Weekly meeting 4

Dr. Doina Bein Friday, June 16, 10:30am-12pm

Surveys to be completed

To be done today, before starting research:

CIC-PCUBED Pre-event survey:

https://fullerton.gualtrics.com/jfe/form/SV 6YIVSkC6hLxbunA

Project 1: Data Science

What you need to do: topics & objectives

Objective 1: Learn Python using some textbook or some online courses such as

(https://www.codecademy.com/learn/learn-python). Shared by Stephanie Pocci: Learn Python in a couple hours. This YouTuber does a very beginner-friendly crash course about the capabilities of Python and its uses. Here is the link:

https://www.youtube.com/watch?v=rfscVS0vtbw

Objective 2: Learn how to use Jupyter Notebook. Start here http://jupyter-notebook-beginner-guide.readthedocs.io/en/latest/what_is_jupyter.html

Objective 3: For data science, find a suitable dataset and start training some neural network using with Google tensorflow.

Logistics for all students

- Who is participating: <u>list of current research students</u> and their availability
- Research will be conducted virtually during the week with in-person meetings throughout the week
- Zoom meetings for me to teach new topics and for you to participate in open discussions
- Support:
 - If needed, you can meet me
 Zoom: Mon, Tu, Wed from 8:30-10:25 am
 IN PERSON: Mon, Tu, Wed from 8:30-9:30 am, Thursday 8:30-10am or
 by email
 - CIC-PCUBED peer mentor: (tentative) <u>availability</u>

Logistics for all students (contd.)

- Make a copy of this GDoc Work schedule, share the Gdoc copy with me, and maintain it weekly and daily; due at the end of Week 2
- Before the end of week 3, make a copy and maintain your
 Proposed work by individual or teams of up to three; due by the end of Week 3
- Complete your <u>availability here</u>; try to have it consistent over the 7 weeks such that it will be easy to partner in the project
- Group projects: to be decided; sample list <u>here</u>
- Oral or poster presentations: tentatively scheduled for Friday,
 July 28, from 8:30am-12:30 pm and if needed, from 1:30-4 pm

Please checkout:

- Other websites and ebooks
- Websites with free datasets
- If you find good, free resources, please share it by email or during weekly meetings
- Next meeting: I will lecture on ZOOM on Data Science: Thursday, June 22, from 10:30am-12pm

Progress on Learning Python

- Free course: https://www.codecademy.com/learn/learn-python
- Free course: https://www.kaggle.com/learn/python
- Youtube video (about 4 hours):
 https://www.youtube.com/watch?v=rfscVS0vtbw

Data Science

k-means Clustering

 The k-means cost function asks to minimize the sum of squared Euclidean distances of data points to their closest prototype centers:

$$e_k(X;C) = \sum_{i=1}^n \min_{j \in \{1, \dots k\}} ||x_i - c_j||^2$$

• k-means cost function seeks compact globular clusters of small variances: the cost of a single cluster $e_1(G) = e_1(G, c)$ is minimized when we choose for the cluster prototype its center of mass c, called the centroid:

$$c(G) = \underset{c}{\operatorname{argmin}} \sum_{x \in G} ||x - c||^2 = \frac{1}{G} \sum_{x \in G} x$$

where |G| denotes the cardinality of G, that is the number of elements contained in group G and $\operatorname{argmin}_x f(x)$ to denote the argument that yields the minimum in case this minimum value is unique

- If instead of choosing the squared Euclidean distance, we had chosen the ordinary Euclidean distance, one obtains the socalled Fermat-Weber point that generalizes the notion of median. It is thus also called the geometric median.
- Although the Fermat-Weber point is unique and often used in operations research for facility location problems, it does not admit a closed-form solution, but can be arbitrarily finely approximated
- k-median clustering is the clustering obtained by minimizing the cost function

