14. Clustering

Motivation

- Supervised learning is good for interaction with humans, but labels from a supervisor are hard to obtain
- Clustering is unsupervised learning, i.e. it tries to lear only from the data
- Main idea: find a similarity measure and group similar data objects together
- Clustering is a very old research field, many approaches have been suggested
- Main problem in most methods: how to find a good number of clusters





The full posterior of the Gaussian Mixture Model is

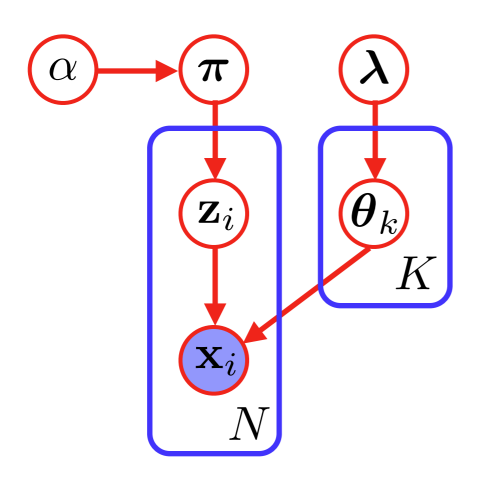
$$p(X, Z, \boldsymbol{\mu}, \Sigma, \boldsymbol{\pi}) = p(X \mid Z, \boldsymbol{\mu}, \Sigma) p(Z \mid \boldsymbol{\pi}) p(\boldsymbol{\pi} \mid \alpha) p(\boldsymbol{\mu}, \Sigma \mid \boldsymbol{\lambda})$$

data likelihood (Gaussian)

correspondence prob. (Multinomial)

mixture prior (Dirichlet)

parameter prior (Gauss-IW)



In this model, we use:

$$ullet$$
 $oldsymbol{\mu}=(oldsymbol{\mu}_1,\ldots,oldsymbol{\mu}_K)$

•
$$\Sigma = (\Sigma_1, \dots, \Sigma_K)$$

•
$$(\boldsymbol{\mu}_k, \Sigma_k) = \boldsymbol{\theta}_k$$

Simplification for now:

- Assume Σ_k are known
- Thus: $\theta_k = \mu_k$



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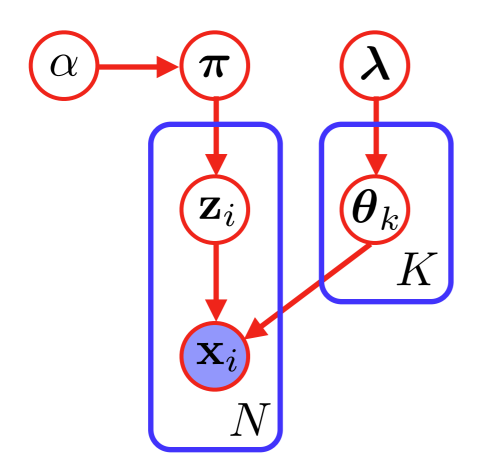
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Given this model, we can create new samples:

- 1. Sample π , θ_k from priors
- 2. Sample corresp. z_i
- 3. Sample data point x_i



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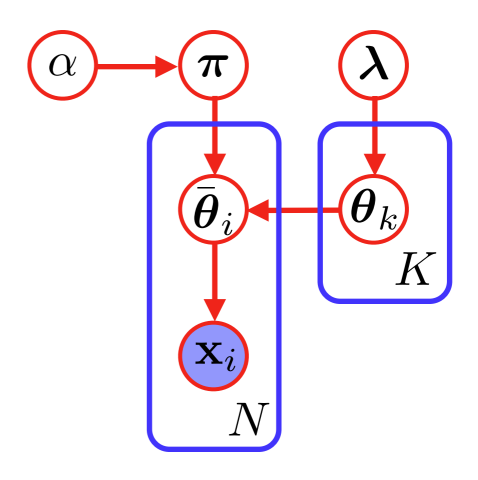
parameter prior (Gauss-IW)

An equivalent formulation of this model is this:

- 1. Sample π , θ_k from priors
- 2. Sample params $\bar{\theta}_i$ from:

$$p(\bar{\boldsymbol{\theta}}_i \mid \boldsymbol{\pi}, \boldsymbol{\theta}_k) = \sum_{k=1}^K \pi_k \delta(\boldsymbol{\theta}_k, \bar{\boldsymbol{\theta}}_i)$$

3. Sample data point x_i





What is the difference in that model?

