCI Breast Cancer data. Once again, the normalized eigengap method was used, and the presented algorithm performed well, although the advantage was not as obvious in this case (see Figure 9). However, in none of our tests has it ranked worse than k-means++. The results are displayed as a boxplot generated from 200 iterations of each algorithm using a different seed, in order to better grasp the variability lying in the seed dependent nature of the k-means++, Expectation Maximization and Spectral Bridges algorithms.

Since the Spectral Bridges algorithm is expected to excel at discerning complex and intricate cluster structures, an array of four toy datasets was collected, as illustrated in Figure 10.

Multiple algorithms, including the proposed one, were benchmarked in the exact same manner as for the UCI Breast Cancer data. The results show that the proposed method outperforms all tested algorithms (DBSCAN, k-means++, Expectation Maximization, GIT, and Ward Clustering) while requiring few hyperparameters. As previously discussed, DBSCAN's parameters were optimized using the ground truth labels to represent a best-case scenario; however, in practical applications, suboptimal performance is more likely. Despite this optimization, the Spectral-Bridge algorithm still demonstrates superior ability to capture and represent the underlying cluster structures.

4.7 Noise robustness

To evaluate the noise robustness of the algorithm, two experimental setups were devised: one involved introducing Gaussian-distributed perturbations to the data, and the other involved concatenating uniformly distributed points within a predefined rectangular region (determined by the span of the dataset) to the existing dataset. As illustrated in Figure 12, the tests demonstrate that in both scenarios, the algorithm exhibits a high degree of insensitivity to noise.

5 Conclusive remarks

Spectral Bridges is an original clustering algorithm which presents a novel approach by integrating the strengths of traditional k-means and spectral clustering frameworks. This algorithm utilizes a simple affinity measure for spectral clustering, which is derived from the minimal margin between pairs of Voronoï regions.

The algorithm demonstrates scalability, handling large datasets efficiently through a balanced computational complexity between the k-means clustering and eigendecomposition steps. As a non-



(a) ARI and NMI scores of k-means++ (pink), EM (green), Ward Clustering (red), GIT (blue), and Spectral Bridges (purple) on the UCI Breast Cancer dataset.

Metric	Algorithm	UCI Breast Cancer		
ARI	KM	0.6675		
	EM	0.7718		
	WC	0.5750		
	GIT	0.2138		
	SB	0.6985		
NMI	KM	0.5502		
	EM	0.6586		
	WC	0.4569		
	GIT	0.2576		
	SB	0.5787		

Figure 9: Comparison of clustering algorithms on the UCI Breast Cancer dataset with ARI and NMI metrics. The highest mean scores among tested algorithms are in bold.

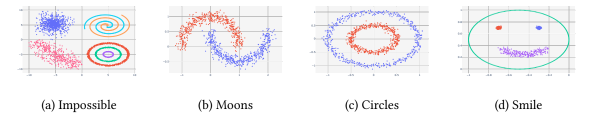
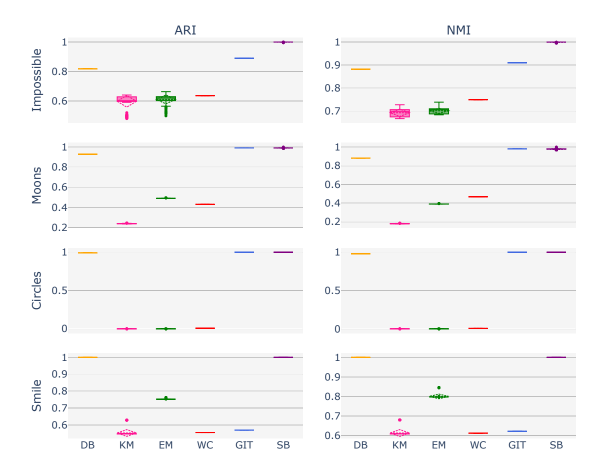


Figure 10: Four toy datasets.



