Lecture 20: Unsupervised learning I - Clustering

Isabel Valera

Machine Learning Group
Department of Mathematics and Computer Science
Saarland University, Saarbrücken, Germany

28.06.2021

Bibliography ●O

- Bibliography
- 2 Introduction
- K-means
- 4 Hierarchical clustering
- Density-based clustering

Main references

- Bishop Chapter 9 & 12
- EML Chapter 14

- Bibliography
- 2 Introduction
- K-means
- 4 Hierarchical clustering
- Density-based clustering

Unsupervised learning

Unsupervised Learning:

Given a set of input points $(X_i)_{i=1}^n$:

- Clustering: Construction of a grouping of the points into sets of similar points, the so called clusters. (Today!)
- Density Estimation: Estimation of the distribution of the input points over the input space X. Related to outlier detection.
 (Partially today!)
- Dimensionality Reduction: Construction of a mapping $\phi: \mathcal{X} \to \mathbb{R}^m$, where the dimensionality m of the target space is usually much smaller than that of the input space \mathcal{X} . Generally, the mapping should preserve properties of the input space \mathcal{X} , e.g., distances.

Clustering

Clustering approaches aim at grouping of the observed data into sets of *similar* points, the so called *clusters*. In general,

- there is not a broadly accepted objective for clustering without specifying a suitable objective, clustering is ill-defined,
- clustering objective depends usually on application,
- in clustering the modeling aspect is even more important than in supervised learning – thus, do not use a clustering method if you do not understand what the objective implies!

- Bibliography
- 2 Introduction
- K-means
- 4 Hierarchical clustering
- Density-based clustering

K-means

K-means clustering is a prototype based clustering approach:

- Goal: find prototypes (centroids) μ_i , i = 1, ..., k which represent the data in an optimal way (what does that mean?),
- **Objective:** denote by C_i the *i*-th cluster (set of points) which is represented by the prototype μ_i ,

$$\underset{(C_1,\boldsymbol{\mu}_1),...,(C_k,\boldsymbol{\mu}_k)}{\arg\min} \sum_{i=1}^k \sum_{\mathbf{x}_i \in C_i} \|\mathbf{x}_j - \boldsymbol{\mu}_i\|^2,$$

where $\|\cdot\|$ is the Euclidean norm,

- True Goal:
 - 1 finds sphere-like clusters in the data,
 - heavily influenced by outliers,
 - non-sphere like clusters are hard to fit.

K-means algorithm

K-means clustering:

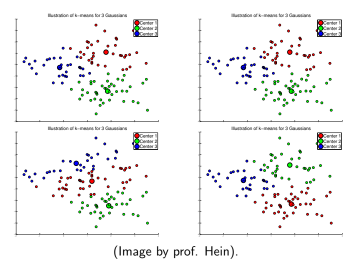
- k-means is combinatorial optimization problem,
- simple iterative algorithm converges fast but finds only local minimum.

Lloyd's algorithm for k-means clustering:

- lacktriangle initialize centers μ_i ,
- **4 while** changing μ_i , $i=1,\ldots,k$ (i.e, iterate until clusters stop changing),
 - group all samples according to closest μ_i , i = 1, ..., k
 - ② recompute μ_i as the mean of the observations in cluster C_i for i = 1, ..., k
- \bullet return μ_1,\ldots,μ_k ,

Steps are optimal for fixed centroids

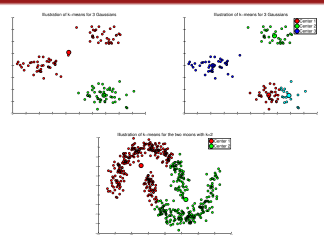
Example I



Four independent runs for k-means algorithm for the same dataset



Example II



Left: k is chosen too small. Middle: k is chosen too large. Right: The two moons dataset - clusters are not of spherical shape (thus, decreasing k not useful to find true k). (Image by Prof. Hein)



- Bibliography
- 2 Introduction
- 3 K-means
- 4 Hierarchical clustering
- Density-based clustering

Hierarchical clustering

Hierarchical clustering generates a hierarchical grouping of the n data points. Two approaches:

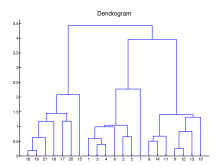
- agglomerative: start with all n points as individual clusters and consecutively join cluster which are most similar;
- divisive: start with one cluster containing all n points and consecutively divide the clusters so that they are most dissimilar.

Generates a tree structure on the data – the **dendrogram**.

Dendrogram

Definition

A **dendrogram** is a binary tree with a distinguished root, that has the data points as its leaves. The height where two clusters are merged is equal to their dissimilarity.



Agglomerative hierarchical clustering

Requirement: a distance measure between point sets.

Definition

A dissimilarity measure D between finite subsets of X is defined as $D: 2^{\mathcal{X}} \times 2^{\mathcal{X}} \to \mathbb{R}$ with

- $D(A,B) \geq 0$ for all $A,B \subseteq \mathcal{X}$,
- D(A, B) = 0 if and only if A = B,
- D(A, B) = D(B, A).

Note: triangle inequality not required – not necessarily a metric.

