

Expectation-Maximization (EM)

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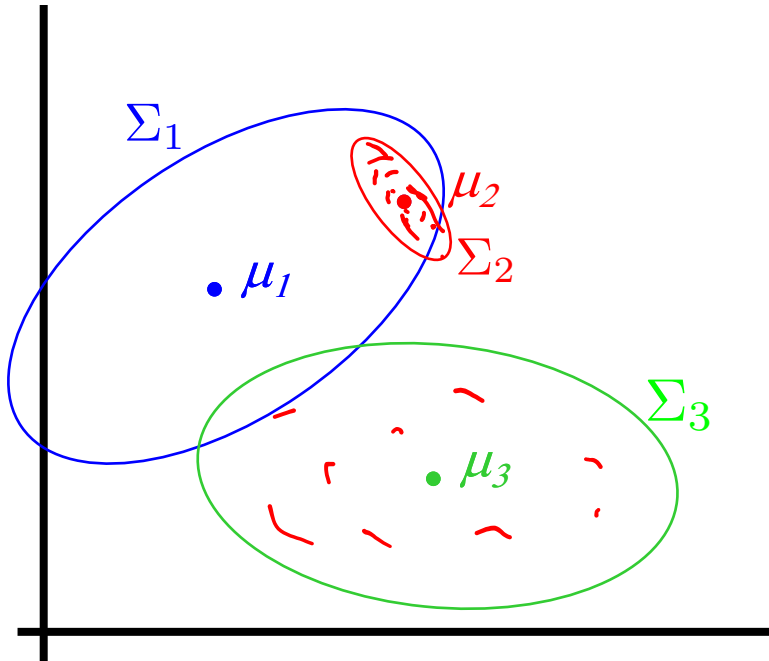
Machine Learning 10-315
Nov 17, 2021



Some slides courtesy of Eric Xing, Carlos Guestrin

Mixture models (Gaussian)

$$\underbrace{x_1, \dots, x_m}_{\text{data}} \sim \underbrace{p(x)} = \sum_{i=1}^k \underbrace{p(x|Y=i)}_{\text{Mixture component}} \underbrace{P(Y=i)}_{\text{Mixture proportion, } p_i}$$



Gaussian mixture model

$$\underbrace{p(x|Y=i)} \sim \mathcal{N}(\underbrace{\mu_i}, \underbrace{\Sigma_i})$$

Parameters: $\underbrace{\{p_i, \mu_i, \Sigma_i\}_{i=1}^K}$

- How to estimate parameters? Max Likelihood
But don't know labels Y (recall Gaussian Bayes classifier)

Expectation-Maximization (EM)

A general algorithm to deal with hidden data, but we will study it in the context of unsupervised learning (hidden labels)

- No need to choose step size as in Gradient methods. ✓
- EM is an Iterative algorithm with two linked steps:
 - E-step: fill-in hidden data (Y) using inference ✓
 - M-step: apply standard MLE/MAP method to estimate parameters $\{\mu_i, \Sigma_i\}_{i=1}^k$ ✓
- This procedure monotonically improves the marginal likelihood of observed data (or leaves it unchanged). Thus it always converges to a local optimum of the likelihood.

EM for spherical, same variance GMMs

same mixture proportions

Initialize: $\mu_1, \mu_2, \dots, \mu_K$ randomly

$$\{\tilde{p}_i, \mu_i, \tilde{\Sigma}_i\}_{i=1}^K$$

$\underbrace{\tilde{p}_i}_{\frac{1}{K}} \quad \uparrow \quad \underbrace{\tilde{\Sigma}_i}_{\sigma^2 I}$

E-step

Compute “expected” classes of all datapoints for each class

In K-means “E-step”
we do hard assignment

EM does soft assignment

$$\begin{aligned}
 & \xrightarrow{i=1, \dots, K} P(y=i | x_j, \mu_1, \dots, \mu_K) \propto \underbrace{\exp\left(-\frac{1}{2\sigma^2} \|x_j - \mu_i\|^2\right)}_{p(x_j | y=i)} \underbrace{P(y=i)}_{1/K} \\
 & \propto \underbrace{p(x_j | y)}_{p(x_j | y=i)} \underbrace{P(y)}_{1/K}
 \end{aligned}$$

M-step

Compute Max. like μ given our data’s class membership distributions (weights)

$$\mu_i = \frac{\sum_{j=1}^m P(y=i | x_j) x_j}{\sum_{j=1}^m P(y=i | x_j)} \quad \text{with weights } w_j$$

$$\sum_{j=1}^m w_j x_j$$

Exactly same as MLE with weighted data

Iterate.

EM for general GMMs

Iterate. On iteration t let our estimates be

$$\rightarrow \lambda_t = \{ \underbrace{\mu_1^{(t)}, \mu_2^{(t)} \dots \mu_k^{(t)}}_{\text{means}}, \underbrace{\Sigma_1^{(t)}, \Sigma_2^{(t)} \dots \Sigma_k^{(t)}}_{\text{covariances}}, \underbrace{p_1^{(t)}, p_2^{(t)} \dots p_k^{(t)}}_{\text{weights}} \}$$

$p_i^{(t)}$ is shorthand for estimate of $P(y=i)$ on t 'th iteration

E-step

Compute "expected" classes of all datapoints for each class

Just evaluate a Gaussian at x_j

$$\begin{matrix} i=1 \dots K \\ j=1 \dots m \end{matrix} \rightarrow \underbrace{P(y=i|x_j, \lambda_t)}_{\propto p(x_j|y=i)} \propto \underbrace{p_i^{(t)}}_{P(y=i)} \underbrace{p(x_j|\mu_i^{(t)}, \Sigma_i^{(t)})}_{p(x_j|y=i)}$$

M-step

Compute MLEs given our data's class membership distributions (weights)

$$\mu_i^{(t+1)} = \frac{\sum_j P(y=i|x_j, \lambda_t) x_j}{\sum_j P(y=i|x_j, \lambda_t)} \quad \Sigma_i^{(t+1)} = \frac{\sum_j P(y=i|x_j, \lambda_t) (x_j - \mu_i^{(t+1)}) (x_j - \mu_i^{(t+1)})^T}{\sum_j P(y=i|x_j, \lambda_t)}$$

