Statistical Methods in AI (CSE/ECE 471)

Lecture-9: Unsupervised Learning (k-means, GMM)



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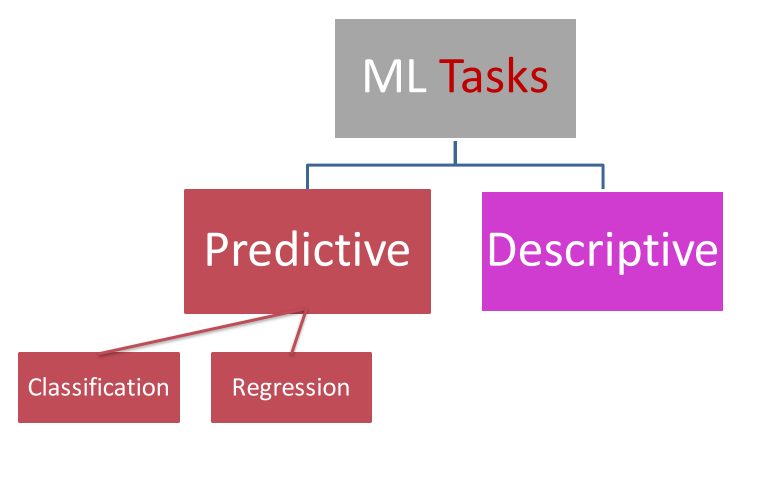
Vineet Gandhi (v.gandhi@iiit.ac.in)

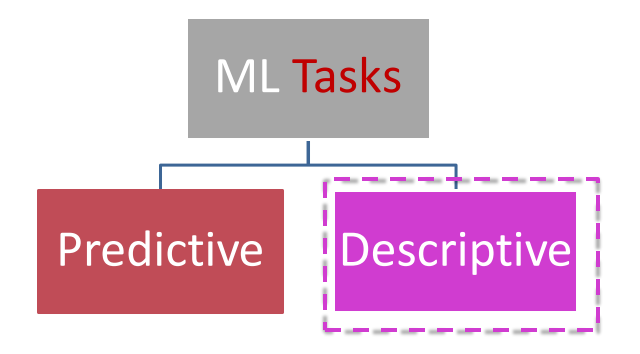




Center for Visual Information Technology (CVIT)

IIIT Hyderabad





ML::Tasks → Descriptive

- Study/Exploit the 'structure' of data
 - Clustering
 - Dimensionality Reduction
 - Density Estimation
- Also studied as 'Unsupervised Learning'
 - Input' data without paired 'Output'

Unsupervised Learning

Task: Given $X \in \mathcal{X}$, learn f(X).

Unsupervised Learning → Clustering

Group similar things e.g. images [Goldberger et al.]



- Determine groups of people in image above
 - based on clothing styles
 - gender, age, etc



features



• Determine moving objects in videos

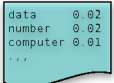
Topic Modelling

Topics

0.04 gene 0.02 genetic 0.01

```
0.02
life
         0.01
evolve
organism 0.01
```

```
0.04
brain
          0.02
neuron
          0.01
nerve
```



Documents

Topic proportions and assignments

Seeking Life's Bare (Genetic) Necessities

COLD SPRING HARBOR, NEW YORK-How many genes does an organism need to survive? Last week at the genome meeting here, * two genome researchers with radically different approaches presented complainerstary views of the basic genes needed for life. - sus answer may be more than just a One research team, using computer analyses to compute known genomes, concluded that today's organisms can be sustained with just 250 genes, and that the earliest life forms required a more 128 genes. The

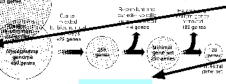
other researcher mapped genes in a stimple parasite and estimated that for this organism. 800 genesors: plenty to do the job—but that anything short of 100 wouldn't be enough.

Although the numbers don't match precisely, those predictions.

* Genome Mapping and Sequencing, Cold Spring Harbor, New York. May 8 to 12.