- ullet there is one parameter $ar{ heta}_i$ for each observation \mathbf{x}_i
- intuitively: we first sample the location of the cluster and then the data that corresponds to it

In general, we use the notation:

$$m{\pi} \sim ext{Dir}(rac{lpha}{K}\mathbf{1})$$
 $m{ heta}_k \sim ext{H}(m{\lambda})$ "Base distribution"
 $ar{m{ heta}}_i \sim ext{G}(m{\pi}, m{ heta}_k)$ where $G(m{\pi}, m{ heta}_k) = \sum_{K} \pi_k \delta(m{ heta}_k, ar{m{ heta}}_i)$

However: We need to know K



The Dirichlet Process

- So far, we assumed that K is known
- To extend that to infinity, we use a trick:

Definition: A Dirichlet process (DP) is a distribution over probability measures G, i.e. $G(\theta) \ge 0$ and

$$\int G(\theta)d\theta = 1$$
. If for any partition (T_1, \dots, T_K) it holds:

$$(G(T_1),\ldots,G(T_K)) \sim \text{Dir}(\alpha H(T_1),\ldots,\alpha H(T_K))$$

then *G* is sampled from a Dirichlet process.

Notation: $G \sim \mathrm{DP}(\alpha, H)$

where α is the concentration parameter and H is the base measure





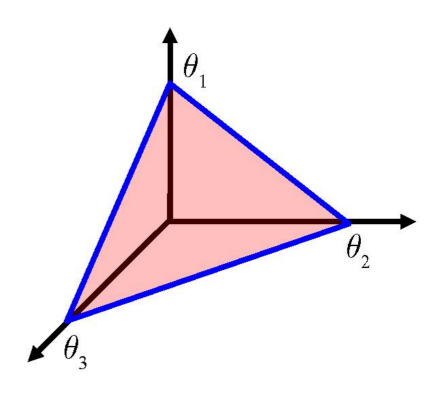
Repetition

The Dirichlet distribution is defined as:

$$\operatorname{Dir}(\boldsymbol{\mu} \mid \boldsymbol{\alpha}) = \frac{\Gamma(\alpha_0)}{\Gamma(\alpha_1) \cdots \Gamma(\alpha_K)} \prod_{k=1}^K \mu_k^{\alpha_k - 1} \qquad \alpha_0 = \sum_{k=1}^K \alpha_k$$

$$0 \le \mu_k \le 1 \qquad \sum_{k=1}^K \mu_k = 1$$

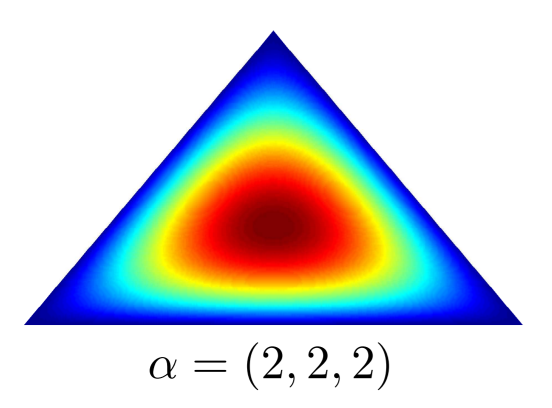
- It is the conjugate prior for the multinomial distribution
- There, the parameter α can be interpreted as the effective number of observations for every state



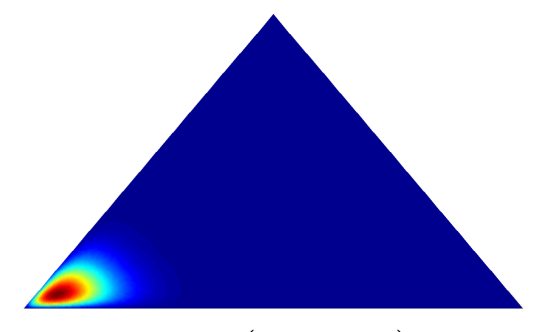
The simplex for K=3



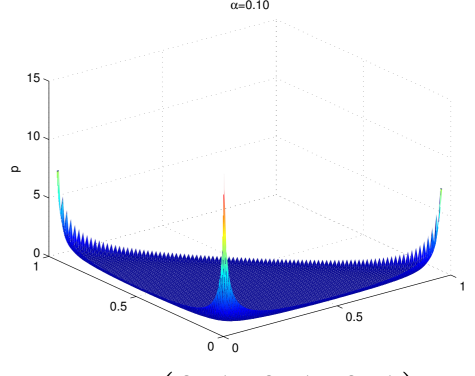
Some Examples



- α_0 controls the strength of the distribution ("peakedness")
- α_k control the location of the peak