$$p_i^{(t+1)} = \frac{\sum_j P(y=i|x_j, \lambda_t)}{m}$$

m = #data points

$$\boxed{\frac{m_i}{m}}$$

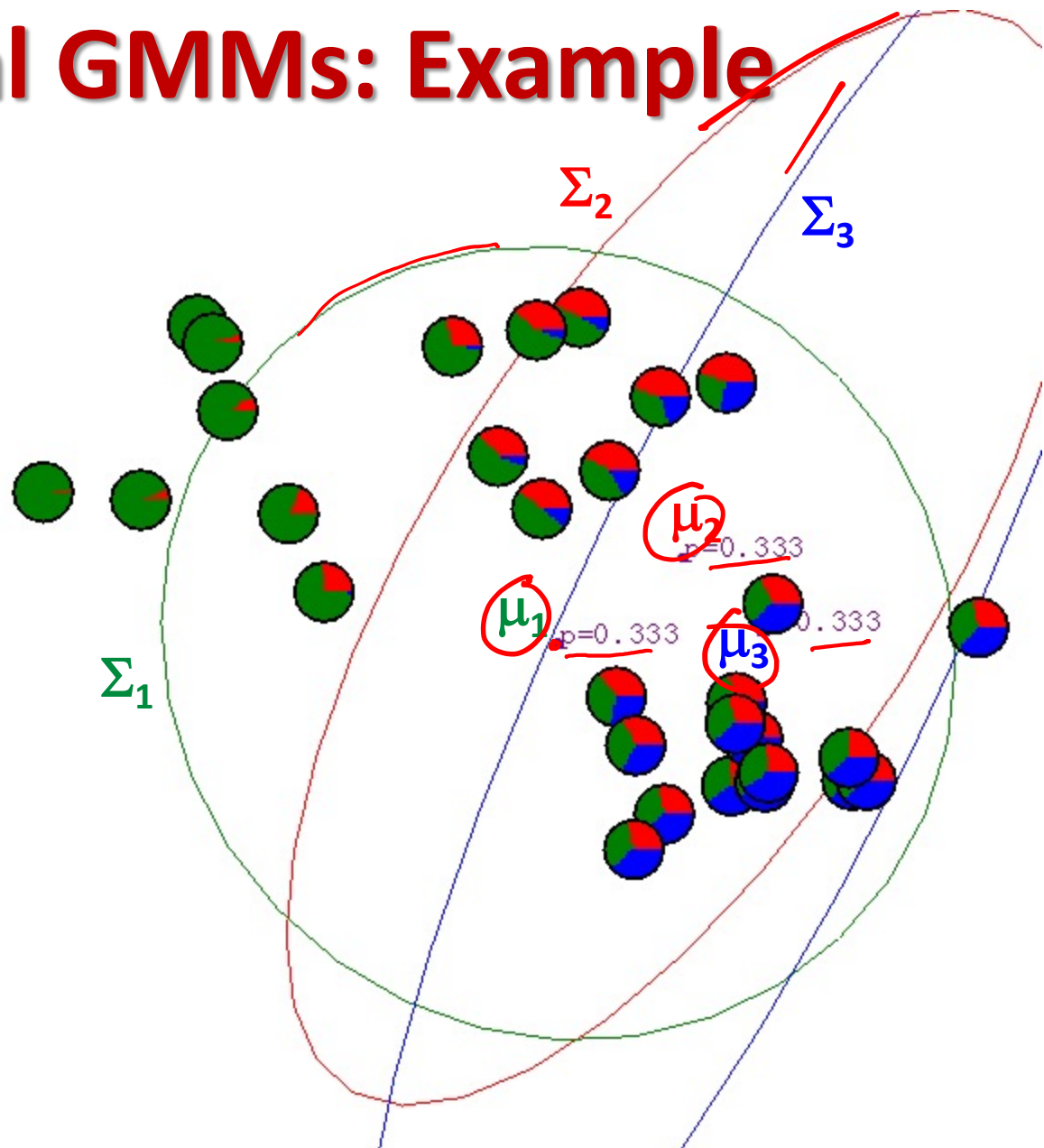
EM for general GMMs: Example

$K=3$

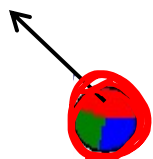
p_1, p_2, p_3

μ_1, μ_2, μ_3

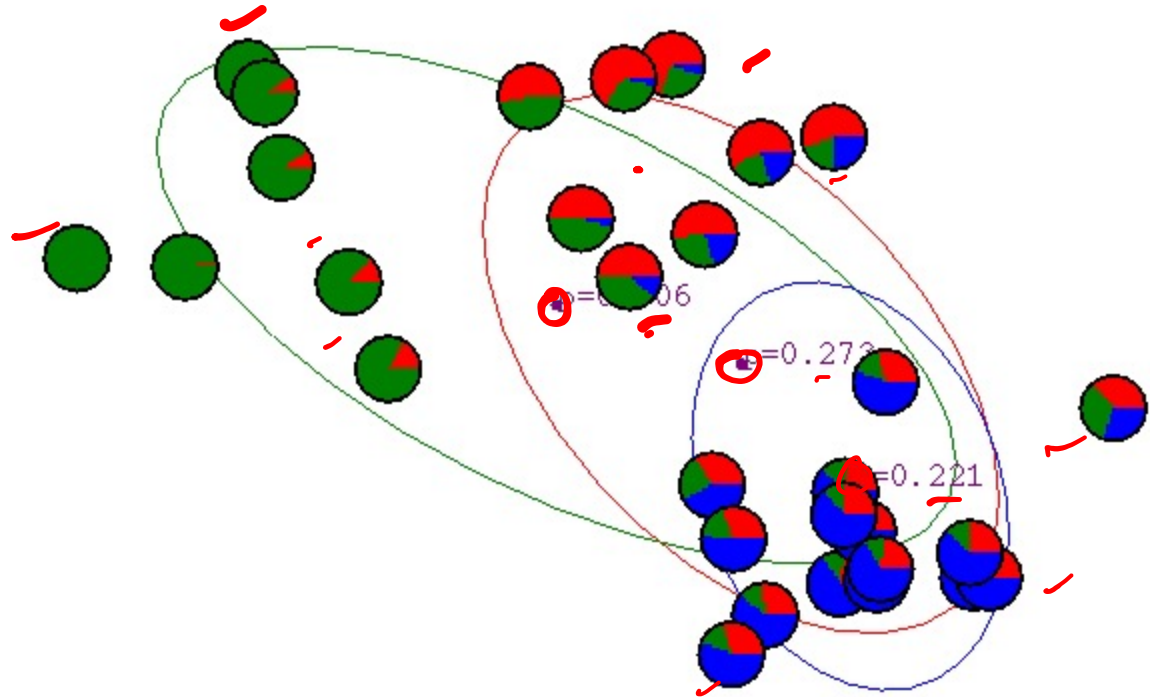
$\Sigma_1, \Sigma_2, \Sigma_3$



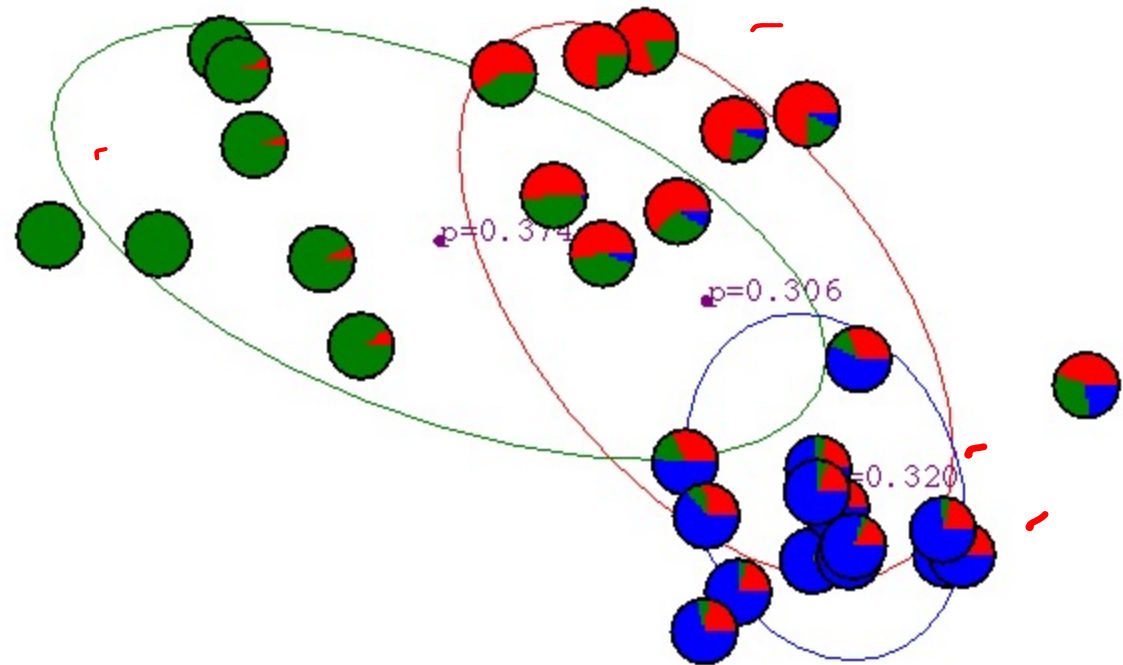
$$P(y = \bullet | x_j, \mu_1, \mu_2, \mu_3, \Sigma_1, \Sigma_2, \Sigma_3, p_1, p_2, p_3)$$



