# Unsupervised Learning: Clustering

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## Outline

#### Introduction

Principal Component Analysis

Model-based clustering: EM algorithm for Gaussian Mixtures

k-means, k-medoids and variants

(Agglomerative) Hierarchical Cluster Analysis

Other methods

# Clustering

#### Goal

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Goal: automatically discover clusters in the data

- Model-based clustering:
  - assume that the data was generated by a model
  - try to recover the original model from the data
- Model-free clustering:
  - no assumption on the mechanism producing data
  - vector quantization
  - cluster should be homogeneous and different from one another
  - data-driven loss function to minimize

Introduction

# Model-based clustering: choice of distance

- Data often in  $\mathbb{R}^p$  (or projected to)
- Distance sometimes natural, sometimes not
- Often: need to normalize first
- Default choice: Euclidian distance  $d(x,x') = \sqrt{\sum (x_i x_i')^2}$
- Other possible norms:  $L^1, L^{\infty}$ , etc.
- Mahalanobis distance:  $d(x, x') = \sqrt{(x x')\Sigma^{-1}(x x')}$
- Categorial data: typically  $\chi^2$  distance

### Ressources

- The Elements of Statistical Learning, *T. Hastie, R. Friedman, J. Tibshirani*, Springer
- Data Mining , S. Tufféry, Technip
- WikiStat P. Besse et al. http://wikistat.fr/
- Interesting overview at http: //scikit-learn.org/stable/modules/clustering.html
- Interesting demos on https://www.toptal.com/ machine-learning/clustering-algorithms

EM 000000 k-means

HCA 0000

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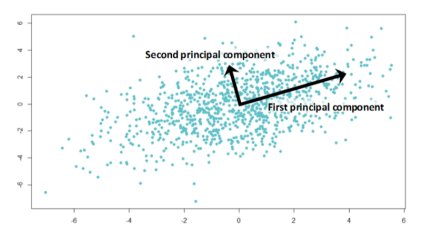
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## Principal Component Analysis

- Observation  $X_1, \ldots, X_n$  in  $\mathbb{R}^p$ , centered
- PCA = Dimension reduction tool (for visual clustering)
- Idea: project onto a subspace  $R^d \subset \mathbb{R}^p$  so as to save as much variance as possible
- Big Picture: orthogonal linear transformation such that the first component has highest variance, then the second, etc.
- ullet First component:  $w_1 = \mathrm{argmax}_{\|w\|=1} \sum (x_i \cdot w)^2$  is the eigenvector of  $X^TX$  corresponding to the highest eigenvalue.
- Similar reasonning for the next components in the orthogonal of  $w_1$ .

### PCA: visualization



Src: [https://techannouncer.com/

 ${\tt global-pca-unit-market-2017-adelte-airmak-industries-amss-ltd-cavotec-airport-division-ciat-effeti/]}$ 

## PCA algorithm

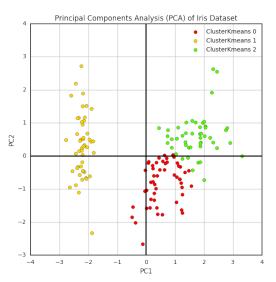
#### **PCA**

- Center all variables
- Compute the  $p \times p$  empirical covariance matrix  $X^T X$ .
- Compute the components  $W_d$  = the d first eigenvectors of  $X^TX$  in decreasing order of the eigenvalues
- Return the projection of X onto the d first components  $T_d = X W_d$ .

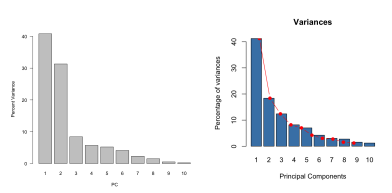
#### Then:

- either vizualize clusters (2d or 3d plots)
- or use another clustering algorithm on the lower-dimensionnal data  $T_d$  (dimension reduction)

# Example: IRIS



## PCA scree plot



 ${\tt Src:\ http://strata.uga.edu/8370/lecturenotes/principalComponents.html\ and}$ 

http://www.sthda.com/english/wiki/print.php?id=207

# Independent Component Analysis (ICA)

- similar idea, but search for *independent* (instead of uncorrelated) components
- computationally more demanding, but iterative (entropy-based) algorithm exists
- often used for blind source separation
- see also Non-negative Matrix Factorization (NMF)
- for vizualization, see also t-SNE: t-distributed Stochastic Neighbor Embedding

### Commands

#### scikitlearn:

class sklearn.decomposition.PCA(n\_components=None,
copy=True, whiten=False, svd\_solver=?auto?, tol=0.0,
iterated\_power=?auto?, random\_state=None)

ICA: sklearn.decomposition.FastICA

**R:** package stats

prcomp() et princomp() [fonction de base, package stats],
PCA() [package FactoMineR],
dudi.pca() [package ade4],
epPCA() [package ExPosition]

ICA: ica

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# Model-based approach: Gaussian Mixtures

- Observations  $X_1, \ldots, X_n$  in  $\mathbb{R}^p$
- K cluster centers:  $\mu_1, \ldots, \mu_K$
- Cluster k has probability  $p_k$
- Points in cluster k have law  $\mathcal{N}(\mu_k, \Sigma)$
- Given a sample  $(X_1, ..., X_n)$ , how to estimate the cluster centers  $(\mu_k)_k$  and how identify the clusters?