$$\alpha = (20, 2, 2)$$



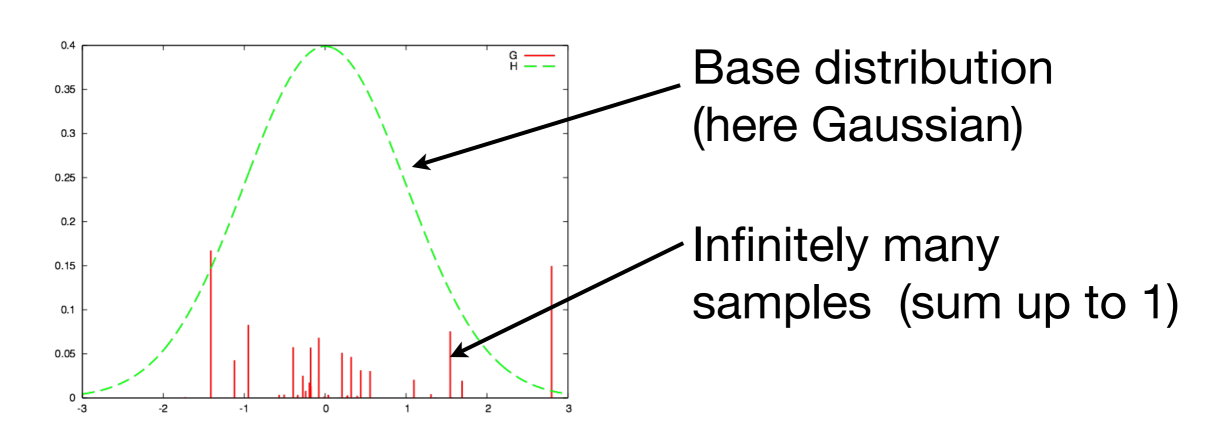
$$\alpha = (0.1, 0.1, 0.1)$$





Intuitive Interpretation

- Every sample from a Dirichlet distribution is a vector of K positive values that sum up to 1, i.e. the sample itself is a finite distribution
- Accordingly, a sample from a Dirichlet process is an infinite (but still discrete!) distribution



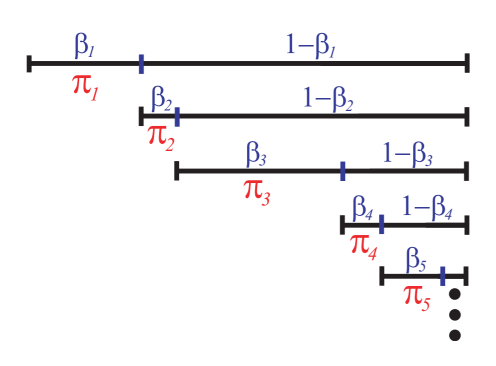
Construction of a Dirichlet Process

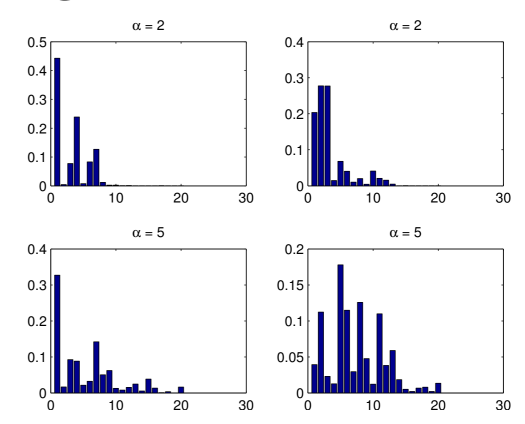
- The Dirichlet process is only defined implicitly, i.e.
 we can test whether a given probability measure is
 sampled from a DP, but we can not yet construct
 one.
- A DP can be constructed using the "stickbreaking" analogy:
 - imagine a stick of length 1
 - we select a random number β between 0 and 1 from a Beta-distribution
 - we break the stick at $\pi = \beta$ * length-of-stick
 - we repeat this infinitely often





The Stick-Breaking Construction





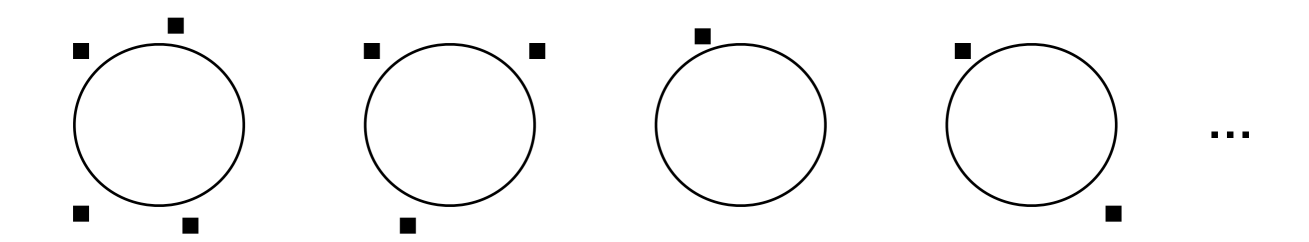
formally, we have

$$eta_k \sim \mathrm{Beta}(1, lpha)$$
 $\pi_k = eta_k \prod_{l=1}^{k-1} (1 - eta_l) = eta_k (1 - \sum_{l=1}^{k-1} \pi_l)$

now we define

$$G(\boldsymbol{\theta}) = \sum_{k=1}^{\infty} \pi_k \delta(\boldsymbol{\theta}_k, \boldsymbol{\theta})$$
 $\boldsymbol{\theta}_k \sim H$ then: $G \sim \mathrm{DP}(\alpha, H)$

The Chinese Restaurant Process



- Consider a restaurant with infinitely many tables
- Everytime a new customer comes in, he sits at an occupied table with probability proportional to the number of people sitting at that table, but he may choose to sit on a new table with decreasing probability as more customers enter the room.