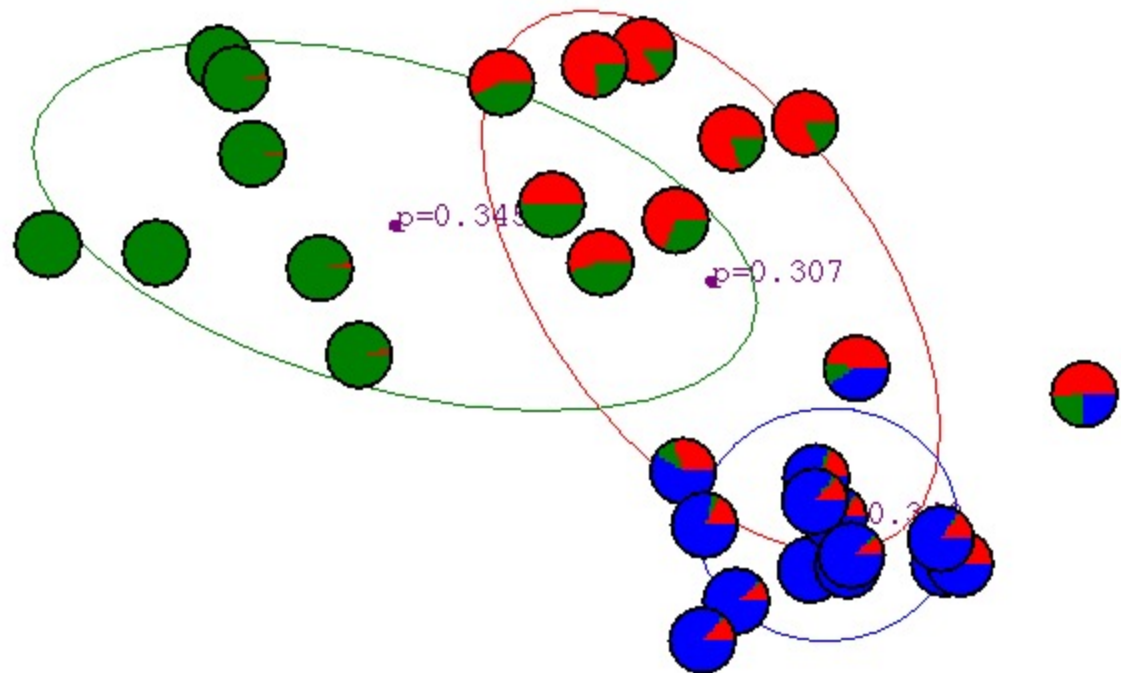
After 1st iteration



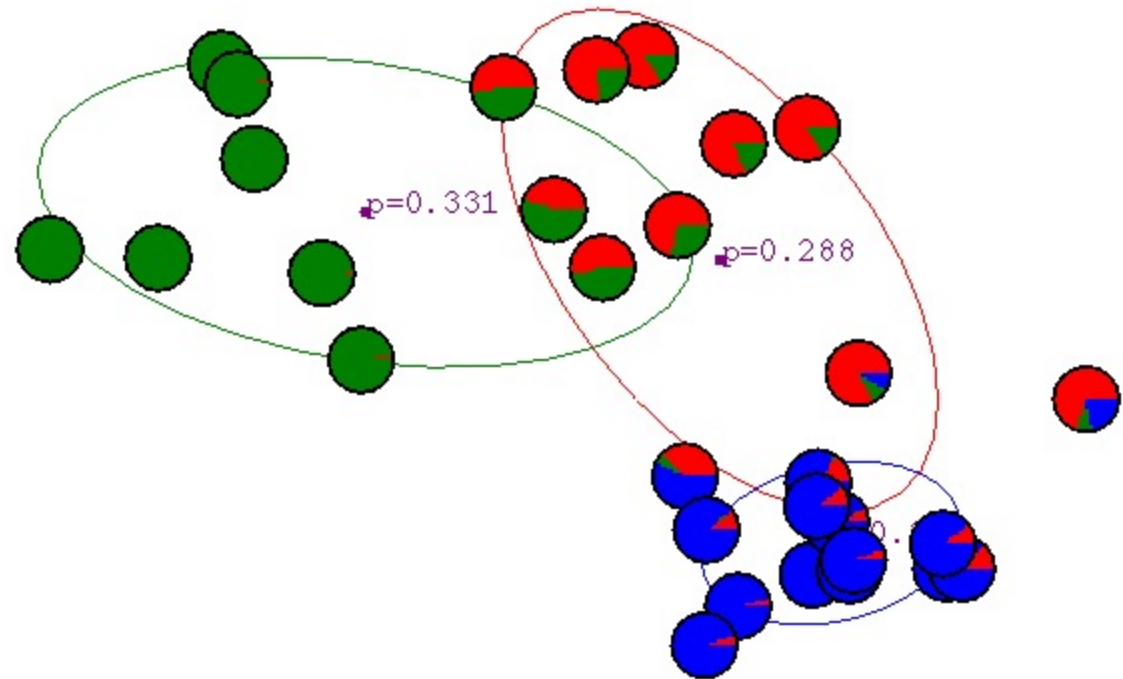
After 2nd iteration



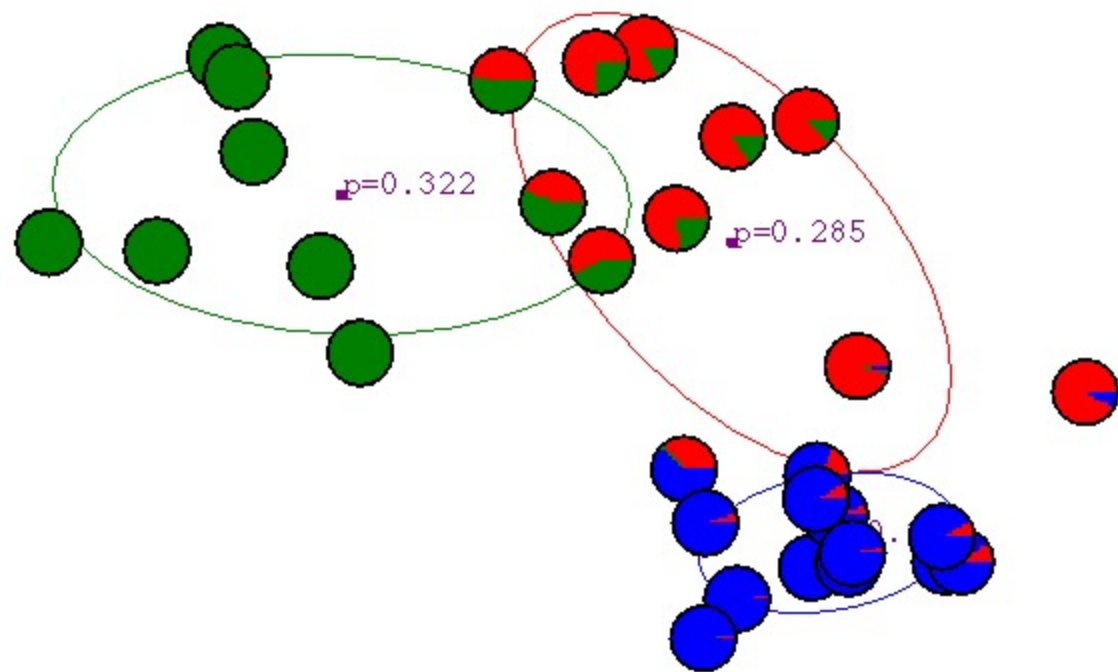
After 3rd iteration



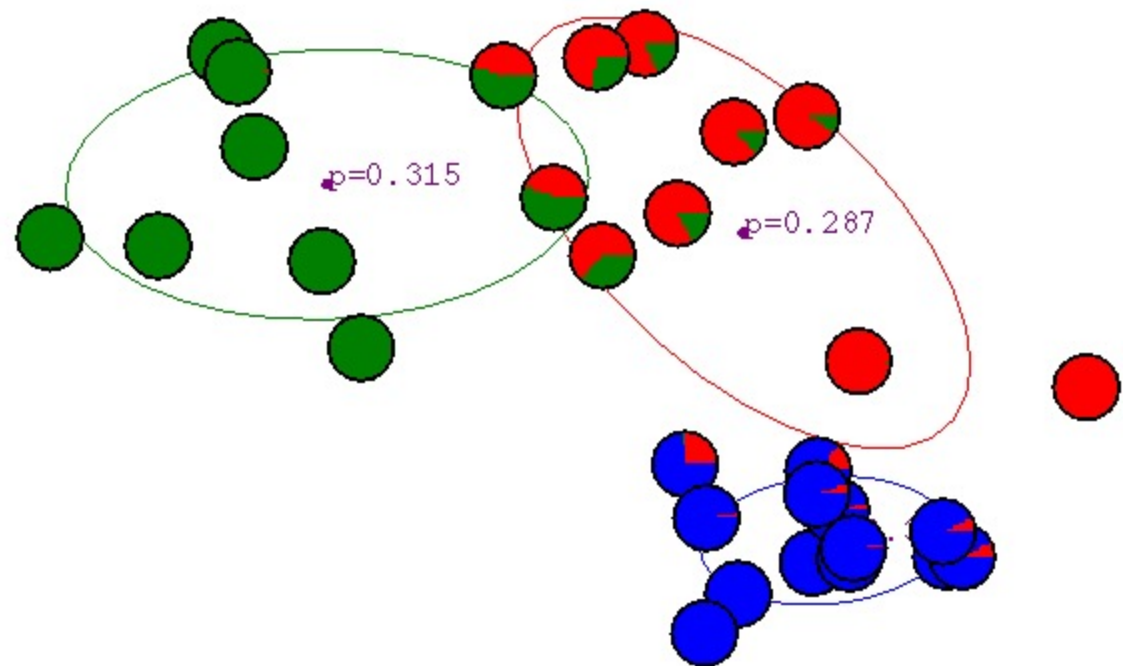
After 4th iteration



