# Quantum Transport Clustering

The package quantum\_transport\_clustering contains three major class objects:

- GraphMethods: construct undirected graphs, and compute and encapsulate their graph Laplacians
- SpectralClustering: perform correct spectral clustering on undirected graph Laplacians
- QuantumTransportClustering: perform quantum transport clustering on undirected graph Laplacians

#### Usage example

```
import quantum_transport_clustering as qtc
graph_ = qtc.GraphMethods(data)
...
spec = qtc.SpectralClustering(n_clusters=3, norm_method='row')
...
shot = qtc.QuantumTransportClustering(n_clusters=3, Hamiltonian=Lap_)
```

### **Graph Methods**

```
1 quantum_transport_clustering.GraphMethods(data_, graph_embedded=True, edt_tau=None, eps_quant=None, normed=True, compute_lap=True)
```

The Class GraphMethods is able to

- Generate Gaussian RBF adjacency matrix using Euclidean distances of the data distribution
- Compute Graph Lapalcian (symmetrically normalized by default)
- Store the raw data as well as adjacency matrix and graph Laplacian

#### **Parameters**

data_	If graph_embedded = True, data_ is a numby array of shape $(n_{\text{feature}}, m_{\text{sample}})$ , or $m_{\text{sample}}$ points in $\mathbb{R}^{n_{\text{feature}}}$ . If graph_embedded = False, data_ is a numby array of shape $(m_{\text{sample}}, m_{\text{sample}})$ representing the adjacency of a graph with $m_{\text{sample}}$ nodes.
graph_embedded	bool, optional. If True, assume the graph is embedded in a Euclidean space. If False, assume the input data set is an adjacency matrix not <i>a priori</i> embedded in a Euclidean space.
edt_tau	int, $\tau > 0$ , optional. If specified, it is the number of iterations of effective

	dissimilarity transformation (EDT). Neglected if graph_embedded = False.
eps_quant	float, in range $0<\varepsilon<100$ , optional. The the quantile of distance distribution. If not specified, $\varepsilon=1$ . Neglected if graph_embedded = False.
normed	bool, optional. If False, graph Laplacian is $L=D-A$ where $D$ is degree diagonal matrix, and $A$ the adjacency matrix. If True, graph Laplacian will be normalized $H=D^{-\frac{1}{2}}LD^{-\frac{1}{2}}$ .
compute_lap	bool, optional. If True, graph Laplacian will be computed upon initialization.
Returns	

numpy array of shape (  $\emph{m}_{sample}$  ,  $\emph{m}_{sample}$  ). The graph Laplacian matrix  $\emph{L}$  or  $\emph{H}$ . Lap\_

#### Example:

```
graph_ = qtc.GraphMethods(data)
2 laplacian_matrix_ = graph_.Lap_
```

## Spectral Clustering

quantum\_transport\_clustering.SpectralClustering(n\_clusters, norm\_method='row', isExact=True)

Perform correct spectral clustering on undirected graph Laplacians. Requires numpy >= 1.13.

#### **Parameters**

n_clusters	int, $n_{ m cluster} > 0$ , the number of clusters.
norm_method	None, "row", or "deg". If None, the spectral embedding is not normalized. If "row", the spectral embedding is $L^2$ -normalized by row where each row represent a node. If "deg", the spectral embedding is normalized by degree vector.
isExact	bool. If True, exact eigenvalues and eigenvectors will be computed. If False, first (small) $n_{\text{cluster}}$ eigenvalues and eigenvectors will be computed.

#### Methods

Lap\_ is the symmetric graph Laplacian. First, the eigenvalues and eigenstates are fit(Lap\_)

computed. Next, perform spectral embedding and *k*-means.

#### **Returns**

labels\_ An integer-valued numpy array of shape  $(m_{\text{sample}})$ . The class labels associated with each node.

#### Example:

```
spec = qtc.SpectralClustering(n_clusters=3, norm_method='row')
spec.fit(laplacian_matrix)
spec_labels_ = spec.labels_
```

### **Quantum Transport Clustering**

```
quantum_transport_clustering.QuantumTransportClustering(n_clusters, Hamiltonian,
s=1.0, isExact=True, n_eigs = None)
```

Perform quantum transport clustering on undirected graph Laplacians.

#### **Parameters**

n_clusters	int , $n_{ m cluster} > 0$ , the number of clusters.
Hamiltonian	numpy array of shape ( $m_{ m sample}$ , $m_{ m sample}$ ). The symmetric graph Laplacian matrix $H$ .
S	float, $\tilde{s}>0$ , optional. The actual $s$ -parameter of Laplace transform will be $s=\tilde{s}\times(E_{n_{\mathrm{cluster}}-1}-E_0)/(n_{\mathrm{cluster}}-1)$ , where $E_n$ are eigenvalues of $H$ .
isExact	bool, optional. If True, exact eigenvalues and eigenvectors of $\boldsymbol{H}$ will be computed. If False, first n_eigs low energy states will be computed approximated.
n_eigs	int, $n_{\rm eigs} > 0$ , optional. If n_eigs not specified and isExact = False, $n_{\rm eigs} = 10 \times n_{\rm cluster}$ . If n_eigs is specified and isExact = True, then first $n_{\rm eigs}$ low exact energy state will be used to perform quantum transport clustering. The latter case can be used to speed up the clustering processes.

#### Methods

```
Grind() Grind(s=None, grind='medium', method_='diff', init_nodes_=None) Option grind can be "coarse", "medium", "fine", "micro", or custom. Option method can be diff or
```

	kmeans corresponding to direct difference and $k$ -means methods. If grind="custom", then init_nodes_ is the custom python list of initialization nodes. Method Grind() produces the array Omega_ or the $\Omega$ -matrix which contains the raw class labels.
Espresso()	Perform "direct extraction method" on $\Omega$ . This method creates attribute labels_ as the predicted class labels.
Coldbrew()	Compute "consensus matrix" $C$ based on $\Omega$ . This method creates attribute

#### Returns

Omega_	An integer-valued numpy array of shape $(m_{\text{sample}}, m_{\text{initialization}})$ . The raw class labels of $m_{\text{samples}}$ from quantum transport from $m_{\text{initialization}}$ nodes.
labels_	An integer-valued numpy array of shape $(m_{\text{sample}})$ . The final prediction by Espresso().
consensus_matrix_	A float-valued numpy array of shape $(m_{\rm sample}, m_{\rm sample})$ . The consensus matrix computed by Coldbrew().

### Example:

```
shot = qtc.QuantumTransportClustering(n_clusters=3, Hamiltonian=Lap_) #
initialization

Omg_ = shot.Grind() # generate raw class label

# One may extract the eigevalues by attribute shot.Heigval

shot.Espresso() # direct extraction method

class_labels_ = shot.labels_

shot.Coldbrew() # generate consensus matrix

C_matrix_ = shot.consensus_matrix_
```