"are not all that far apart," especially incomparison to the 75,000 genes ut the hunasagenome, notes Six Andersson/a University in Same at the arrived at 800 result of But coming up with a s numbers same, comicularly sequenced. "It may be a way of organizaany newly sequenced genome," explains Aready Mushegian, a computational me-

localar biologist at the National Center. for Biotechnology Information C in Bethesda, Maryland, Comparing a

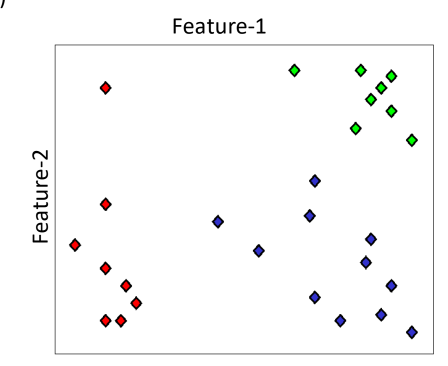


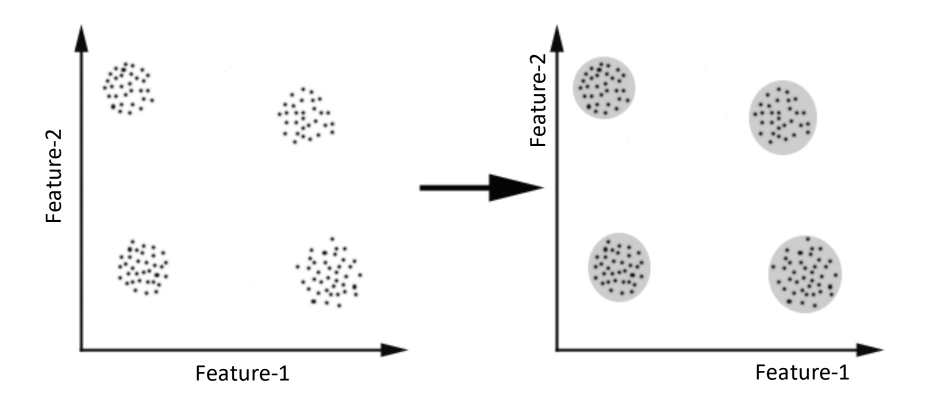
Stripping down. Computer analysis yields an estimate of the minimum modern and ancient genomes

SCIENCE • VOL 272 • 24 MAY 1996

What is Clustering?

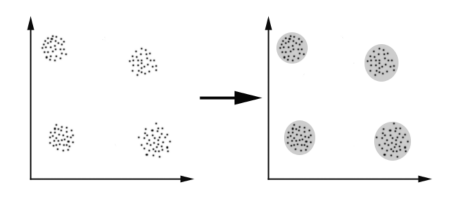
- Organizing data into groups s.t.
 - High intra-group similarity (within members of a cluster)
 - Low inter-group similarity (across clusters)





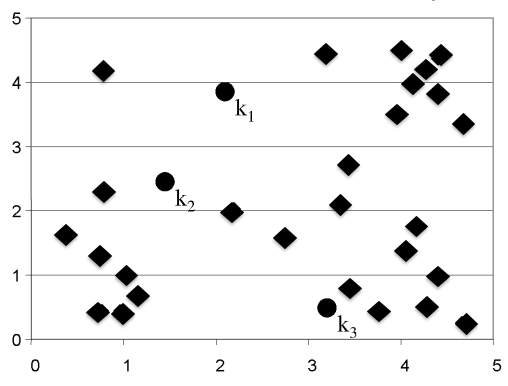
Approach-1: Assume that we (somehow) know the number of clusters

- # of clusters = K (often a 'guess')
- How to represent a cluster?
 - Option-1: Cluster members
 - Option-2: A suitable statistic of the cluster members
- Suitable statistic = Mean

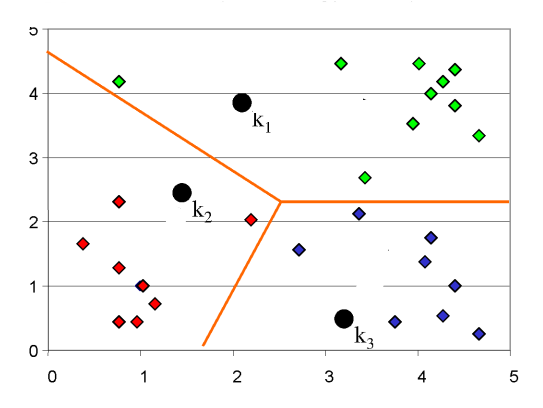


K-means Clustering: Initialization

Decide *K*, and initialize *K* centers (randomly)