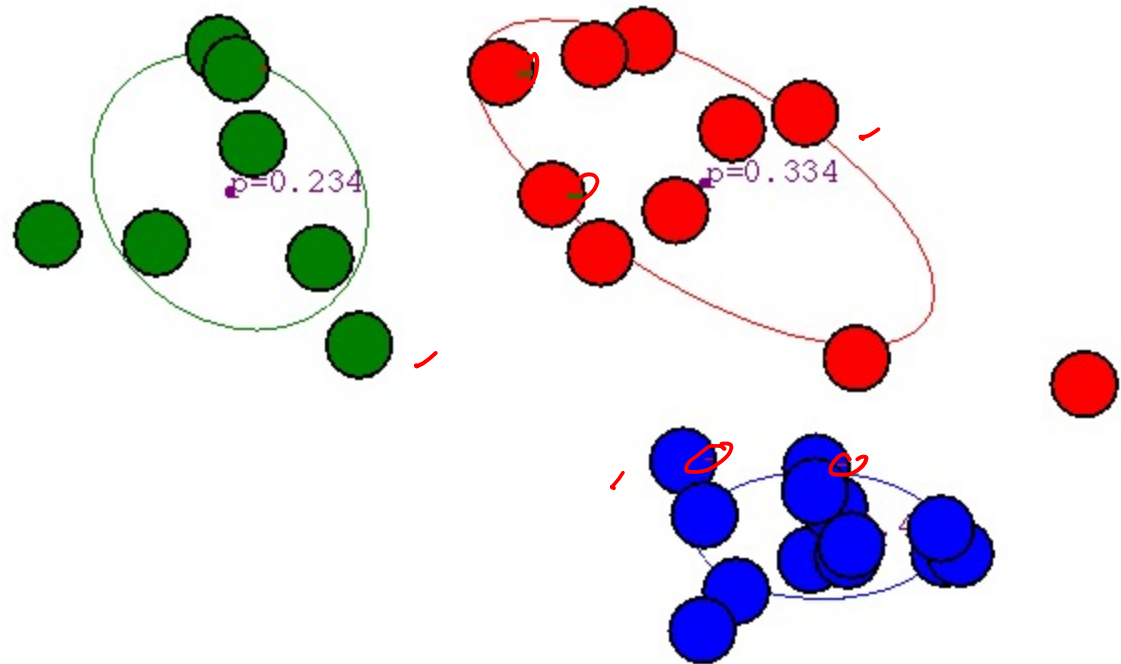
After 5th iteration



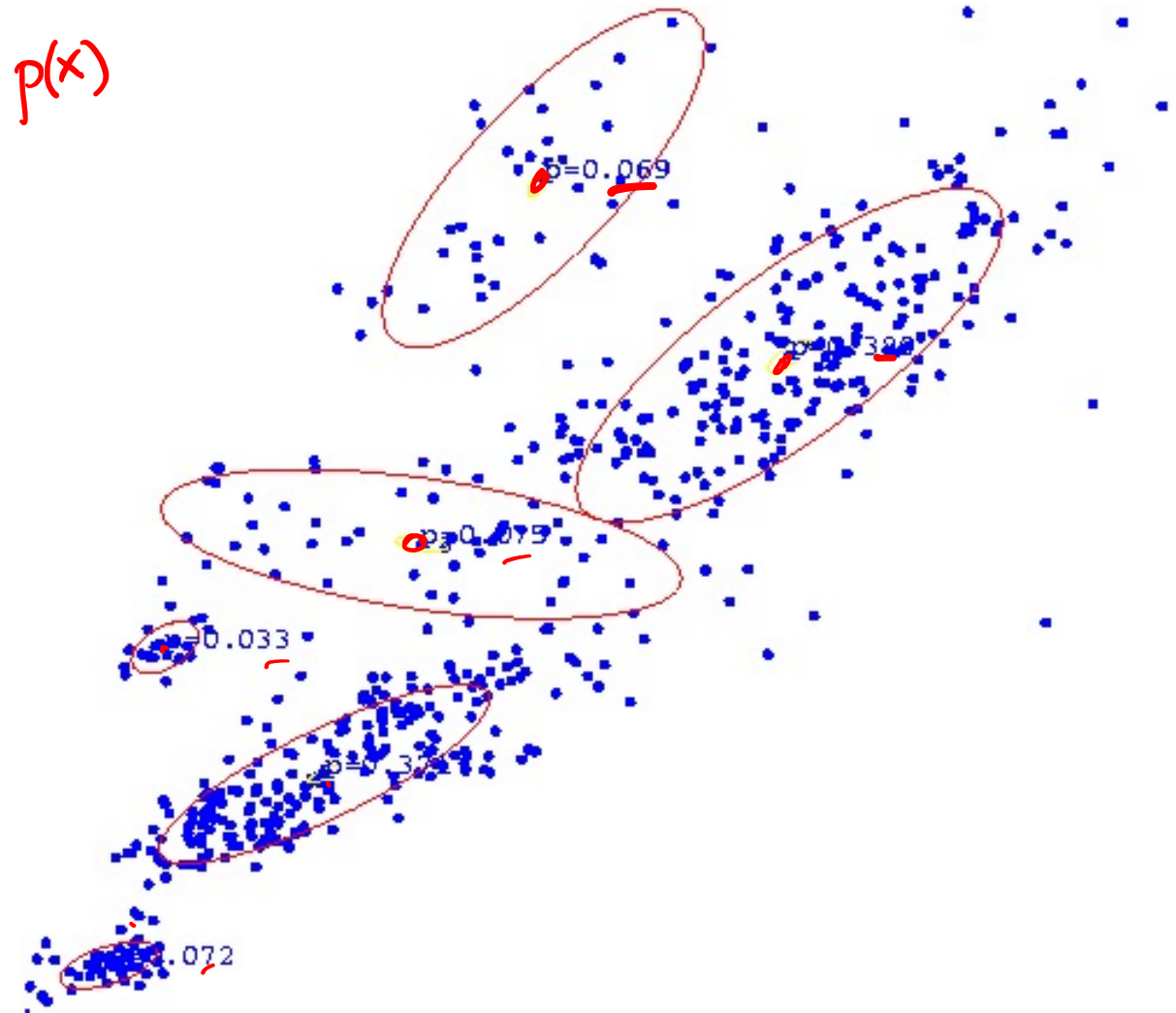
After 6th iteration



After 20th iteration



GMM clustering of assay data



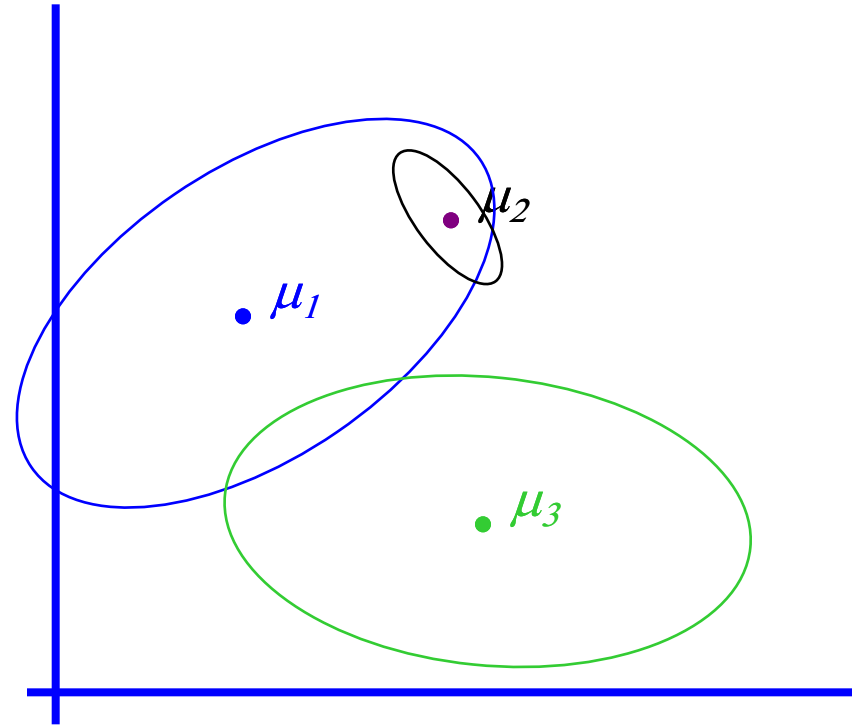
General GMM

GMM – Gaussian Mixture Model (Multi-modal distribution)

