# Skeleton Clustering : A Dimension free Density-Aided Clustering

Kaustav Paul Sourav Biswas

Indian Statistical Institute, Kolkata

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# Traditional Clustering Methods

#### k-means clustering:

- Unable to detect non-convex clusters.
- The center of a non-convex cluster falls outside the cluster itself and may come close to observations from a different cluster.
- In high dimension k-means algorithm may assign all the points to a single cluster.

#### Density Based Clustering:

- To estimate the underlying PDF and detect clusters based on the PDF.
- The rate of convergence for the density estimates is  $\mathcal{O}_{\mathbb{P}}(n^{-\frac{1}{d+4}})$

## • Hierarchical Clustering:

- Problem with non-convex clusters persists.
- If any pair of the points in two different clusters lie very close to each other, the two clusters may get merged in this method.

# Skeleton Clustering Framework.

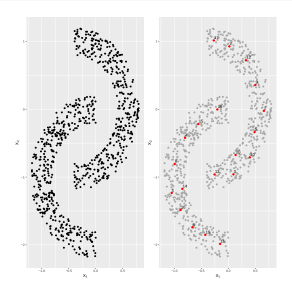
**Input**: Observations  $X_1, X_2, \dots, X_N$ , final number of clusters S.

- **Knot construction**: Perform k—means clustering with a large number k; the centers are the knots.
- **2 Edge construction** : Apply approximate Delaunay triangulation to the knots. Generally we choose  $k = |\sqrt{n}|$
- Edge weights construction: Add weights to each edge using either Voronoi density, Face density or Tube density similarity measure.
- Knots segmentation: Use linkage criterion to segment knots into S groups based on the edge weights.
- **Assignment of labels**: Assign a cluster label to each observation based on which knot group the nearest knot belongs to.

## Knot construction

- Some knots are constructed to give a concise representation of the data structure.
- In practice we use k-Means to choose  $k = \lfloor \sqrt{n} \rfloor$  knots, where n is the number of samples.
- Empirically robustness performance with sufficient number of knots.

## **Knot Construction**

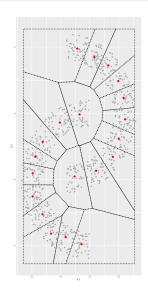


## Edge construction

Let  $c_1, c_2, \ldots, c_k$  be the given knots and we use  $C = \{c_1, c_2, \ldots, c_k\}$  to denote their collection of them.

• The Voronoi cell, or Voronoi region,  $\mathbb{C}_j$  associated with a knot  $c_j$  is the set of all points in  $\mathcal{X}$  whose distance to  $c_j$  is the smallest compared to other knots. That is,  $\mathbb{C}_j = \{ \mathbf{x} \in \mathcal{X} : d(\mathbf{x}, c_j) \leq d(\mathbf{x}, c_\ell) \, \forall \ell \neq j \}$  where  $d(\mathbf{x}, \mathbf{y})$  is the usual Euclidean distance.

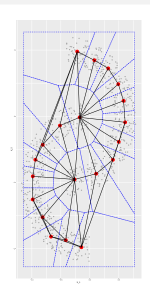
# **Edge Construction**



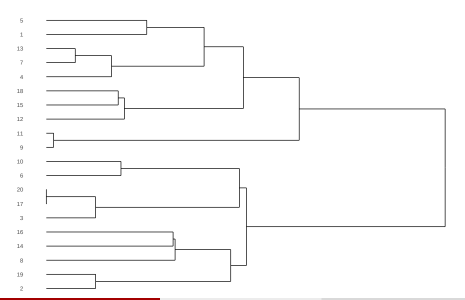
## **Edge Construction**

- We add an edge between a pair of knots if they are neighbors, with the neighboring condition being that the corresponding Voronoi cells share a common boundary.
- Such resulting graph is the Delaunay Triangulation of the set of knots  $\mathcal C$  and we denote is as  $DT(\mathcal C)$ .
- But in case of high dimensional data, it becomes computationally expensive. Therefore, in practice we approximate the exact Delaunay Triangulation with  $\widehat{DT}(\mathcal{C})$  by examining the 2-nearest knots of the sample data points.

# **Edge Construction**



# **Knot Segmentation**



# Final Clustering Result