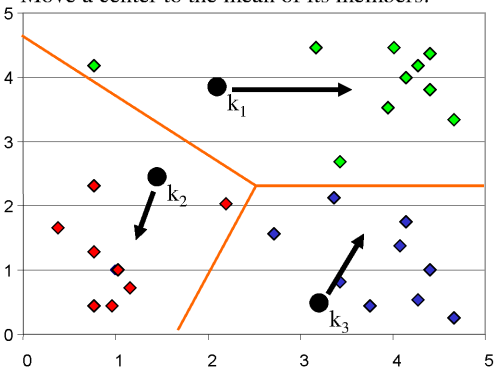


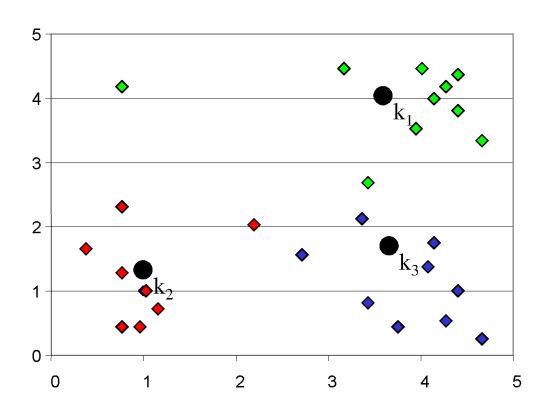
Assign all objects to the nearest center.



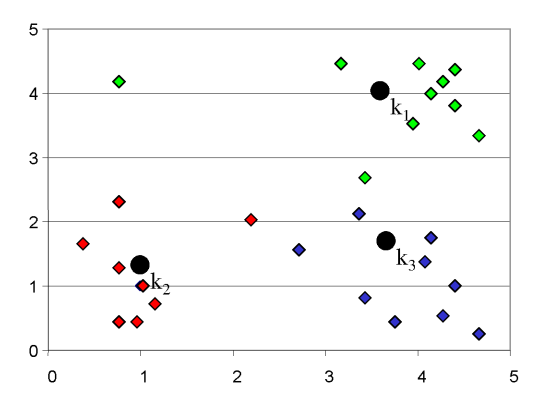
Assign all objects to the nearest center.

Move a center to the mean of its members.

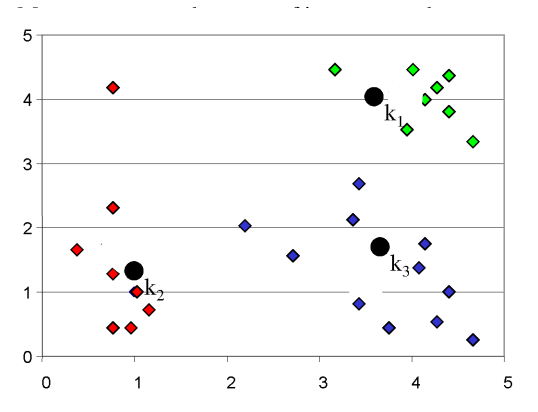




After moving centers, re-assign the objects...

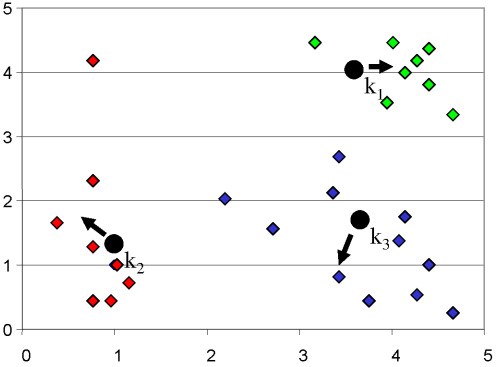


After moving centers, re-assign the objects to nearest centers.



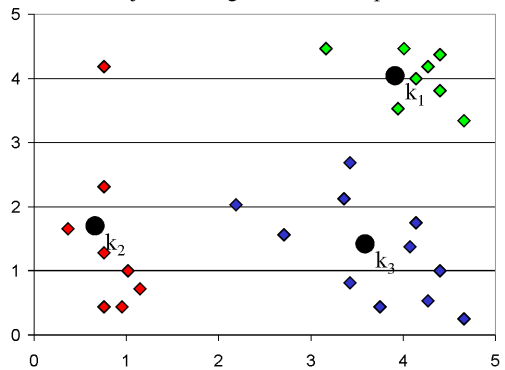
After moving centers, re-assign the objects to nearest centers.