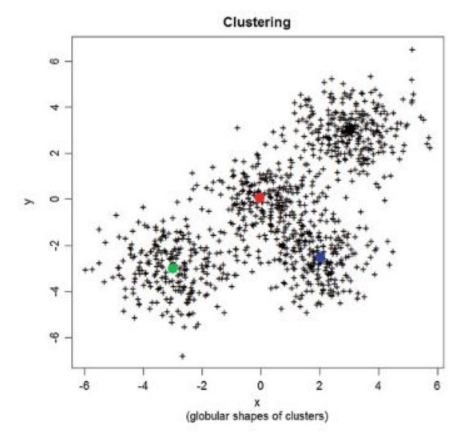
$$\min_{C} \sum_{i=1}^{n} \min_{j \in \{1, \dots k\}} ||x_i - c_j||$$

where ||... ||means the regular Euclidean distance

 Partitions from k-means and k-medians can be very different from each other: the centroid location can be different to the median for a single cluster

- Centroids can be easily corrupted by adding a single outlier point
- We say that the breakdown point of the centroid is 0: A single outlier p₀ diverging to ∞ will impact the centroid to be diverging to ∞ too.
- But the median is more robust since it requires $\left\lfloor \frac{n}{2} \right\rfloor$ outliers (that is, about 50% of outliers) to steer the median point to ∞ .
- Therefore k-median clustering is often preferred when there are many outliers in data-sets.

Fig. 7.3 The k-means cost function tend to find globular-shaped clusters that minimize the weighted sum of the cluster variances. k-Means clustering is a model-based clustering where each cluster is associated to a prototype: its center of mass, or centroid. Here, we have choosen k = 4 groups for the k-means: Cluster prototypes, centroids, are illustrated with large disks



The k-means Problem (https://www.math.uwaterloo.ca/~cswamy/talks/kmeans-short.p pt)

Given: n points in d-dimensional space

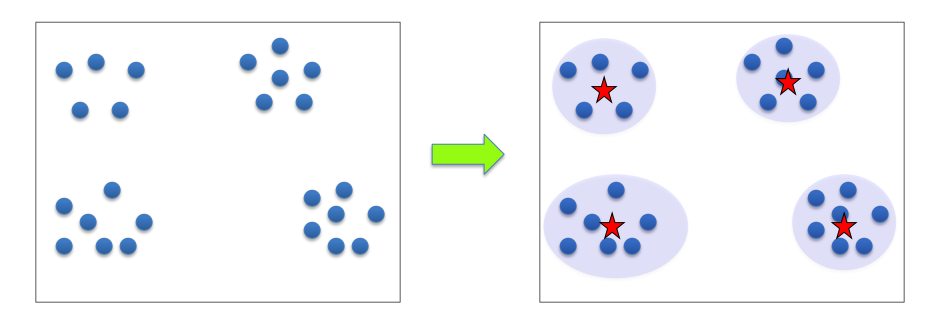
$$X \subseteq R^d$$
: point set with $|X| = n$

- partition X into k clusters X₁,..., X_k
- assign each point in X_i to a common center

Goal: Minimize $\sum_{i} \sum_{x \in X_i} d(x, c_i)^2$

(web.stanford.edu/group/mmds/slides2012/s-speaker-ppt-2012-temp/Bahmani.pptx)

- K-means clustering is a fundamental problem in data analysis and machine learning
- "By far the most popular clustering algorithm used in scientific and industrial applications" [Berkhin '02]
- Identified as one of the top 10 algorithms in data mining [Wu et al '07]



k-Means Optimization Problem

- Finding the center of a single cluster is a particular case of clustering with k
 = 1 cluster.
- With the squared Euclidean distance cost, the center is the mean of the attributes, hence its naming *k-means*.
- Finding the minimum of a k-means cost function is a NP-hard problem when the dimension d > 1 and the number of clusters k > 1.
- When k = 1, we have shown that we can compute the optimal solution (the centroid) in linear time (computing the mean of the group).
- When d = 1, we can compute an optimal k-means solution using dynamic programming: Using O(nk) memory, we can solve the k-means for n scalar values in time $O(n^2k)$

- For NP-hard problems, we seek efficient heuristics to approximate the cost function. We distinguish two classes of such heuristics:
- 1. Global heuristics that do not depend on initialization, and
- 2. Local heuristics that iteratively starts from a solution (a partition) and iteratively improves this partition using "pivot rules."