# Expectation-Maximization Algorithm

• Likelihood: the parameter  $\theta = (p, \mu)$  has likelihood

$$L(\theta) \propto \sum_{z_1,\dots,z_n \in \{1,\dots,K\}^n} \prod_{i=1}^n p_{z_i} e^{-\frac{1}{2}\Sigma^{-1}(X_i - \mu_{z_i})\Sigma^{-1}}$$

- non-convex, very hard to maximize
- Approximate iterative optimization

## Expectation-Maximization Algorithm

Given an estimate  $\theta^j = (p^j, \mu^j)$ , compute

membership weights

$$w_{i,k}^{j} = P_{\theta^{j}}(z_{i} = k|X_{i}) = \frac{p_{k}^{j} e^{-\frac{1}{2}\Sigma^{-1}(X_{i} - \mu_{k}^{j})\Sigma^{-1}}}{\sum_{\ell=1}^{K} p_{\ell}^{j} e^{-\frac{1}{2}\Sigma^{-1}(X_{i} - \mu_{\ell}^{j})\Sigma^{-1}}}$$

updated cluster weights:

$$p_k^{j+1} = \frac{\sum_{i=1}^{n} w_{i,k}^{j}}{n}$$

updated cluster means:

$$\mu_k^{j+1} = \frac{\sum_{i=1}^n w_{i,k}^j X_i}{\sum_{i=1}^n w_{i,k}^j}$$

## Convergence of the Expectation-Maximization Algorithm

### Theorem

The likelihood of the iterates are increasing:

$$L\left(\theta^{j+1}\right) \geq L\left(\theta^{j}\right)$$

Good: converges
Bad: local optimum

### **EM Algorithm**

- ullet randomly initialize  $heta_0$
- compute EM iterations until convergence:
  - membership weights  $(w_{i,k}^j)_{i,k}$
  - updated cluster weights  $(p_k^{j+1})_k$
  - updated cluster means  $(\mu_k^{j+1})_k$
- start again (a few times) to look for a better local optimum

### Commands

#### scikitlearn:

```
class sklearn.mixture.GaussianMixture(n_components=1,
covariance_type=?full?, tol=0.001, reg_covar=1e-06,
max_iter=100, n_init=1, init_params=?kmeans?,
weights_init=None, means_init=None, precisions_init=None,
random_state=None, warm_start=False, verbose=0,
verbose interval=10)
```

#### R:

```
MClust() [package MASS],
GuassianMixtures() [package sBIC]
```

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# Model-free clustering

- Observations  $X_1, \ldots, X_n$  in  $\mathbb{R}^p$ :
- Objective function: for candidate cluster centers  $\mu = (\mu_1, \dots, \mu_K)$  and cluster assignations  $z = (z_1, \dots, z_n)$ :

$$L(\mu, z) = \sum_{k=1}^{K} \sum_{i: z_i = k} \|X_i - \mu_k\|^2 = \sum_{i=1}^{n} \sum_{k=1}^{k} \mathbb{1}\{z_i = k\} \|X_i - \mu_k\|^2$$

• If  $S_k = \{i : z_i = k\}$ ,

$$L(\mu, z) = \sum_{k=1}^{K} |S_k| \, \mathbb{V}ar[S_k]$$

 Minimizing L is equivalent to minimizing pairwise deviations in the clusters:

$$\operatorname{argmin}_{\mu,z} L(\mu,z) = \operatorname{argmin}_{\mu,z} \sum_{k=1}^K \frac{1}{|S_k|} \sum_{i,j \in S_k} \|X_i - X_j\|^2$$

# Lloyd's algorithm

- For a fixed  $\mu$ , optimizing in z is easy: choose  $z_i = \operatorname{argmin}_k \|X_i \mu_k\|$
- BUT optimizing in  $\mu$  is NP-hard!