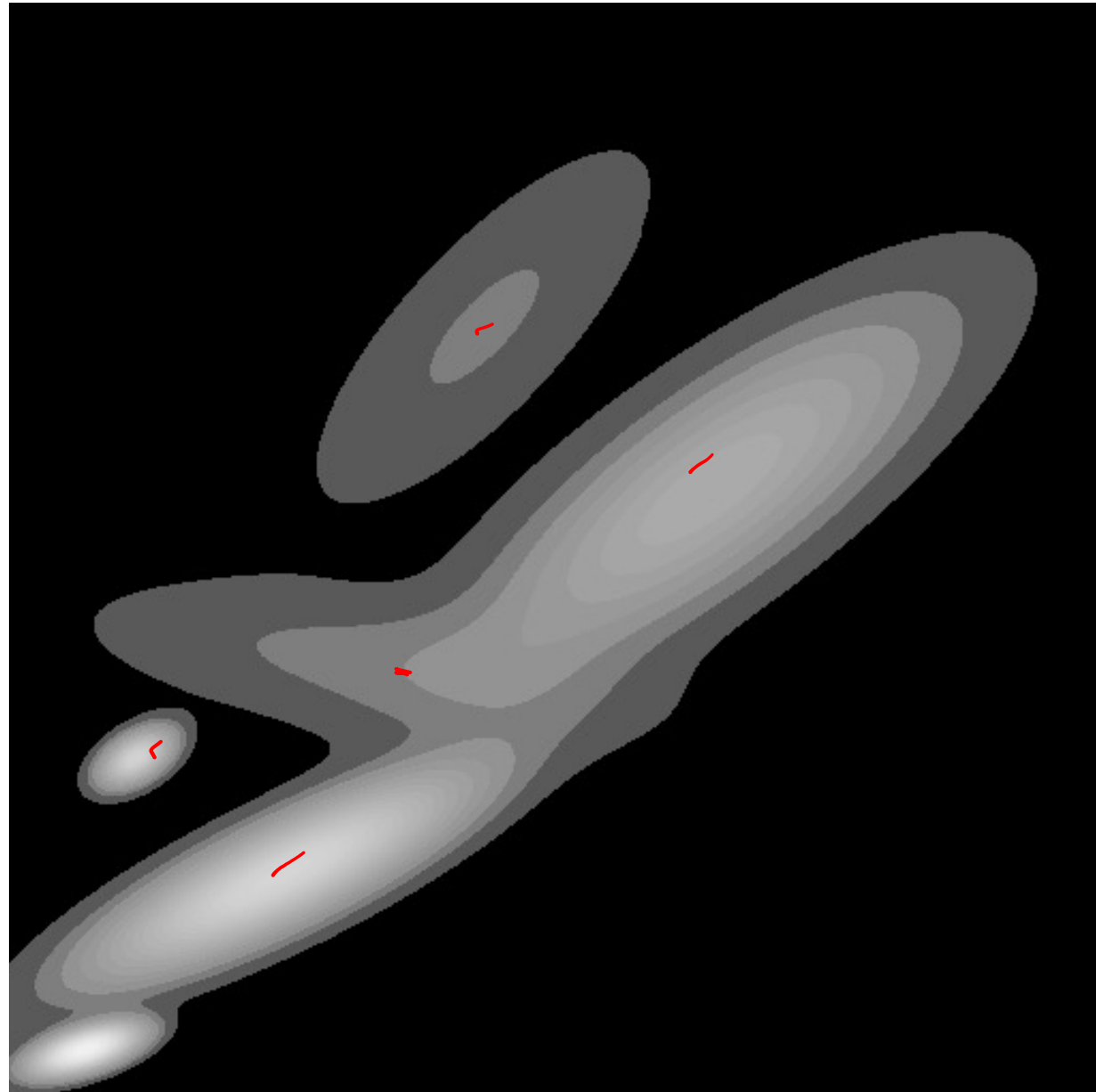
$$p(x) = \sum_i p(x|y=i) P(y=i)$$

↓ ↓
Mixture **Mixture**
component **proportion**

$$p(x|y=i) \sim N(\mu_i, \Sigma_i)$$



Resulting Density Estimator



$$p(x|y=i) \sim \text{GMM}$$

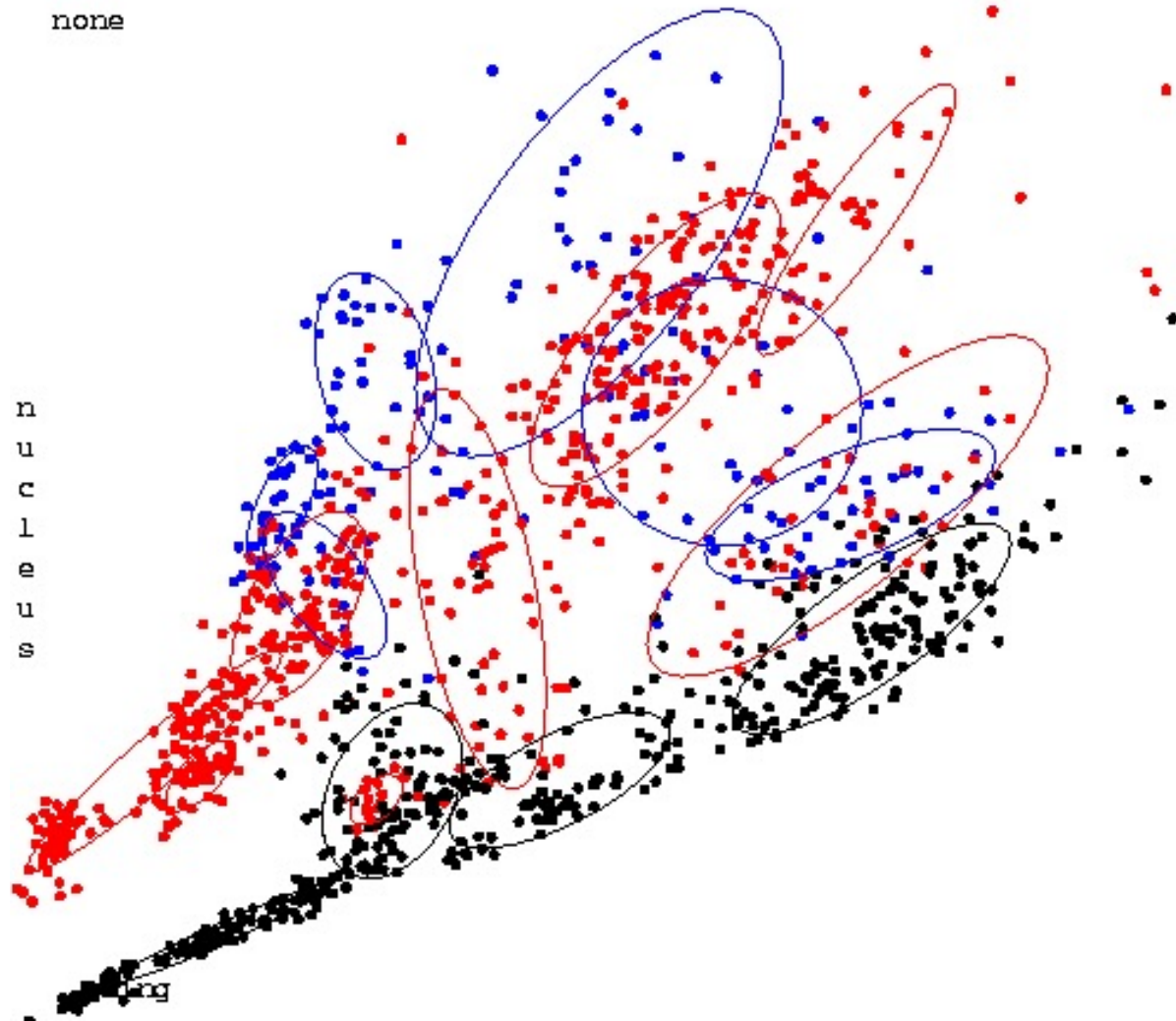
Compound =

IL-1

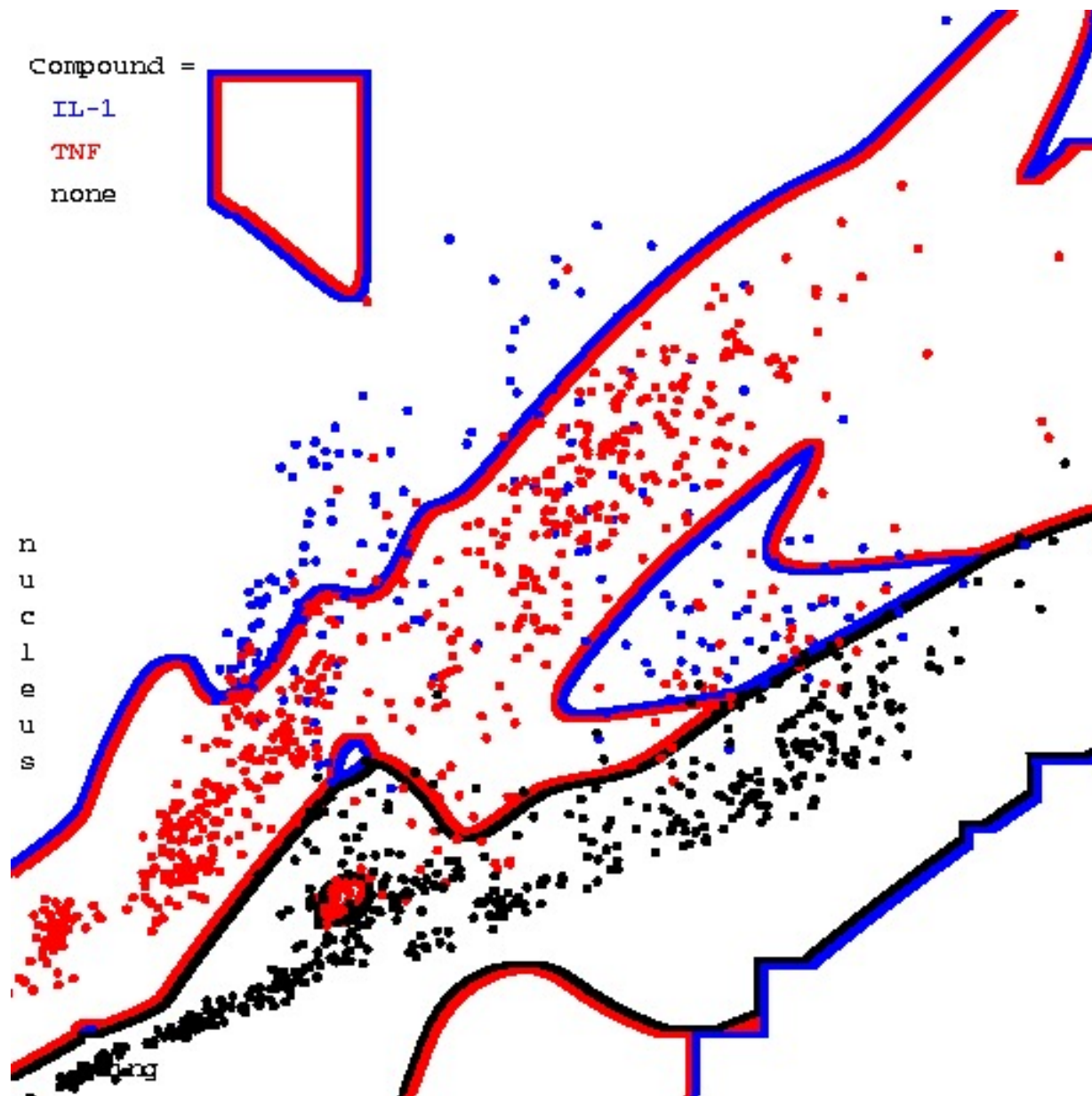
TNF

none

**Three
classes of
assay**
(each learned with
it's own mixture
model)



Resulting Bayes Classifier

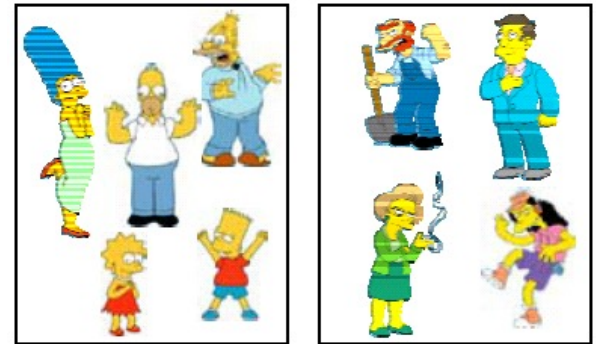


Summary: EM Algorithm

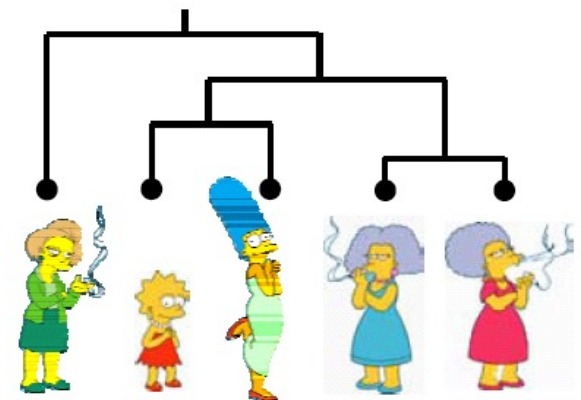
- A way of maximizing likelihood function for hidden variable models. Finds MLE of parameters when the original (hard) problem can be broken up into two (easy) pieces:
 1. Estimate some “missing” or “unobserved” data from observed data and current parameters.
 2. Using this “complete” data, find the maximum likelihood parameter estimates.
- Alternate between filling in the latent variables using the best guess (posterior) and updating the parameters based on this guess:
 1. E-step: soft cluster assignment for each data point
 2. M-step: update parameters of each mixture component
- EM can get stuck in local minima.
- BUT very popular in practice.

Clustering Algorithms

- Partition algorithms
 - K means clustering ✓
 - Mixture-Model based clustering ✓



- Hierarchical algorithms ←
 - Single-linkage
 - Average-linkage
 - Complete-linkage
 - Centroid-based



Hierarchical Clustering

- Bottom-Up Agglomerative Clustering

Starts with each object in a separate cluster, and repeat:

- Joins the most similar pair of clusters,
- Update the similarity of the new cluster to others until there is only one cluster.