Lloyd's Batched k-Means Local Heuristic

▲ Lloyd's heuristic (1957): from a given input and initialization:

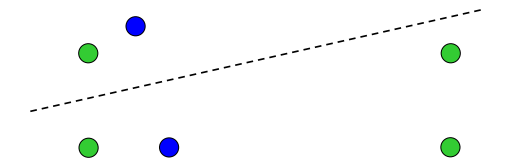
Input: $\{x_1, x_2, ... x_n\}$, each $x_1 \in \mathbb{R}^d$, the number of partitions k

Initialization: Start with k cluster centers $\{c_1, c_2, ..., c_k\}$ (typically chosen uniformly at random from data points)

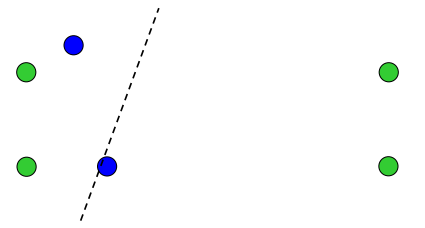
iteratively repeat until convergence the following two steps:

- 1. **Assign points to clusters**: (each $x_i \in X$ is assigned to its closest cluster center) for each $x_i \in X$, let $l_i = \operatorname{argmin}_l \|x_i c_i\|^2$, and define the k cluster groups as $G_j = \{x_i : l_i = j \}$ with $n_j = |G_j|$, the number of elements of X falling into the jth cluster.
- 2. **Update centers** (perform an Expected Maximization-type local search): For all $j \in \{1, ..., k\}$, update the centers to their cluster centroids : $c_j = \frac{1}{n} \sum_{x \in G_j} x$ (or the barycenters $c_j = \frac{1}{\sum_{x \in G_j} w(x)} \sum_{x \in G_j} w(x) x$ for weighted data-sets).
- A possible convergence criteria: cluster centers do not change anymore

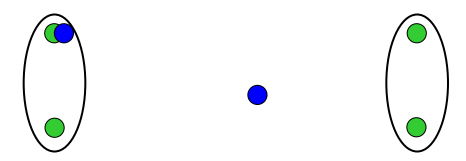
- b) Heuristics: Lloyd's method invented in 1957 and remains an extremely popular heuristic even today
 - 1) Start with k initial / "seed" centers c_1, \ldots, c_k .
 - 2) Iterate the following Lloyd step
 - a) Assign each point to nearest center c_i to obtain clustering $X_1, ..., X_k$.
 - b) Update $c_i \leftarrow ctr(X_i) = \sum_{x \in X_i} x/|X_i|$.



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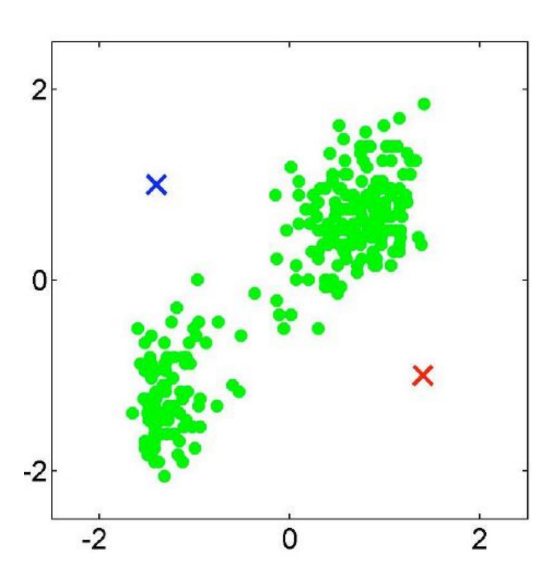
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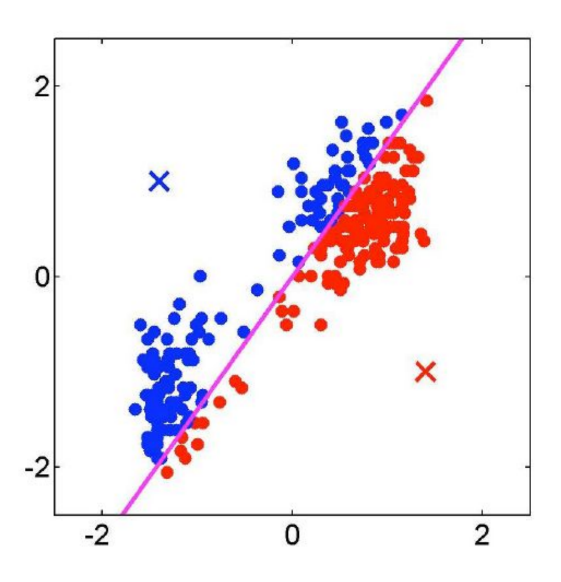
Other example