The Chinese Restaurant Process

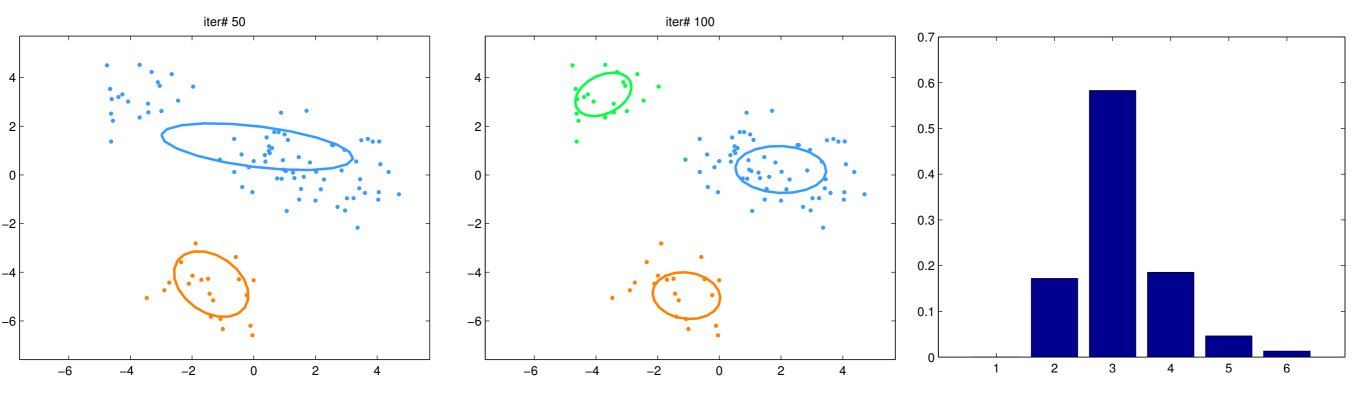
 It can be shown that the probability for a new customer is

$$p(\bar{\boldsymbol{\theta}}_{N+1} = \boldsymbol{\theta} \mid \bar{\boldsymbol{\theta}}_{1:N}, \alpha, H) = \frac{1}{\alpha + N} \left(\alpha H(\boldsymbol{\theta}) + \sum_{k=1}^{K} N_k \delta(\bar{\boldsymbol{\theta}}_k, \boldsymbol{\theta}) \right)$$

- This means that currently occupied tables are more likely to get new customers (rich get richer)
- The number of occupied tables grows logarithmically with the number of customers

The DP for Mixture Modeling

- Using the stick-breaking construction, we see that we can extend the mixture model clustering to the situation where K goes to infinity
- The algorithm can be implemented using Gibbs sampling



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- Often, we are only given a similarity matrix for the data points
- The idea of Affinity Propagation is to determine cluster centers ("exemplars") that explain other data points in an optimal way
- This is similar to k-medoids, but the algorithm is more robust against local minima
- Idea: each data point must choose another data point as its exemplar; some points will choose themselves as exemplar
- The number of clusters is then found automatically





- Input: similarity values s(i,j)
- Initialize the responsibilities r(i,j), and the availabilities a(i,j) to 0
- do until convergence:
 - recompute the responsibilities:

$$r(i,j) = s(i,j) - \max_{j' \neq j} \{a(i,j') + s(i,j')\}$$

recompute the availabilities:

$$a(i,j) = \min \left\{ 0, r(j,j) + \sum_{i' \notin \{i,j\}} \max\{0, r(i',j)\} \right\}$$