Move a center to the mean of its new members.



K-means Clustering: Finished! Re-assign and move centers, until ...

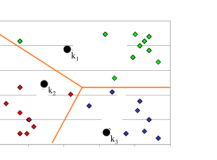
no objects changed membership.



$$\{x^{(1)}, \dots, x^{(m)}\}$$
 $x^{(i)} \in \mathbb{R}^n$

The k-means clustering algorithm is as follows:

- 1. Initialize cluster centroids $\mu_1, \mu_2, \dots, \mu_k \in \mathbb{R}^n$ randomly.



2. Repeat until convergence: {

For every i, set $c^{(i)} := \arg\min_{i} ||x^{(i)} - \mu_{i}||^{2}.$ For each j, set

$$\mu_{j} := \frac{\sum_{i=1}^{m} 1\{c^{(i)} = j\}x^{(i)}}{\sum_{i=1}^{m} 1\{c^{(i)} = j\}}.$$

Assignment step: Assign each data point to the closest cluster

Refitting step: Move each cluster center to the center of the data assigned to it

$$\{x^{(1)}, \dots, x^{(m)}\} \qquad x^{(i)} \in \mathbb{R}^n$$

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- 1. Initialize cluster centroids $\mu_1, \mu_2, \dots, \mu_k \in \mathbb{R}^n$ randomly.
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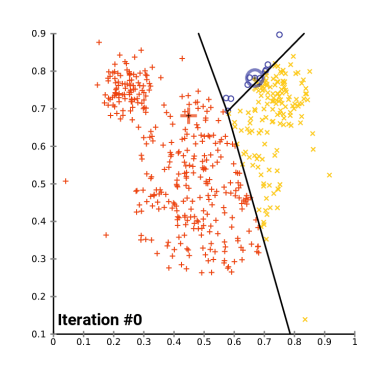
For every
$$i$$
, set

$$c^{(i)} := \arg\min_{i} ||x^{(i)} - \mu_j||^2.$$

For each j, set

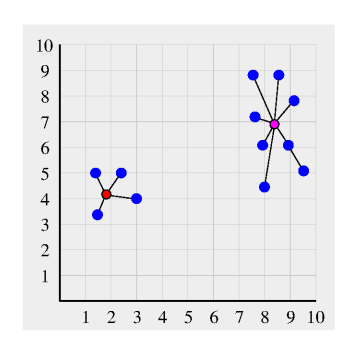
$$\mu_j := \frac{\sum_{i=1}^m 1\{c^{(i)} = j\}x^{(i)}}{\sum_{i=1}^m 1\{c^{(i)} = j\}}.$$





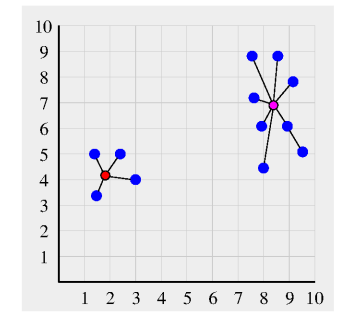
Why K-means Works

- What is a good partition?
- High intra-cluster similarity



Why K-means Works

- What is a good partition?
- High intra-cluster similarity
- K-means optimizes



$$J = \sum_{k=1}^{K} \sum_{i=1}^{n_k} ||x_{ki} - \mu_k||^2$$

Why does K-means work?