(a) ARI and NMI scores of Spectral Bridges and competitors on standard synthetic toy datasets.

Metric	Algo- rithm	Impossible	Moons	Circles	Smile
ARI	DB	0.8180	0.9294	0.9920	1.0000
	KM	0.5984	0.2390	-0.0009	0.5522
	EM	0.6112	0.4897	-0.0009	0.7526
	WC	0.6368	0.4298	0.0068	0.5548
	GIT	0.8895	0.9920	1.0000	0.5693
	SB	0.9996	0.9912	1.0000	1.0000
NMI	DB	0.8823	0.8825	0.9803	1.0000
	KM	0.6936	0.1807	0.0000	0.6126
	EM	0.7021	0.3905	0.0001	0.8001
	WC	0.7505	0.4690	0.0056	0.6120
	GIT	0.9103	0.9812	1.0000	0.6210
	SB	0.9995	0.9787	1.0000	1.0000

Figure 11: Comparison of clustering algorithms on toy datasets with ARI and NMI metrics. The highest mean scores among tested algorithms are in bold.

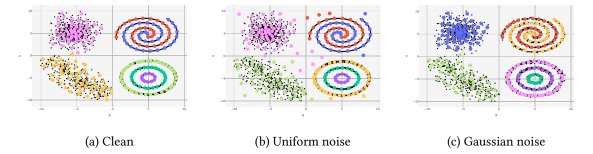


Figure 12: Three representations of the algorithm's predicted cluster centers are displayed as colored dots, with each point of the Impossible dataset shown as a small black dot. In the left graph, the dataset is unmodified. In the center graph, 250 uniformly distributed samples were added. In the right graph, Gaussian noise perturbations with $\sigma = 0.1$ were applied.

parametric method, Spectral Bridges does not rely on strong assumptions about data distribution, enhancing its versatility across various data types. It performs exceptionally well with both synthetic and real-world data and consistently outperforms conventional clustering algorithms such as k-means, DBSCAN, and mixture models.

The design of Spectral Bridges ensures robustness to noise, a significant advantage in real-world applications. Additionally, the algorithm requires minimal hyperparameters, primarily the number of Voronoï regions, making it straightforward to tune and deploy.

Furthermore, Spectral Bridges can be kernelized, allowing it to handle data in similarity space directly, which enhances its flexibility and applicability. Overall, Spectral Bridges is a powerful, robust, and scalable clustering algorithm that offers significant improvements over traditional methods, making it an excellent tool for advanced clustering tasks across numerous domains.

6 Appendix

6.1 Derivation of the bridge affinity

We denote a bridge as a segment connecting two centroids μ_k and μ_l . The inertia of a bridge between \mathcal{V}_k and \mathcal{V}_l is defined as

$$B_{kl} = \sum_{\boldsymbol{x}_i \in \mathcal{V}_k \cup \mathcal{V}_l} \|\boldsymbol{x}_i - \boldsymbol{p}_{kl}(\boldsymbol{x}_i)\|^2,$$

where

$$\boldsymbol{p}_{kl}(\boldsymbol{x}_i) = \boldsymbol{\mu}_k + t_i(\boldsymbol{\mu}_l - \boldsymbol{\mu}_k),$$

with

$$t_i = \min\left(1, \max\left(0, \frac{\langle \mathbf{x}_i - \mathbf{\mu}_k | \mathbf{\mu}_l - \mathbf{\mu}_k \rangle}{\|\mathbf{\mu}_l - \mathbf{\mu}_k\|^2}\right)\right).$$

 B_{kl} , the bridge inertia between centroids k and l, can be expressed as the sum of three terms, which represents the projection onto each centroids and onto the segment:

$$B_{kl} = \sum_{i|t_i=0} \|\mathbf{x}_i - \mathbf{\mu}_k\|^2 + \sum_{i|t_i=1} \|\mathbf{x}_i - \mathbf{\mu}_l\|^2 + \sum_{i|t_i\in[0,1[} \|\mathbf{x}_i - \mathbf{p}_{kl}(\mathbf{x}_i)\|^2.$$