Greedy - less accurate but simple to implement

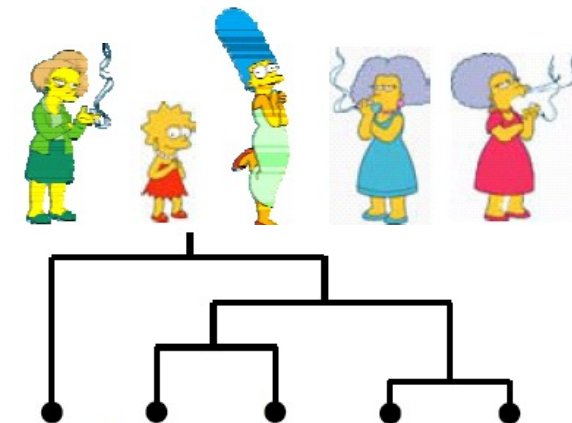


- Top-Down divisive

Starts with all the data in a single cluster, and repeat:

- Split each cluster into two using a partition algorithm
- Until each object is a separate cluster.

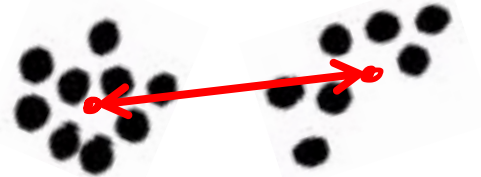
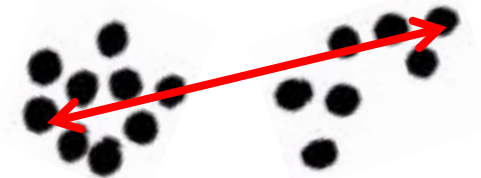
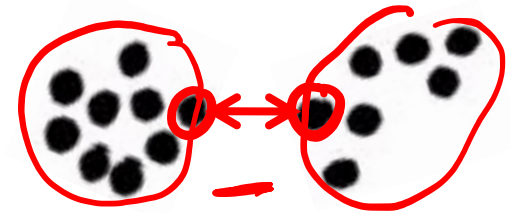
More accurate but complex to implement



Bottom-up Agglomerative clustering

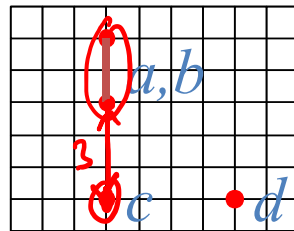
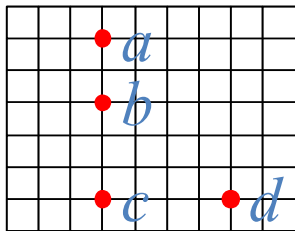
Different algorithms differ in how the similarities are defined (and hence updated) between two clusters

- Single-Linkage
 - Nearest Neighbor: similarity between their closest members.
- Complete-Linkage
 - Furthest Neighbor: similarity between their furthest members.
- Centroid
 - Similarity between the centers of gravity
- Average-Linkage
 - Average similarity of all cross-cluster pairs.

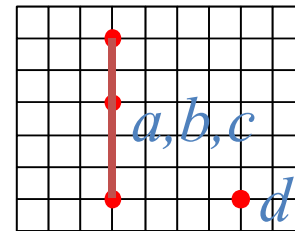


Single-Linkage Method

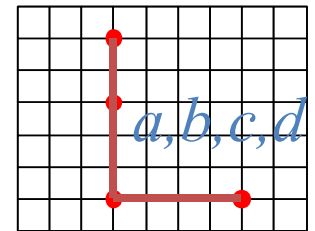
Euclidean Distance



(1)



(2)



(3)

	<i>b</i>	<i>c</i>	<i>d</i>
<i>a</i>	2	5	6
<i>b</i>		3	5
<i>c</i>			4

	<i>b</i>	<i>c</i>	<i>d</i>
<i>a</i>	2	5	6
<i>b</i>		3	5
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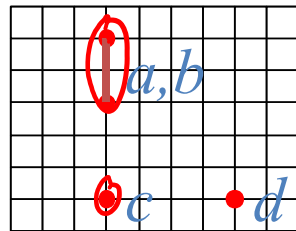
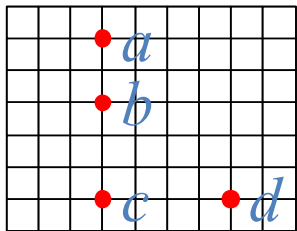
	<i>c</i>	<i>d</i>
<i>a, b</i>	3	5
<i>c</i>		4

	<i>d</i>
<i>a, b, c</i>	4

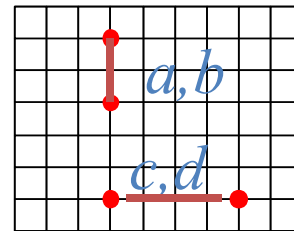
Distance Matrix

Complete-Linkage Method

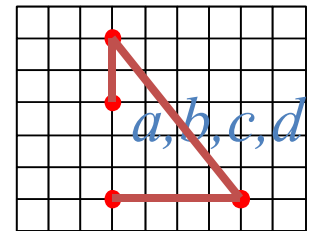
Euclidean Distance



(1)



(2)



(3)

	<i>b</i>	<i>c</i>	<i>d</i>
<i>a</i>	2	5	6
<i>b</i>		3	5
<i>c</i>			4

	<i>b</i>	<i>c</i>	<i>d</i>
<i>a</i>	2	5	6
<i>b</i>		3	5
<i>c</i>			4

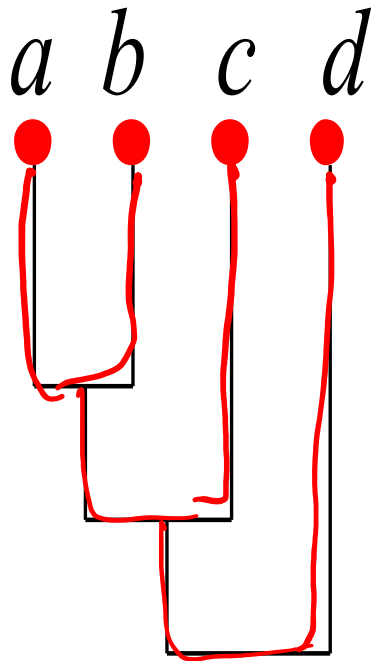
	<i>c</i>	<i>d</i>
<i>a, b</i>	5	6
<i>c</i>		4