• the j that maximizes r(i,j) + a(i,j) is the exemplar of i

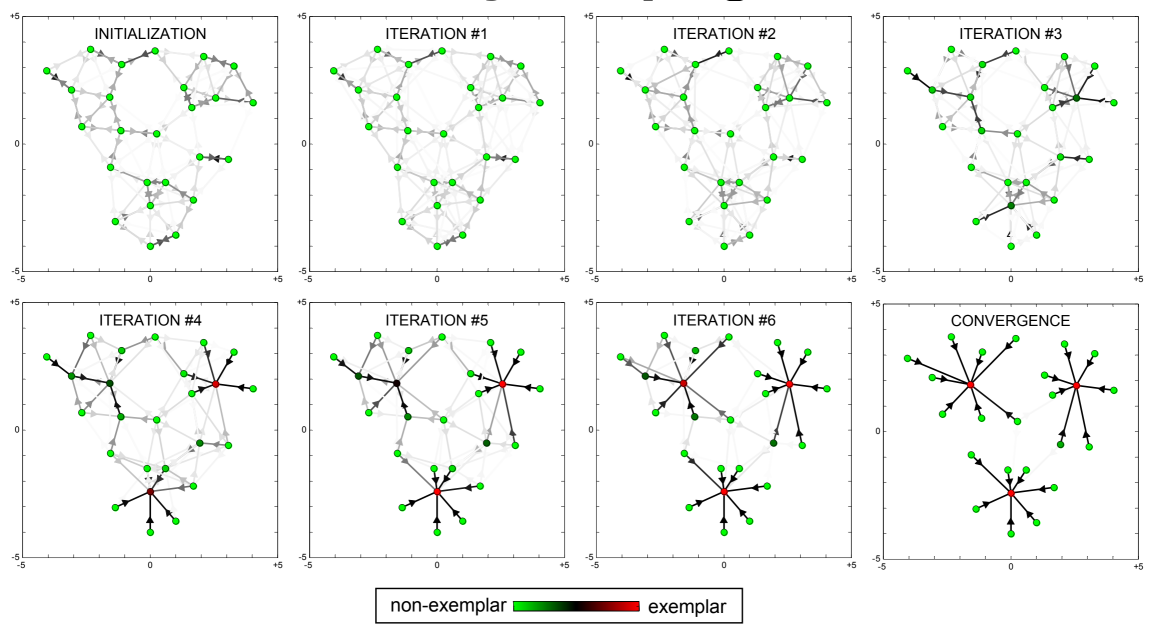




- Intuitively:
 - responsibility measures how much i thinks that j would be a good exemplar
 - availability measures how strongly j things it should be an exemplar for i
- The algorithm can be shown to be equivalent to max-product loopy belief propagation
- Convergence is not guaranteed, but with "damping" oscillations can be avoided
- The number of clusters can be controlled by the "self-similarity"





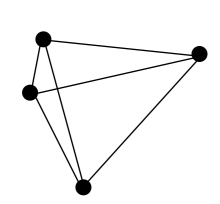


- Colours: how much each point wants to be an exemplar
- Edge strengths: how much a point wants to belong to a cluster

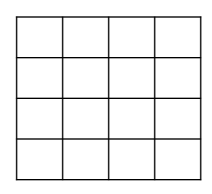




- Consider an undirected graph that connects all data points
- The edge weights are the similarities ("closeness")
- We define the weighted degree d_i of a node as the sum of all outgoing edges

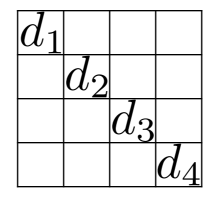


W =



$$d_i = \sum_{j=1}^{N} w_{ij}$$

$$D =$$



• The Graph Laplacian is defined as:

$$L = D - W$$

- This matrix has the following properties:
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- This matrix has the following properties:
 - the 1 vector is eigenvector with eigenvector 0
 - the matrix is symmetric and positive semi-definite
- With these properties we can show:

Theorem: The set of eigenvectors of L with eigenvalue 0 is spanned by the indicator vectors $\mathbf{1}_{A_1}, \ldots, \mathbf{1}_{A_K}$, where A_k are the K connected components of the graph.

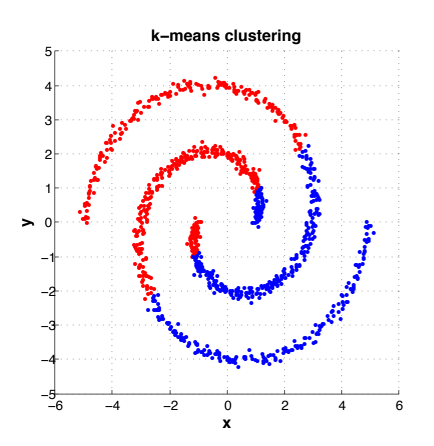


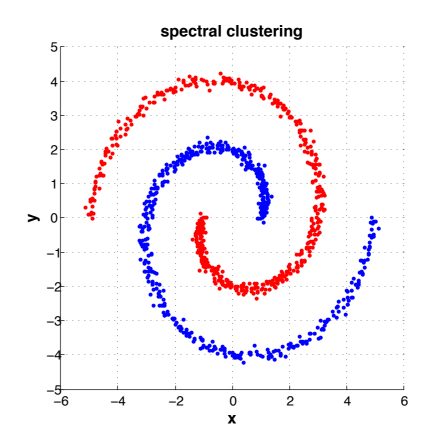
The Algorithm

- Input: Similarity matrix W
- Compute L = D W
- Compute the eigenvectors that correspond to the K smallest eigenvalues
- Stack these vectors as rows in a matrix U
- Treat each row of U as a K-dim data point
- Cluster the N rows with K-means clustering
- The indices of the rows that correspond to the resulting clusters are those of the original data points.