Repeat until convergence: {

For every i, set

$$c^{(i)} := \arg\min_{i} ||x^{(i)} - \mu_j||^2.$$

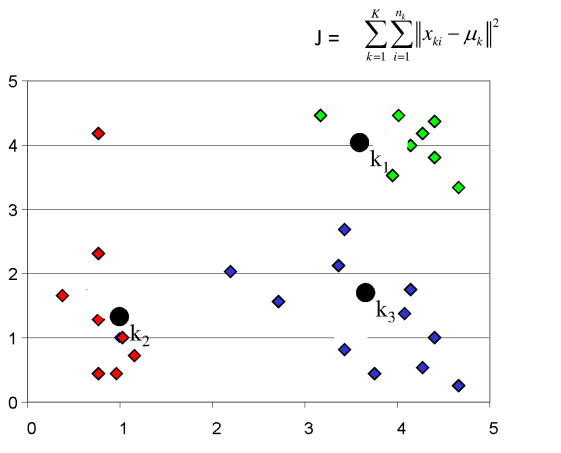
For each j, set

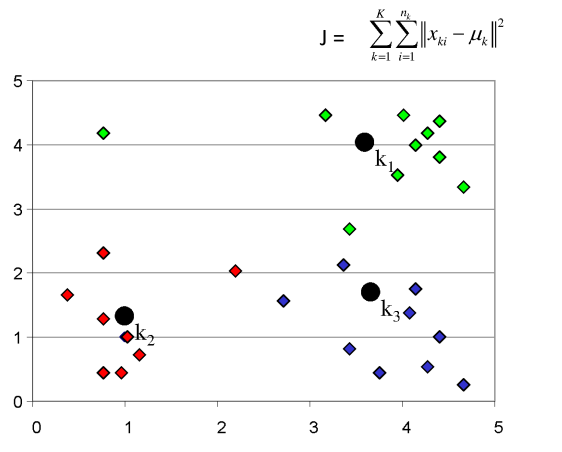
$$\mu_j := \frac{\sum_{i=1}^m 1\{c^{(i)} = j\}x^{(i)}}{\sum_{i=1}^m 1\{c^{(i)} = j\}}.$$

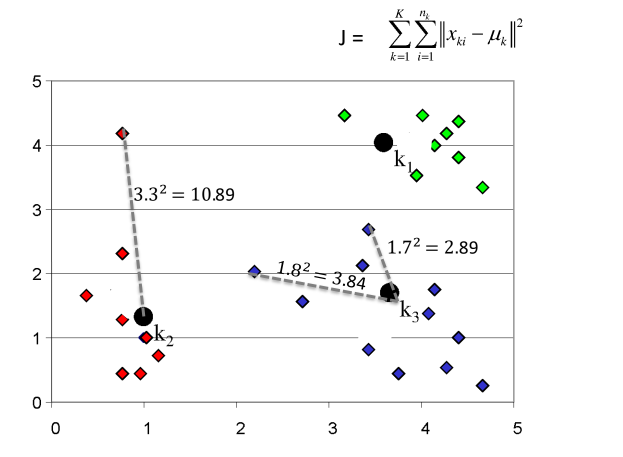
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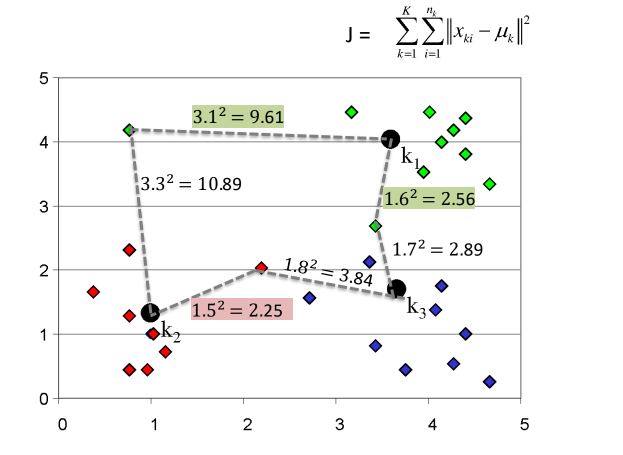
$$J = \sum_{k=1}^{K} \sum_{i=1}^{n_k} ||x_{ki} - \mu_k||^2$$

• Whenever an assignment is changed, the sum squared distances *J* of data points from their assigned cluster centers is reduced.









Why does K-means work?

Repeat until convergence: {

For every
$$i$$
, set
$$c^{(i)} := \arg\min_i ||x^{(i)} - \mu_j||^2.$$

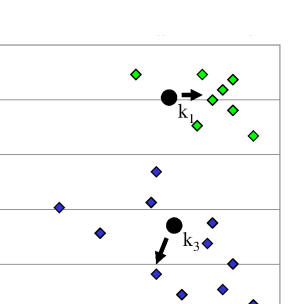
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For each j, set

each
$$j$$
, set
$$\mu_j := \frac{\sum_{i=1}^m 1\{c^{(i)} = j\}x^{(i)}}{\sum_{i=1}^m 1\{c^{(i)} = j\}}.$$

 $J = \sum_{k=1}^{K} \sum_{i=1}^{n_k} ||x_{ki} - \mu_k||^2$

• Whenever an assignment is changed, the sum squared distances J of data points from their assigned cluster centers is reduced. Whenever a cluster center is moved, J is reduced.