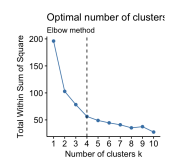
#### k-means

- randomly initialize  $\theta_0$
- compute Lloyd's iterations until convergence:
  - membership variables  $z_i^j = \operatorname{argmin}\{\|X_i \mu_k^j\|$
  - updated cluster weights  $N_k^j = \sum_{i=1}^n \mathbb{1}\{z_i^j = k\}$
  - updated cluster means  $\mu_k^{j+1} = \frac{\sum_{i:z_i^j=k}^j X_i}{N_k^j}$
- start again (a few times) to look for a better local optimum

### Comments on k-means

- k-means is a "hard" version of EM for mixtures! (when variances tends to 0)
- Complexity: linear in n and k (fast)
- Requires only a dissimilarity measure (not necessarily a distance)
- Quality of solution found depends on (random) initialization
- Not robust to outliers

 Problem: how to choose k?
 [Src: http://www.sthda.com/english/articles/ 29-cluster-validation-essentials/]



## Improved Initialization: k-means++

[Arthur, D.; Vassilvitskii, S. (2007). "k-means++: the advantages of careful seeding" (PDF). Proceedings of the eighteenth annual ACM-SIAM symposium on Discrete algorithms.]

Idea: enforce distant cluster centers from the start

#### k-means++

- Choose first center  $\mu_1^1$  at random
- for j = 2 to K, repeat:
  - compute  $D_i^j = \min_{\ell < j} \|X_i \mu_\ell^1\|$
  - choose  $\mu_j^1 = X_i$  with probability proportionnal to  $D_i^j$
- theoretical guarantee of  $O(\log k)$  approximation from the start
- still linear complexity
- often a dramatic improvement in practice

## k-medoids: robustness vs computation time

#### k-medoids

- randomly initialize  $\theta_0$  with K points of the dataset
- iterate until convergence:
  - membership variables  $z_i^j = \operatorname{argmin}\{\|X_i \mu_k^j\|$
  - updated cluster medoids  $\mu_k^{j+1} = \operatorname{argmin}_{i \in S_k} \sum_{\ell \in S_k} \|X_\ell X_i\|$
- start again (a few times) to look for a better local optimum
- Robust to outliers (cf median versus mean)
- BUT computation time is quadratic in n
- k-means is k-medoids with  $||X_{\ell} X_i||^2$  instead of  $||X_{\ell} X_i||$

## Commands

#### scikitlearn:

```
class sklearn.cluster.KMeans(
n_clusters=8, init=?k-means++?, n_init=10, max_iter=300,
tol=0.0001, precompute_distances='auto', verbose=0,
random_state=None, copy_x=True, n_jobs=1, algorithm='auto'
```

### **R:** package stats

```
kmeans(x, centers, iter.max = 10, nstart = 1,
algorithm = c("Hartigan-Wong", "Lloyd", "Forgy",
"MacQueen"), trace=FALSE)
```

Variants to be found in various packages: ClusterR, kmed, etc.

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# Agglomerative clustering

- greedy bottom-up algorithm
- requires a distance (idssimilarity) between observations ||x - x'||
- choice of distance between clusters:
  - complete linkage:  $d(A, B) = \max\{||x x'|| : x \in A, x' \in B\}$

  - single linkage:  $d(A, B) = \min \{ ||x x'|| : x \in A, x' \in B \}$  average linkage distance:  $d(A, B) = \frac{1}{|A||B|} \sum_{x \in A} \sum_{x' \in B} ||x x'||$
  - Ward distance for Euclidian mean:

$$d(A, B) = \frac{|A||B|}{n(|A| + |B|)} ||\bar{A} - \bar{B}||$$

- sum of intra-cluster variance
- etc.

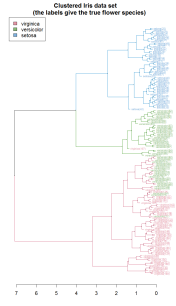
## HCA algorithm

#### HCA

- Initialization: all observations are clusters  $\{X_1\}, \ldots, \{X_n\}$
- As long as there are at least two clusters:
  - add a link between two clusters with smallest distance
  - merge them for the next iterations
- Return the *dendrogram* = hierarchy of clusters

Property of Ward for Euclidian distance: interclass variance decreasing with the number of classes

# Dendrogram



Author: Talgalili

https://commons.wikimedia.org/wiki/File:

Iris\_dendrogram.png

### Pros and Cons

- No need to specify the number of clusters in advance
- A relevant choice can be deduced from the observation of the dendrogram (and practical needs)
- Computational complexity in  $O(n^2)$
- Does not find an optimal solution

### Commands

#### scikitlearn:

```
class sklearn.cluster.AgglomerativeClustering(n_clusters=2
affinity=?euclidean?, memory=None, connectivity=None,
compute_full_tree=?auto?, linkage=?ward?,
pooling_func=<function mean>)
```