An Example





- Spectral clustering can handle complex problems such as this one
- The complexity of the algorithm is O(N³), because it has to solve an eigenvector problem
- But there are efficient variants of the algorithm

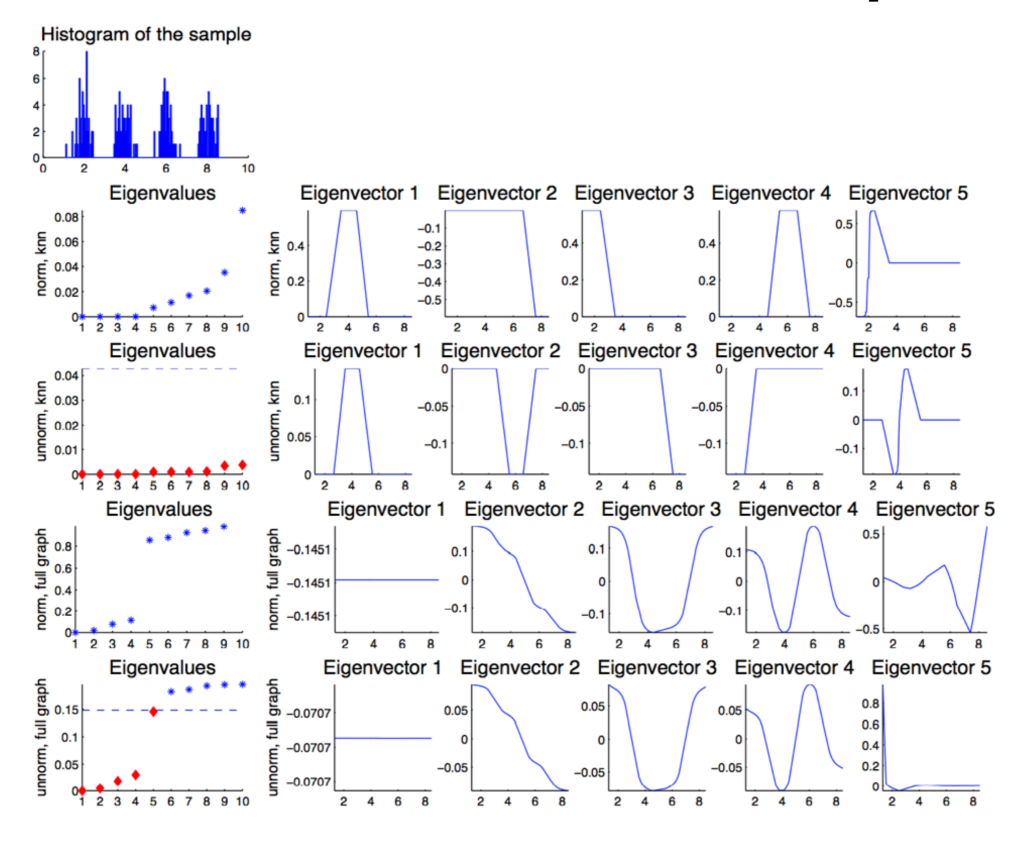


Further Remarks

- To account for nodes that are highly connected, we can use a normalized version of the graph Laplacian
- Two different methods exist:
 - $L_{rw} = D^{-1}L = I D^{-1}W$
 - $L_{sym} = D^{-\frac{1}{2}}LD^{-\frac{1}{2}} = I D^{-\frac{1}{2}}WD^{-\frac{1}{2}}$
- These have similar eigenspaces than the original Laplacian L
- Clustering results tend to be better than with the unnormalized Laplacian