For every i, set

$$c^{(i)} := \arg\min_{i} ||x^{(i)} - \mu_{i}||^{2}.$$

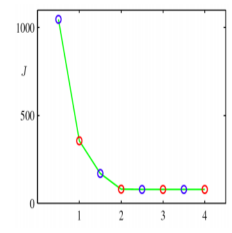
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$$J = \sum_{k=1}^{K} \sum_{k=1}^{n_k} ||x_{ki} - \mu_k||^2$$

- Whenever an assignment is changed, the sum squared distances J of data points from their assigned cluster centers is reduced.
- Whenever a cluster center is moved, J is reduced.
- Test for convergence: If the assignments do not change in the assignment step, we have converged (to at least a local minimum).



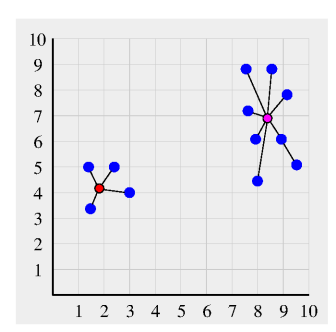
Why K-means Works

- What is a good partition?
- High intra-cluster similarity
- K-means optimizes
 - the average distance to members of the same cluster

$$\sum_{k=1}^{K} \frac{1}{n_k} \sum_{i=1}^{n_k} \sum_{j=1}^{n_k} \left\| x_{ki} - x_{kj} \right\|^2$$

which is twice the total distance to centers,

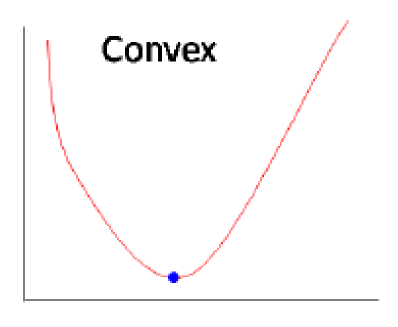
$$J = \sum_{k=1}^{K} \sum_{i=1}^{n_k} ||x_{ki} - \mu_k||^2$$



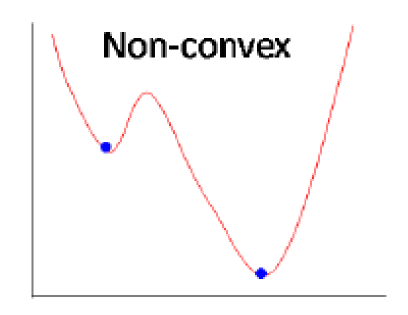
Algorithm *k-means*

- 1. Decide on a value for K, the
- 2. Initialize the *K* cluster center necessary).
- Use one of the distance / similarity functions we discussed earlier
- 3. Decide the class memberships of the *N* objects by assigning them to the nearest cluster center.
- 4. Re-estimate the *K* cluster centers, by assuming the memberships found above are correct.
- 5. Repeat 3 and 4 until none of the *N* objects changed membership in the last iteration Average / median of class members

Convex and Nonconvex functions

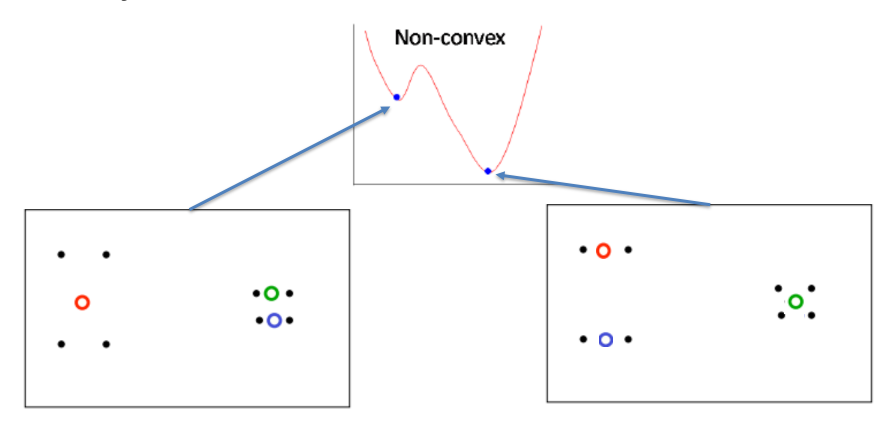


Unique minimum



Multiple minima

Objective function for k-means is non-convex



Let \vec{x}_i , i = 1, 2, ..., n be the data points and $\vec{\mu}_j$, j = 1, 2, ..., k be the k mean values.