#### R:

```
hclust(d, method = "complete", members = NULL)
```

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# Affinity Propagation

- Message-passing algorithm
- affinity function s(x, x'), for example  $s(x, x') = -\|x x'\|^2$
- s(x,x) = input preference: the lower, the higher the chances to be an exemplar
- responsibility matrix R:  $r(i, k) = \text{how well-suited } X_k \text{ can}$  serve as an exemplar for  $X_i$  (wrt other candidate exemplars)
- availability matrix A:  $a(i, k) = \text{how appropriate it is for } X_i \text{ to pick } X_k \text{ as an examplar, taking into account other points preferences for } X_k \text{ as an exemplar}$

# Affinity Propagation

## **Affinity Propagation**

- Initialize R and A with 0
- Repeat until convergence:
  - update responsibilities:  $r(i, k) \leftarrow s(i, k) - \max_{k' \neq k} a(i, k') + s(i, k')$
  - update availabilities:

$$a(i,k) \leftarrow \min \left\{ 0, r(k,k) + \sum_{i' \notin \{i,k\}} \max(0, r(i',k)) \right\} \text{ for } i \neq k$$
and 
$$a(k,k) \leftarrow \sum_{i' \neq k} \max(0, r(i',k))$$

• Pick exemplars as maximizers of r(i, i) + a(i, i)

# Affinity Propagation: pros/cons

- No need to specify the number of clusters
- ... but a parameter plays the same role
- quadratic time complexity
- some improvement over k-means in some cases

## Commands

#### scikitlearn:

```
class sklearn.cluster.AffinityPropagation(damping=0.5,
max_iter=200, convergence_iter=15, copy=True,
preference=None, affinity=?euclidean?, verbose=False)
```

### R: package APCluster

```
apcluster(s, x, p=NA, q=NA, maxits=1000, convits=100, lam=0.9, includeSim=FALSE, details=FALSE, nonoise=FALSE, seed=NA)
```

# Spectral Clustering

- Similarity matrix  $S_{i,j}$ , for example  $S_{i,j} = -\|X_i X_i\|^2$
- Idea: use standard clustering method on eigenvectors of the (normalized) Laplacian matrix

$$L = Id - D^{-1/2} S D^{-1/2}$$

where D is diagonal with  $D_{i,i} = \sum_{j} S_{i,j}$ 

 Intuition: if S is diagonal by blocks, the eigenvectors are the indicators of the blocks

## Commands

#### scikitlearn:

```
class sklearn.cluster.SpectralClustering(n_clusters=8, eigen_solver=None, random_state=None, n_init=10, gamma=1.0, affinity=?rbf?, n_neighbors=10, eigen_tol=0.0, assign_labels=?kmeans?, degree=3, coef0=1, kernel_params=None, n_jobs=1)
```

```
R: package kernlab
```

```
specc(x, data = NULL, na.action = na.omit, ...)
```

### **DBscan**

- DBscan = density-based spatial clustering of applications with noise
- ullet parameters: radius  $\epsilon$  and minimal cluster size minSize

#### **DBscan**

Repeat as long as at least one point has not been visited:

- pick an unvisited point X<sub>i</sub> at random
- ullet if it has less than minSize  $\epsilon$ -neighbors, mark it as outlier
- other, form the cluster of all points that can be reached by jumps of at most  $\epsilon$  starting from  $X_i$

### DBscan: comments

- simple and fast
- no need to specify number of clusters
- Problem: sensitive to the choice of parameters
- ullet choosing the right parameters  $\epsilon$  and  $\emph{minSize}$  propertly is hard
- choice of  $\epsilon$ : such that the proportion of outliers is at most 10% (say)
- choice of minSize: such that at least 90% have at least minSize neighbors
- unable to handle clusters with very different densities

### Commands

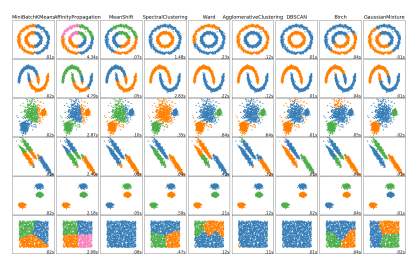
#### scikitlearn:

```
class sklearn.cluster.DBSCAN(eps=0.5, min_samples=5,
metric='euclidean', metric_params=None, algorithm='auto',
leaf_size=30, p=None, n_jobs=1)
```

#### R: package dbscan

```
dbscan(x, eps, minPts = 5, weights = NULL,
borderPoints = TRUE, ...)
```

## Which algorithm to choose?



 $Src: \ [http://scikit-learn.org/stable/auto\_examples/cluster/plot\_cluster\_comparison.html] \\$ 

## Clustering time series

#### Features:

- mean
- trend
- auto-correlation coefficients
- inter-series correlation
- etc.

 $\implies$  it depends on the nature of the problem and on the goal of the clustering!