(https://www.cs.utah.edu/~piyush/teaching/4-10-print.pdf)

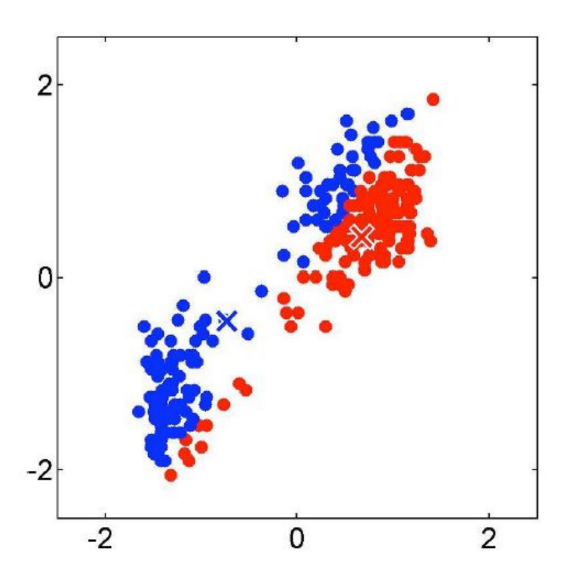
Initialization (assume K = 2)



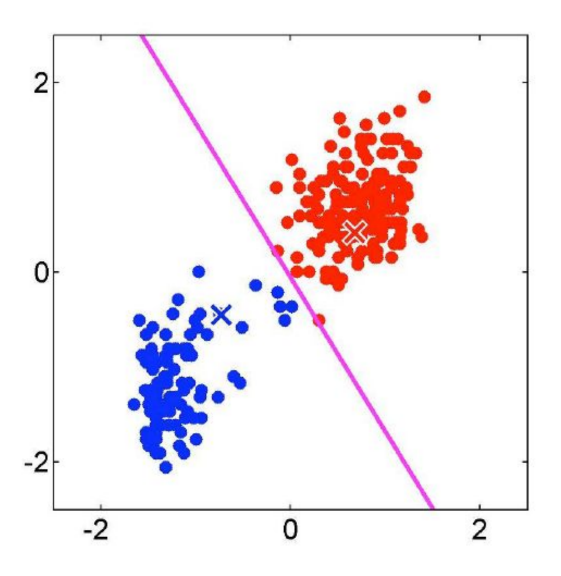
Iteration 1: Assign points to clusters



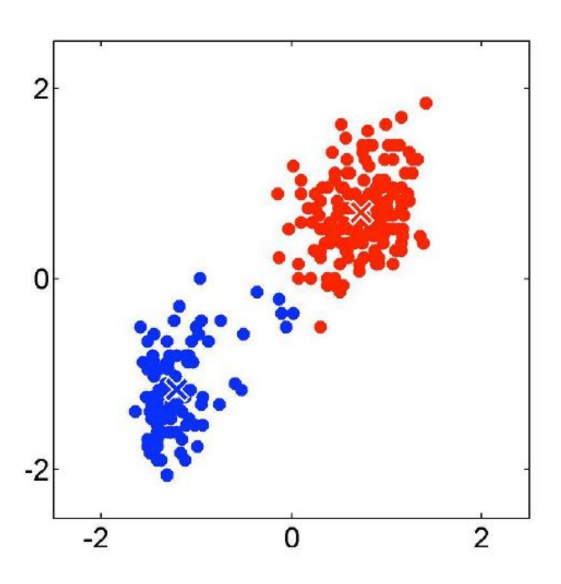
Iteration 1: Update centers



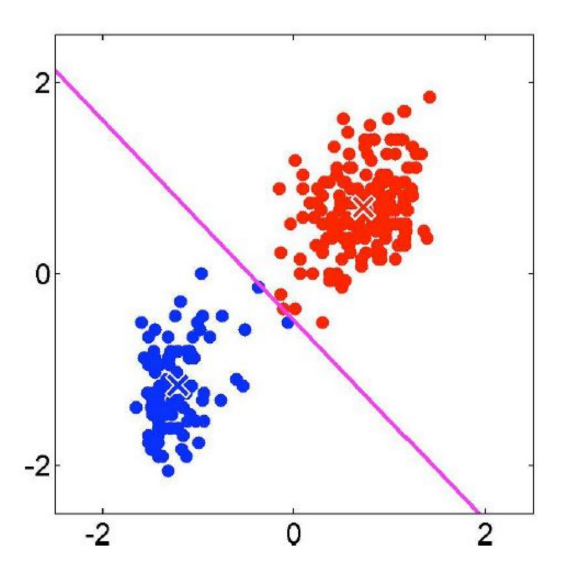
Iteration 2: Assign points to clusters



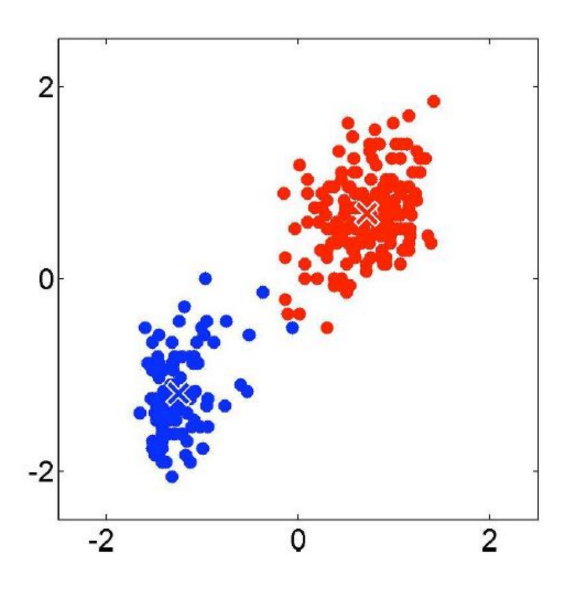
Iteration 2: Update centers



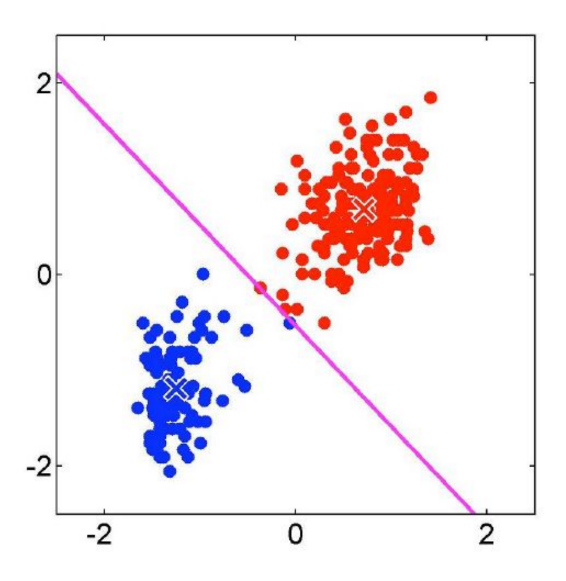
Iteration 3: Assign points to clusters



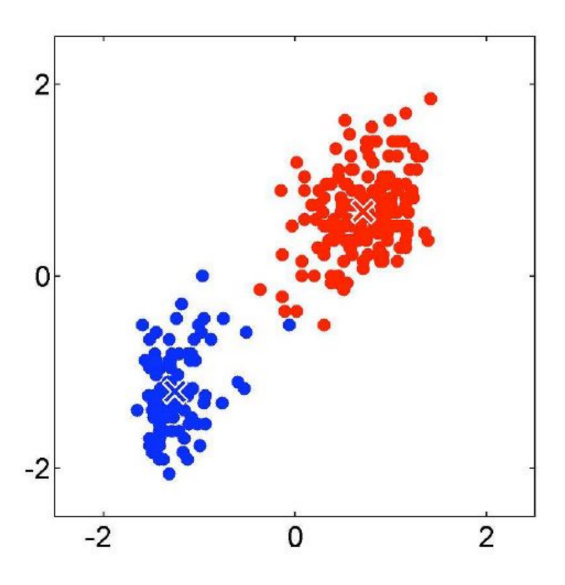
Iteration 3: Update centers



Iteration 4: Assign points to clusters



Iteration 4: Update centers



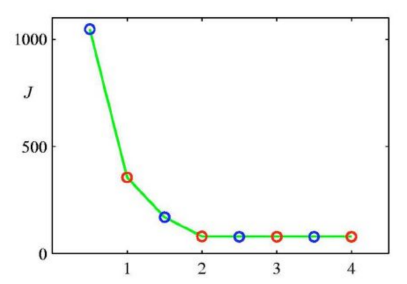
Choosing the number of clusters K

(https://www.cs.utah.edu/~piyush/teaching/4-10-print.pdf)

• Recall that k-means objective is to minimize the cost function, aka the sum of distances of points from their cluster centers

$$J(C,r) = \sum_{i=1}^{n} \sum_{j=1}^{n} r_{nk} ||x_i - c_j||^2$$