minimize
$$\sum_{i=1}^{n} \min_{j=1..k} ||\vec{x}_i - \vec{\mu}_j||^2$$

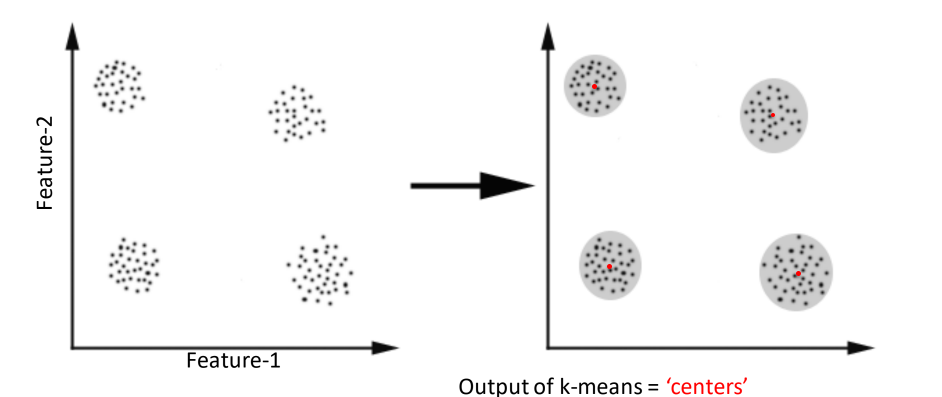
$$\min(|x_i - \mu_1|^2, |x_i - \mu_2|^2)$$

$$\max_{j=1..k} ||\vec{x}_i - \vec{\mu}_j||^2$$

K-means++: Improving K-means initialization

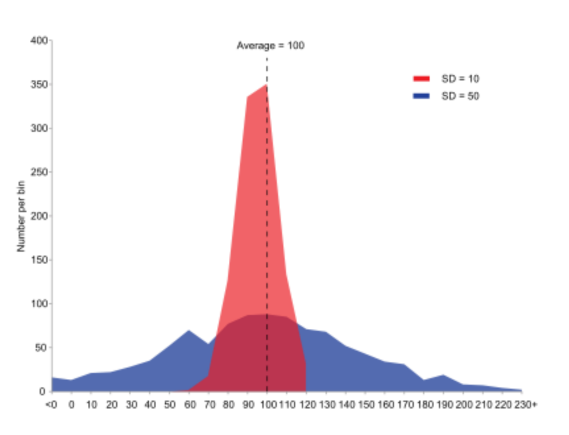
- Common way to improve k-means smart initialization!
- General idea try to get good coverage of the data.
- k-means++ algorithm:
 - 1. Pick the first center randomly
 - 2. For all points $\mathbf{x}^{(n)}$ set $d^{(n)}$ to be the distance to closest center.
 - 3. Pick the new center to be at $\mathbf{x}^{(n)}$ with probability proportional to $d^{(n)2}$
 - 4. Repeat steps 2+3 until you have k centers

Perspective: Clustering as a 'summary' of input data



... but only these are not sufficient to summarize

Mean, Standard Deviation and Variance (1-D)



Encodes spread wrt mean

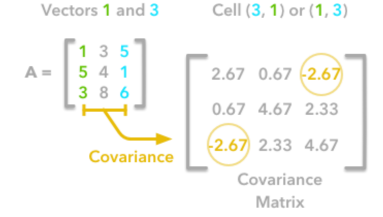
Sample Variance

$$s^2 = \frac{\sum (x - \bar{x})^2}{n - 1}$$

Sample Standard Deviation

$$s^{2} = \frac{\sum (x - \bar{x})^{2}}{n - 1}$$
 $s = \sqrt{\frac{\sum (x - \bar{x})^{2}}{n - 1}}$

Covariance



Variance:
$$s^{2} = \frac{\sum (\overline{X} - X_{i})^{2}}{N}$$

$$cov(\boldsymbol{X}, \boldsymbol{Y}) = \frac{1}{n} \sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})$$