Another Small Example





Hierarchical Clustering

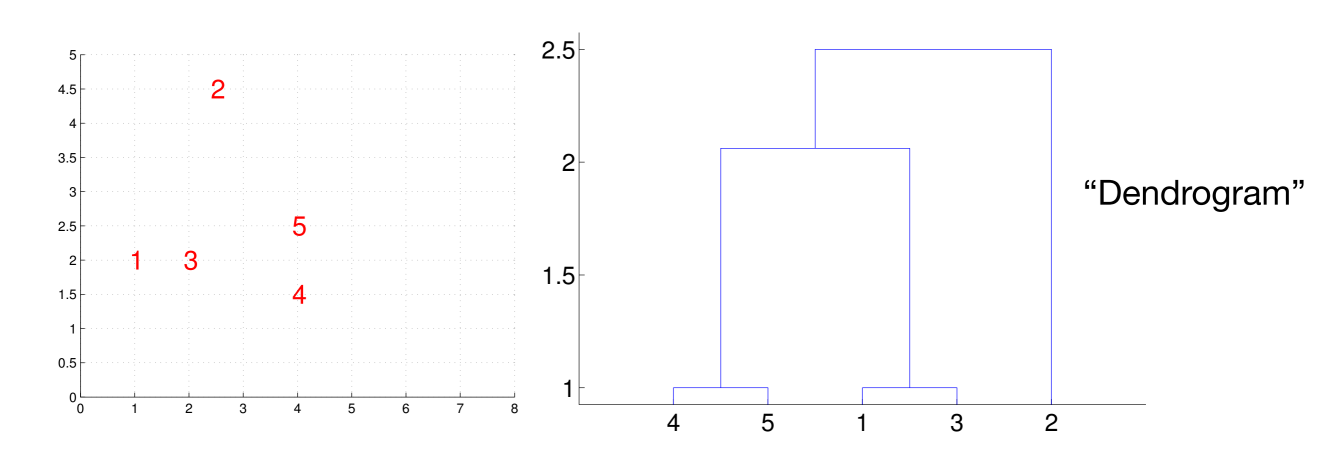
- Often, we want to have nested clusters instead of a "flat" clustering
- Two possible methods:
 - "bottom-up" or agglomerative clustering
 - "top-down" or divisive clustering
- Both methods take a dissimilarity matrix as input
- Bottom-up grows merges points to clusters
- Top-down splits clusters into sub-clusters
- Both are heuristics, there is no clear objective function
- They always produce a clustering (also for noise)





Agglomerative Clustering

- Start with N clusters, each contains exactly one data point
- At each step, merge the two most similar groups
- Repeat until there is a single group





Linkage

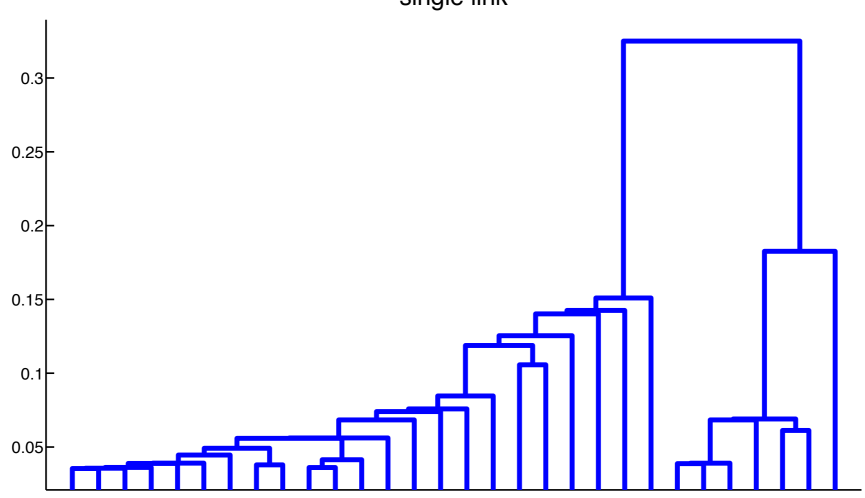
- In agglomerative clustering, it is important to define a distance measure between two clusters
- There are three different methods:
 - Single linkage: considers the two closest elements from both clusters and uses their distance
 - Complete linkage: considers the two farthest elements from both clusters
 - Average linkage: uses the average distance between pairs of points from both clusters
- Depending on the application, one linkage should be preferred over the other





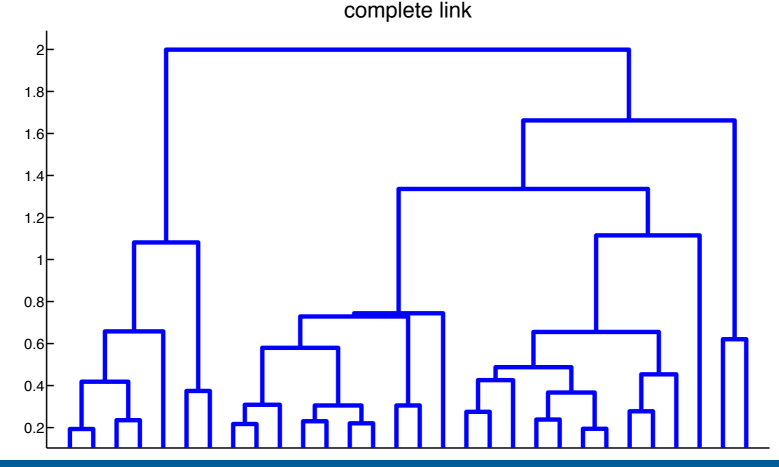
Single Linkage

- The distance is based on $d_{SL}(G,H) = \min_{i \in G, i' \in H} d_{i,i'}$
- The resulting dendrogram is a minimum spanning tree, i.e. it minimizes the sum of the edge weights
- Thus: we can compute the clustering in O(N²) time