	<i>c, d</i>
<i>a, b</i>	6

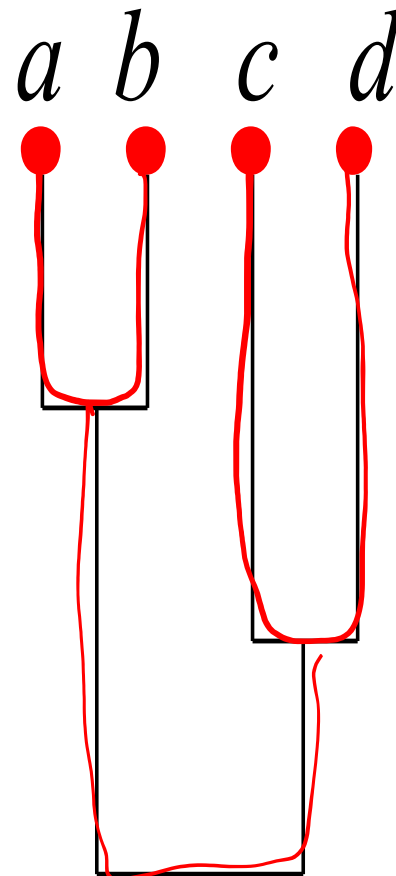
Distance Matrix

Dendrograms

Single-Linkage

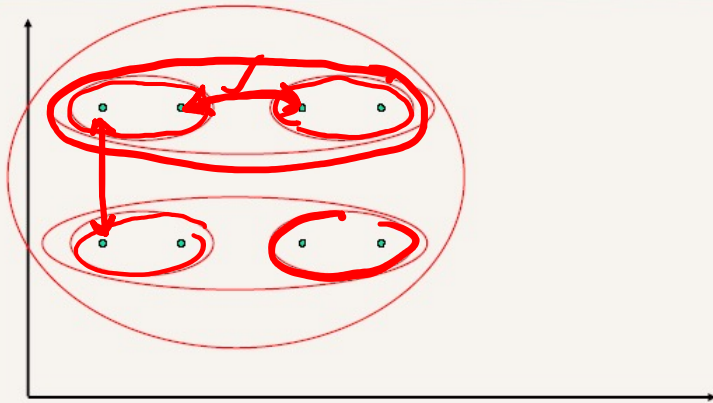


Complete-Linkage

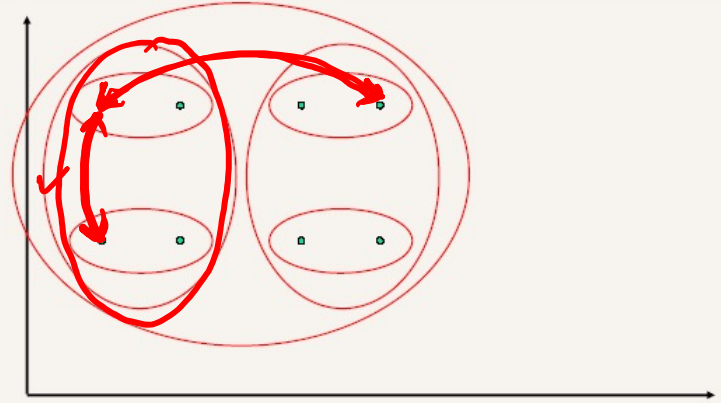


Another Example

Single Link Example



Complete Link Example



Single vs. Complete Linkage

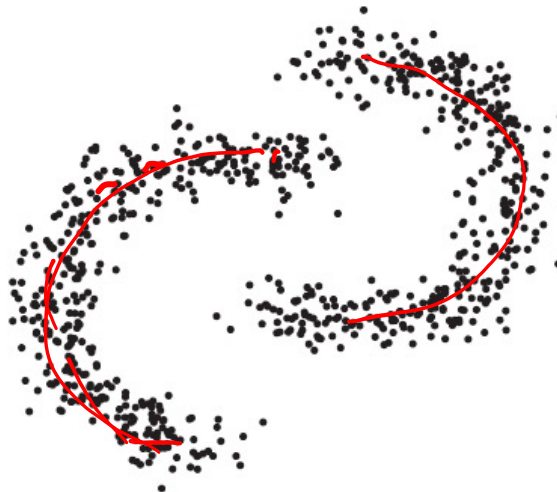
Shape of clusters

Single-linkage

allows anisotropic and
non-convex shapes

Complete-linkage

assumes isotopic, convex
shapes



Computational Complexity

- All hierarchical clustering methods need to compute similarity of all pairs of n individual instances which is $O(n^2)$.
- At each iteration,
 - Sort similarities to find largest one $O(n^2 \log n)$.
 - Update similarity between merged cluster and other clusters.
Computing similarity to each other cluster can be done in constant time.
- So we get $O(n^2 \log n)$ or $O(n^3)$ (if naïvely implemented)

Computational Complexity (K-means)

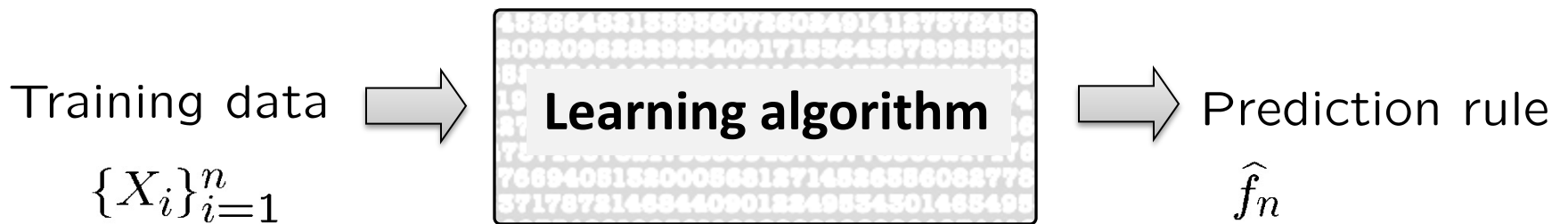
- At each iteration,
 - Computing distance between each of the n objects and the K cluster centers is $O(\underline{Kn})$.
 - Computing cluster centers: Each object gets added once to some cluster: $O(n)$.
- Assume these two steps are each done once for $\underline{l}$ iterations: $O(\underline{lKn})$.

What you need to know...

- Partition based clustering algorithms
 - K-means ✓
 - Coordinate descent
 - Seeding
 - Choosing K
 - Mixture models ✓
 - EM algorithm
- Hierarchical clustering algorithms
 - Single-linkage
 - Complete-linkage
 - Centroid-linkage
 - Average-linkage

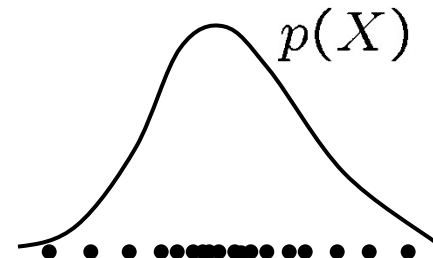
Unsupervised Learning

“Learning from unlabeled/unannotated data” (without supervision)



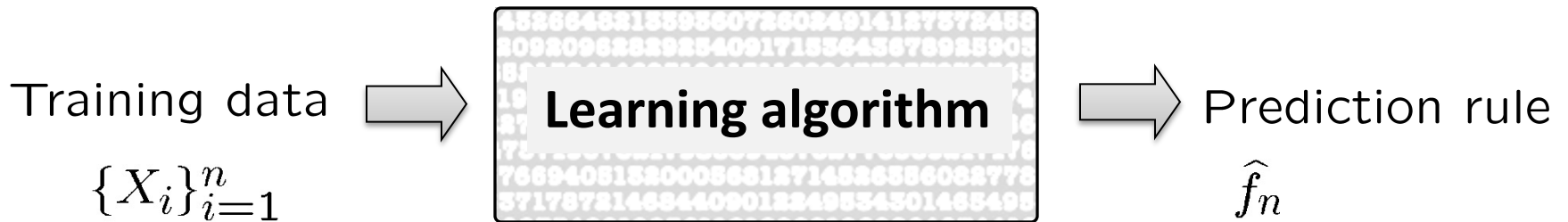
What can we predict from unlabeled data?

- Density estimation



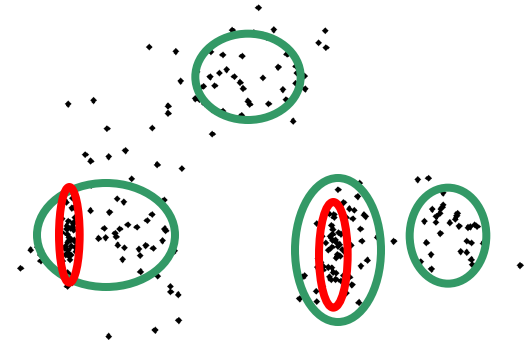
Unsupervised Learning

“Learning from unlabeled/unannotated data” (without supervision)



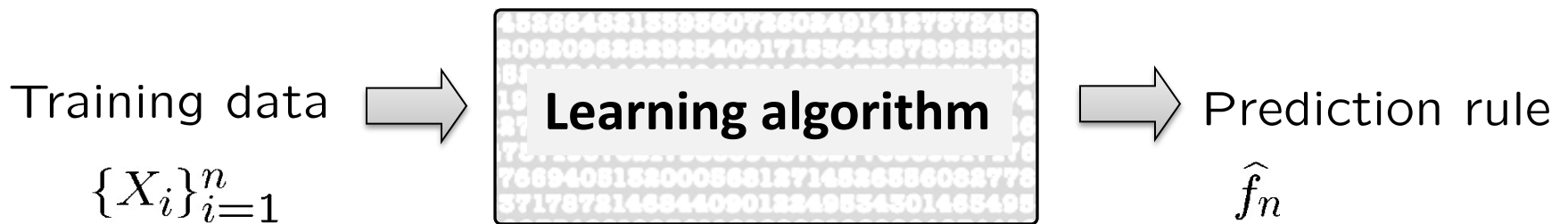
What can we predict from unlabeled data?

- Density estimation
- Groups or clusters in the data



Unsupervised Learning

“Learning from unlabeled/unannotated data” (without supervision)



What can we predict from unlabeled data?

- Density estimation
- Groups or clusters in the data
- Dimensionality reduction