- One way to select the number of clusters k is to try different values of k, plot the k-means objective versus k, and choose the "elbow-point"
- One drawback of this method is that it is computationally very expensive, and sometimes (depending on the data-sets), the inflexion point between the sharp decrease and the plateau is not so well defined



For the plot to the left, k = 2 is the elbow point

Initialization Issues:

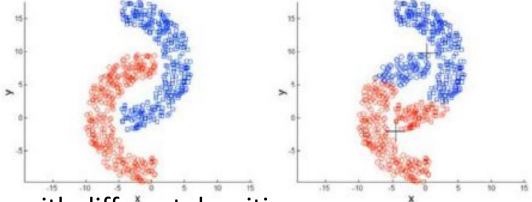
(https://www.cs.utah.edu/~piyush/teaching/4-10-print.pdf)

- K-means is extremely sensitive to cluster center initialization
- Bad initialization can lead to
 - Poor convergence speed
 - Bad overall clustering
- Safeguarding measures:
 - Choose first center as one of the examples, second which is the farthest from the first, third which is the farthest from both, and so on.
 - Try multiple initializations and choose the best result
- Other smarter initialization schemes (e.g., look at the K-means++ algorithm by Arthur and Vassilvitskii)

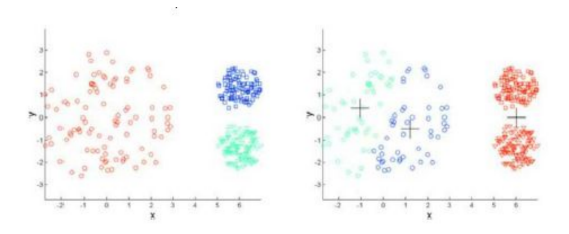
Limitations Illustrated

(https://www.cs.utah.edu/~piyush/teaching/4-10-print.pdf)

- K-Means might not find the best possible assignments and centers.
- Non-convex/non-round-shaped clusters: Standard K-means fails!



Clusters with different densities



k-means++

(cs.haifa.ac.il/hagit/courses/seminars/.../Presentations/Lecture01_K-means+EM.pptx)

- k-Means with smart initial seeding
- Choose one center uniformly at random from among the data points.
- For each data point x, compute D(x), the distance between x and the nearest center that has already been chosen.
- Choose one new data point at random as a new center, using a weighted probability distribution where a point x is chosen with probability proportional to D(x)^2.
- Repeat Steps 2 and 3 until k centers have been chosen.
- Now that the initial centers have been chosen, proceed using standard k-means.