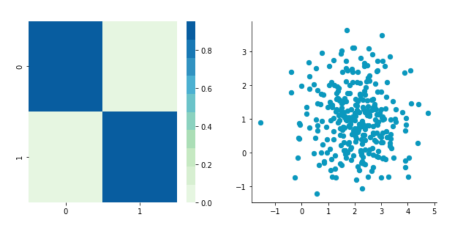
$$Cov(M_a, M_b) = \frac{1}{m} \sum_{i=1}^{m} (q_{i,a} - \overline{q_a})(q_{i,b} - \overline{q_b})$$

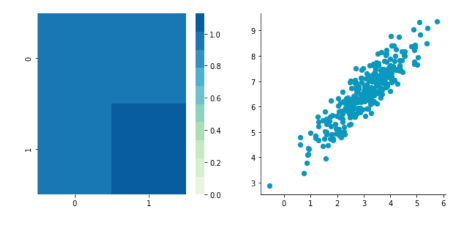
$$C = \begin{pmatrix} cov(M_1, M_1) & cov(M_1, M_2) & \cdots & cov(M_1, M_n) \\ cov(M_2, M_1) & cov(M_2, M_2) & \cdots & cov(M_2, M_n) \\ \vdots & \vdots & \ddots & \vdots \\ cov(M_n, M_1) & cov(M_n, M_2) & \cdots & cov(M_n, M_n) \end{pmatrix}_{\text{non}}$$

n-dimensional Covariance Matrix

Covariance

$$cov(\boldsymbol{X}, \boldsymbol{Y}) = \frac{1}{n} \sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})$$

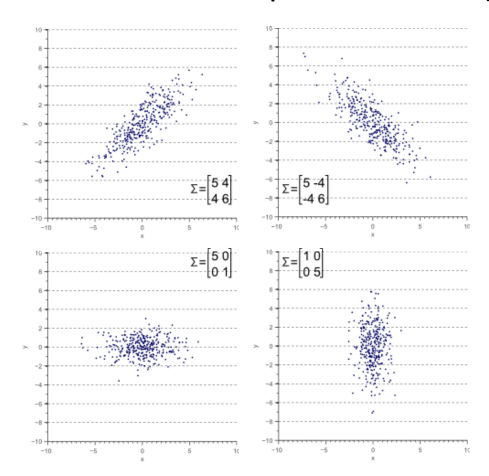




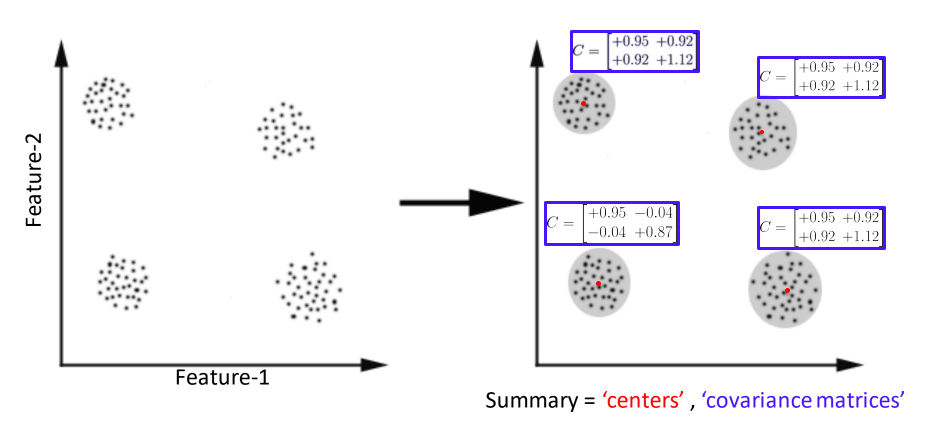
$$C = \begin{bmatrix} +0.95 & -0.04 \\ -0.04 & +0.87 \end{bmatrix}$$

$$C = \begin{bmatrix} +0.95 & +0.92 \\ +0.92 & +1.12 \end{bmatrix}$$

Covariance characterizes spread of data (wrt mean)



Perspective: Clustering as a 'summary' of input data version 2



K-means: Additional issues

- 'Hard' assignments
- Euclidean Favors 'Spherical' clusters
 - Cluster-distribution-adaptive ?