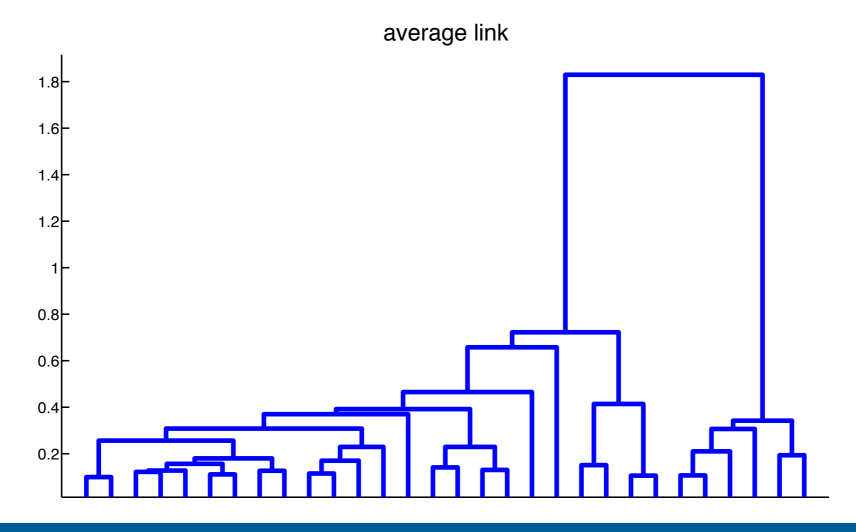
Complete Linkage

- The distance is based on $d_{CL}(G,H) = \max_{i \in G, i' \in H} d_{i,i'}$
- Complete linkage fulfills the compactness property, i.e. all points in a group should be similar to each other
- Tends to produce clusters with smaller diameter



Average Linkage

- The distance is based on $d_{avg}(G,H)=\frac{1}{n_Gn_H}\sum_{i\in G}\sum_{i'\in H}d_{i,i'}$ Is a good compromise between single and
- complete linkage
- However: sensitive to changes on the meas. scale



Divisive Clustering

- Start with all data in a single cluster
- Recursively divide each cluster into two child clusters
- Problem: optimal split is hard to find
- Idea: use the cluster with the largest diameter and use K-means with K = 2
- Or: use minimum-spanning tree and cut links with the largest dissimilarity
- In general two advantages:
 - Can be faster
 - More globally informed (not myopic as bottom-up)





Choosing the Number of Clusters

- As in general, choosing the number of clusters is hard
- When a dendrogram is available, a gap can be detected in the lengths of the links
- This represents the dissimilarity between merged groups
- However: in real data this can be hard to detect
- There are Bayesian techniques to address this problem (Bayesian hierarchical clustering)

Evaluation of Clustering Algorithms

- Clustering is unsupervised: evaluation of the output is hard, because no ground truth is given
- Intuitively, points in a cluster should be similar and points in different clusters dissimilar
- However, better methods use external information, such as labels or a reference clustering
- Then we can compare clusterings with the labels using different metrics, e.g.
 - purity
 - mutual information

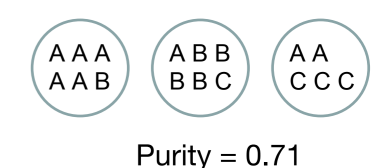


Purity

- Define N_{ij} the number of objects in cluster i that are in class j $_{C}$
- Define $N_i = \sum N_{ij}$ number of objects in cluster i

•
$$p_{ij} = \frac{N_{ij}}{N_i}$$
 $p_i = \max_j p_{ij}$ "Purity"

• overall purity
$$\sum_{i}^{N_{i}} \frac{N_{i}}{N} p_{i}$$



- Purity ranges from 0 (bad) to 1 (good)
- But: a clustering with each object in its own cluster has a purity of 1

Mutual Information

- Let U and V be two clusterings
- Define the probability that a randomly chosen point belongs to cluster u_i in U and to v_i in V

$$p_{UV}(i,j) = \frac{|u_i \cap v_j|}{N}$$

• Also: The prob. that a point is in u_i $p_U(i) = \frac{|u_i|}{N}$ $\mathbb{I}(U,V) = \sum_{i=1}^R \sum_{j=1}^C p_{UV}(i,j) \log \frac{p_{UV}(i,j)}{p_U(i)p_V(j)}$

$$\mathbb{I}(U, V) = \sum_{i=1}^{n} \sum_{j=1}^{C} p_{UV}(i, j) \log \frac{p_{UV}(i, j)}{p_{U}(i) p_{V}(j)}$$

 This can be normalized to account for many small clusters with low entropy





Summary

- Several Clustering methods:
 - Dirichlet process mixture model does not require the number of clusters to be known; full Bayesian
 - Affinity Propagation: iterative approach where exemplars are determined as cluster centers
 - Spectral clustering uses the graph Laplacian and performs an eigenvector analysis
 - Hierarchical approaches can be bottom-up or topdown
- Evaluation methods for Clustering are hard to find
- Some are based on purity or mutual information