Advantages

- This seeding method yields considerable improvement in the final error of k-means
- Takes more time to initialize
- Once initialized, K-Means converges quickly
- Usually faster than K-Means
- 1000 times less prone to error than K-Means

Computing the Similarity of Two Partitions

- We need ground-truth datasets that tell us for each data element is true cluster membership
- Without these ground-truth data sets, we could only perform a subjective or qualitative evaluation of the clustering methods.
- Human eyes can sometimes evaluate whether the obtained clustering is good or not for d < 4
- When ground-truth data-sets are available (say, data-sets annotated by experts), we can compute various metrics that are quantitative values that measure the similarity of two partitions: the one induced by the labels in the ground-truth data-set (assumed to be the optimal clustering by definition!) and the one reported by an automatic clustering algorithm
- The Rand index (1971) computes the similarity of two partitions

Rand Index (1971)

- computes the similarity of two partitions
- Let $G = \bigcup G_i$ and $G' = \bigcup G_i'$ cluster decomposition of the k-means heuristic and of the ground-truth data-set, respectively.
- Compare all the n*(n-1)/2 (n choose 2) pairs (x_i, x_j) of points, and count those that belong to the same cluster (a) from those that are found to belong to different clusters (b).
- The Rand index

$$Rand(G, G') = \frac{a+b}{\binom{n}{2}}$$

with

a:
$$\#\{(i, j) : I(x_i) = I(x_j) \land I'(x_i) = I'(x_j)\}$$

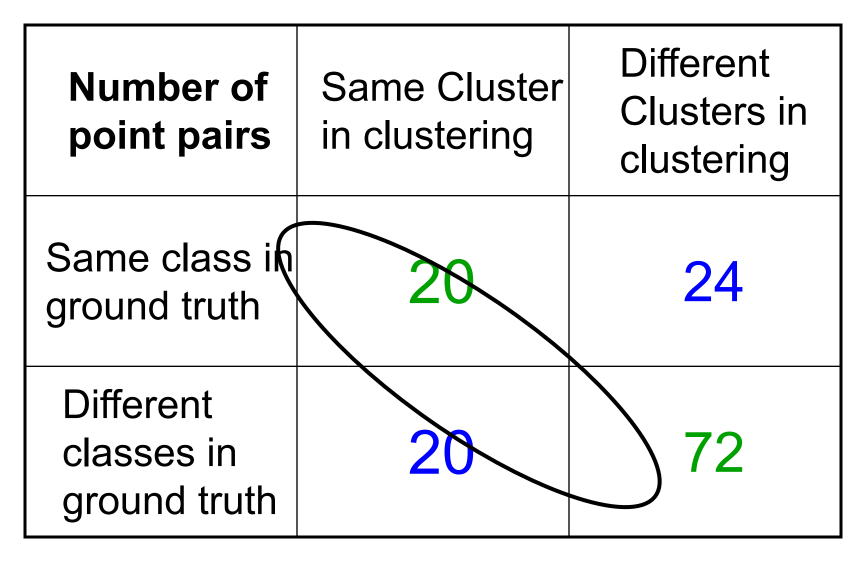
b: $\#\{(i, j) : I(x_i) \neq I(x_i) \land I'(x_i) \neq I'(x_j)\}$,

I and I' are the two cluster labeling functions of the ground-truth clustering and the automatic clustering

Rand index belongs to the interval [0, 1]

Rand Index measures between pair decisions. Here RI = 0.68

(https://web.stanford.edu/class/cs276/handouts/lecture14-clustering.ppt)



Supervised Learning

- In supervised learning, we are given a labeled *training set Z* = $\{(x_i, y_i)\}$ with $y_i \in \pm 1$ (the ground truth labeled data) and the task is to learn a *classifier* so that we can classify new unlabeled observations of a *testing set Q* = $\{x_i'\}_i$.
- When the training set has only two classes, we deal with binary classification, otherwise it is a multi-class classification problem.
- Statistical learning assumes that both the training set and the testing set are independently and identically sampled for an arbitrary but fixed unknown distribution.
- Our focus: learn a target function that can be used to predict the values of a discrete class attribute
- The task is commonly called: Supervised learning, classification, or inductive learning.