The last term may be decomposed in two parts corresponding to the points of the two Voronoï regions which are projected on the segment:

$$\sum_{i|t_i\in]0,1[}\|\boldsymbol{x}_i-\boldsymbol{p}_{kl}(\boldsymbol{x}_i)\|^2 = \sum_{i|t_i\in]0,\frac{1}{n}[}\|\boldsymbol{x}_i-\boldsymbol{p}_{kl}(\boldsymbol{x}_i)\|^2 + \sum_{i|t_i\in[\frac{1}{n},1[}\|\boldsymbol{x}_i-\boldsymbol{p}_{kl}(\boldsymbol{x}_i)\|^2$$

and each part further decomposed using Pythagore

$$\begin{split} \sum_{i|t_i \in]0,\frac{1}{2}[} \| \mathbf{x}_i - \mathbf{p}_{kl}(\mathbf{x}_i) \|^2 &= \sum_{i|t_i \in]0,\frac{1}{2}[} \| \mathbf{x}_i - \mathbf{\mu}_k \|^2 - \sum_{i|t_i \in]0,\frac{1}{2}[} \| \mathbf{\mu}_k - \mathbf{p}_{kl}(\mathbf{x}_i) \|^2 \\ &= \sum_{i|t_i \in]0,\frac{1}{2}[} \| \mathbf{x}_i - \mathbf{\mu}_k \|^2 - \sum_{i|t_i \in]0,\frac{1}{2}[} \| t_i(\mathbf{\mu}_k - \mathbf{\mu}_l) \|^2, \end{split}$$

$$\sum_{i|t_{i}\in]\frac{1}{2},1[}\|\boldsymbol{x}_{i}-\boldsymbol{p}_{kl}(\boldsymbol{x}_{i})\|^{2} = \sum_{i|t_{i}\in]0,\frac{1}{2}[}\|\boldsymbol{x}_{i}-\boldsymbol{\mu}_{l}\|^{2} - \sum_{i|t_{i}\in]0,\frac{1}{2}[}\|\boldsymbol{\mu}_{l}-\boldsymbol{p}_{kl}(\boldsymbol{x}_{i})\|^{2}$$

$$= \sum_{i|t_{i}\in]\frac{1}{2},1[}\|\boldsymbol{x}_{i}-\boldsymbol{\mu}_{k}\|^{2} - \sum_{i|t_{i}\in]0,\frac{1}{2}[}\|(1-t_{i})(\boldsymbol{\mu}_{k}-\boldsymbol{\mu}_{l})\|^{2}$$

Thus

$$\begin{split} B_{kl} - I_{kl} &= \sum_{i|t_i \in]0, \frac{1}{2}[} t_i^2 \| \pmb{\mu}_k - \pmb{\mu}_l \|^2 + \sum_{i|t_i \in]\frac{1}{2}, 1[} (1 - t_i)^2 \| \pmb{\mu}_k - \pmb{\mu}_l \|^2, \\ \frac{B_{kl} - I_{kl}}{\| \pmb{\mu}_k - \pmb{\mu}_l \|^2} &= \sum_{i|t_i \in]0, \frac{1}{2}[} t_i^2 + \sum_{i|t_i \in]\frac{1}{2}, 1[} (1 - t_i)^2, \\ \frac{B_{kl} - I_{kl}}{(n_k + n_l) \| \pmb{\mu}_k - \pmb{\mu}_l \|^2} &= \frac{\sum_{\pmb{x}_i \in \mathcal{V}_k} \langle \pmb{x}_i - \pmb{\mu}_k | \pmb{\mu}_l - \pmb{\mu}_k \rangle_+^2 + \sum_{\pmb{x}_i \in \mathcal{V}_l} \langle \pmb{x}_i - \pmb{\mu}_l | \pmb{\mu}_k - \pmb{\mu}_l \rangle_+^2}{(n_k + n_l) \| \pmb{\mu}_k - \pmb{\mu}_l \|^4}. \end{split}$$

6.2 Code

6.2.1 Implementation

Numerical experiments have been conducted in Python. The python scripts to reproduce the simulations and figures are available at https://github.com/flheight/Spectral-Bridges. The Spectral Bridge algorithm is implemented both in

- Python: https://pypi.org/project/spectral-bridges, and
- R: https://github.com/cambroise/spectral-bridges-Rpackage.