Algorithm:

- Input: set of n points in \mathcal{X} , dissimilarity D between subsets of \mathcal{X} .
- Initialize with n clusters at level n, $C_1^{(n)}, \ldots, C_n^{(n)}$ with $C_i^{(n)} = \{\mathbf{x}_i\}$.
- while *l* > 1. do
 - compute for all I clusters in $C_1^{(I)}, \ldots, C_L^{(I)}$ their dissimilarity $d_{ii} = D(C_i^{(I)}, C_i^{(I)})$
 - @ merge the least dissimilar clusters, with indices $(r,s) = \underset{1 \le i,j \le l, \ i \ne j}{\arg \min} d_{ij}$.
 - **3** for $i \neq r$ and $i \neq s$, $C_i^{(l-1)} = C_i^{(l)}$ and $C_r^{(l-1)} = C_r^{(l)} \cup C_s^{(l)}$
 - **4** height in the dendrogram of the merger between $C_r^{(l)}$ and $C_r^{(l)}$ is

$$\alpha^{(I)} = d_{rs} = \min_{i,j} d_{ij}.$$

- \bullet relabel the clusters of level l-1 from 1 to l-1,
- Output: the sets of clusters $C^{(l)}$ for each level $l=1,\ldots,n$.

Similarity between clusters

Agglomerative clustering: iteratively join *most similar* clusters.

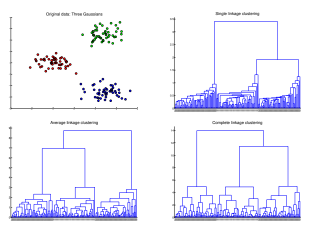
How to measure dissimilarity of clusters C_1 and C_2 ?

- Single-linkage: $d_{\min}(C_1, C_2) = \min_{i \in C_1, j \in C_2} d(\mathbf{x}_i, \mathbf{x}_j)$,
- Complete-linkage: $d_{\max}(C_1, C_2) = \max_{i \in C_1, j \in C_2} d(\mathbf{x}_i, \mathbf{x}_j)$,
- Average-linkage: $d_{\text{avg}}(C_1, C_2) = \frac{1}{|C_1||C_2|} \sum_{i \in C_1, j \in C_2} d(\mathbf{x}_i, \mathbf{x}_j),$

Two clusters are similar:

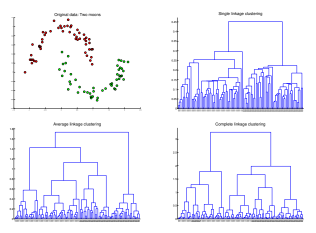
- single linkage: if for all points in each cluster there exists a path so that all points in the path are similar;
- complete-linkage: if all points for both clusters are similar; or
- average-linkage: if on average the points of both clusters are similar.

Example I



Top, left: dataset are three Gaussians. Top, right: single-linkage clustering. Bottom, left: average-linkage clustering. Bottom, right: complete-linkage clustering. (Image by Prof. Hein)

Example II



Top, left: Two moons dataset. Top, right: single-linkage clustering Bottom, left: average-linkage clustering. Bottom, right: complete-linkage clustering. (Image by Prof. Hein)



Problems of dendrograms

Problems of dendrograms

- **instability** small changes in the data can lead to huge changes in the dendrogram,
- hierarchy: multi-scale partitioning but different distance measures are hard to interpret.
- dissimilarity: the dissimilarity of clusters at which one joins clusters encodes their dissimilarity – comparing data using this distance is highly non-intuitive.

Overview

Pros:

- nice hierarchical representation of the data,
- single-linkage has a nice theoretical foundation,
- computationally relatively cheap.

Cons:

- single-linkage and complete very sensitive to data fluctuations,
- complete linkage has problems with non-spherical clusters,
- interpretation of the data requires profound understanding of the cluster similarity measures.

- Bibliography
- 2 Introduction
- 3 K-means
- 4 Hierarchical clustering
- Density-based clustering

Density-based clustering

Probabilistic setting:

- sample $\{\mathbf{x}_i\}_{i=1}^n$ is drawn i.i.d. from probability measure in \mathbb{R}^d ,
- the probability measure has a density in \mathbb{R}^d ,

Clustering model: The simplest approach is to assume that the density *p* is a *Gaussian mixture*, i.e.,

$$ho(\mathbf{x}) = \sum_{k=1}^K \pi_k \mathcal{N}(\mathbf{x}|\boldsymbol{\mu}_k, \boldsymbol{\Sigma}_k),$$

where π_k is teh prior probability of cluster k.

Main difference to approaches up to now is that we have clusters and "background noise".

- The Gaussian mixture model (GMM) assumes that the observed data is an i.i.d. sample from a Gaussian mixture distribution.
- Given observed data $\{\mathbf{x}_i\}_{i=1}^n$, we can get the ML (as well as the MAP) estimate the GMM parameters (i.e., $\{\pi_k, \mu_k, \Sigma_k\}_{k=1}^K$) using the expectation-maximization (EM) algorithm.
- To perfom **model selection** (i.e., select number of components/clusters), one may rely to cross-validation using, e.g., *Bayessian Information Criterion (BIC)* as criteria.
- More details in additional notes and Jupyter notebook.

Observations

- Note that GMMs are an approach for density estimation, that in turn allows us to perform clustering. We, however, need to make the explicit assumption of the likelihood (which is equivalent to select the dismilarity measure for k-means or hierarchical clustering)
- Mixture models can be generalized beyond the Gaussian distribution to accommodate, e.g., categorical or binary data.
- EM algorithm is a general and powerful iterative method to find (local) maximum likelihood or maximum a posteriori (MAP) estimates of parameters in probabilistic models, where the model depends on unobserved latent variables.