6.2.2 Affinity matrix computation

Taking a closer look at the second step of Algorithm 1 , that is the affinity matrix calculation with a $O(n \times m \times d)$ time complexity, most operations can be parallelized leaving a single loop, bundling together m^2 dot products into only m matrix multiplications, thus allowing for an efficient construction in both high and low level programming languages. Though the complexity of the algorithm remains unchanged, libraries such as Basic Linear Algebra Subprograms can render the calculations orders of magnitude faster. Moreover, the symmetrical nature of the bridge affinity can be used to effectively halve the computation time.

The calculation of the affinity matrix is highlighted by the Python code Listing 1. Though it could be even more optimized, the following code snippet is approximately 200 times faster than a naive implementation on a small dataset comprised of n = 3594, d = 2 points, and a value of m = 250.

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Session information

R version 4.4.1 (2024-06-14)
Platform: x86_64-pc-linux-gnu
Running under: Ubuntu 22.04.5 LTS

Matrix products: default

BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.10.0 LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0

locale:

[1] LC_CTYPE=C.UTF-8 LC_NUMERIC=C LC_TIME=C.UTF-8 [4] LC_COLLATE=C.UTF-8 LC_MONETARY=C.UTF-8 LC_MESSAGES=C.UTF-8

[7] LC_PAPER=C.UTF-8 LC_NAME=C LC_ADDRESS=C

[10] LC_TELEPHONE=C LC_MEASUREMENT=C.UTF-8 LC_IDENTIFICATION=C

time zone: Etc/UTC

tzcode source: system (glibc)

attached base packages:

[1] stats graphics grDevices datasets utils methods base

loaded via a namespace (and not attached):

[1] compiler_4.4.1 fastmap_1.1.1 cli_3.6.2 htmltools_0.5.7 [5] tools_4.4.1 yaml_2.3.8 rmarkdown_2.26 knitr_1.45

[5] tools_4.4.1 yaml_2.3.8 rmarkdown_2.26 knitr_1.45 [9] jsonlite_1.8.8 xfun_0.42 digest_0.6.34 rlang_1.1.3

[13] renv_1.0.7 evaluate_0.23

Listing 1 Python code for affinity matrix computation

```
# Initialize the matrix as empty
affinity = np.empty((self.n_nodes, self.n_nodes))
# Center each region
X_centered = [
    np.array(
        X[kmeans.labels_ == i] - kmeans.cluster_centers_[i],
        dtype=np.float32,
        order="F",
    )
    for i in range(self.n_nodes)
٦
# Cardinal calculation
counts = np.array([X_centered[i].shape[0] for i in range(self.n_nodes)])
counts = counts[None, :] + counts[:, None]
# Calculate the affinity
for i in range(self.n_nodes):
    segments = np.asfortranarray(
        kmeans.cluster_centers_ - kmeans.cluster_centers_[i]
    dists = np.einsum("ij,ij->i", segments, segments)
    dists[i] = 1
    projs = sgemm(1.0, X_centered[i], segments, trans_b=True)
    np.clip(projs / dists, 0, None, out=projs)
    projs = np.power(projs, self.p)
    affinity[i] = projs.sum(axis=0)
affinity = np.power((affinity + affinity.T) / counts, 1 / self.p)
affinity -= 0.5 * affinity.max()
# Scale and exponentiate
q10, q90 = np.quantile(affinity, [0.1, 0.9])
gamma = np.log(self.M) / (q90 - q10)
affinity = np.exp(gamma * affinity)
```