# **ML:Clustering**

#### From Coursera

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# **Unsupervised Learning: Introduction**

Unsupervised learning is contrasted from supervised learning because it uses an **unlabeled** training set rather than a labeled one.

In other words, we don't have the vector y of expected results, we only have a dataset of features where we can find structure.

Clustering is good for:

- Market segmentation
- Social network analysis
- Organizing computer clusters
- Astronomical data analysis

## **K-Means Algorithm**

The K-Means Algorithm is the most popular and widely used algorithm for automatically grouping data into coherent subsets.

- 1. Randomly initialize two points in the dataset called the *cluster centroids*.
- 2. Cluster assignment: assign all examples into one of two groups based on which cluster centroid the example is closest to.
- 3. Move centroid: compute the averages for all the points inside each of the two cluster centroid groups, then move the cluster centroid points to those averages.
- 4. Re-run (2) and (3) until we have found our clusters.

Our main variables are:

$$K$$
 (number of clusters)

Training set 
$$x^{(1)}, x^{(2)}, \dots, x^{(m)}$$

Where 
$$x^{(i)} \in \mathbb{R}^n$$

Note that we will not use the  $x_0 = 1$  convention.

#### The algorithm:

```
Randomly initialize K cluster centroids mu(1), mu(2), ..., mu(K)
Repeat:
    for i = 1 to m:
        c(i) := index (from 1 to K) of cluster centroid closest to x(i)
    for k = 1 to K:
        mu(k) := average (mean) of points assigned to cluster k
```

The **first for-loop** is the 'Cluster Assignment' step. We make a vector c where c(i) represents the centroid assigned to example x(i).

We can write the operation of the Cluster Assignment step more mathematically as follows:

$$|c^{(i)} = argmin_k \mid \mid x^{(i)} - \mu_k \mid \mid^2$$

That is, each  $c^{(i)}$  contains the index of the centroid that has minimal distance to  $x^{(i)}$ .

By convention, we square the right-hand-side, which makes the function we are trying to minimize more sharply increasing. It is mostly just a convention. But a convention that helps reduce the computation load because the Euclidean distance requires a square root but it is canceled.

Without the square:

$$||x^{(i)} - \mu_k|| = || \sqrt{(x_1^i - \mu_{1(k)})^2 + (x_2^i - \mu_{2(k)})^2 + (x_3^i - \mu_{3(k)})^2 + \dots} ||$$

With the square:

$$||x^{(i)} - \mu_k||^2 = || \;\; (x_1^i - \mu_{1(k)})^2 + (x_2^i - \mu_{2(k)})^2 + (x_3^i - \mu_{3(k)})^2 + \dots \;\; |$$

...so the square convention serves two purposes, minimize more sharply and less computation.

The **second for-loop** is the 'Move Centroid' step where we move each centroid to the average of its group.

More formally, the equation for this loop is as follows:

$$\mu_k = rac{1}{n} \left[ x^{(k_1)} + x^{(k_2)} + \ldots + x^{(k_n)} 
ight] \in \mathbb{R}^n$$

Where each of  $x^{(k_1)}, x^{(k_2)}, \ldots, x^{(k_n)}$  are the training examples assigned to group  $\mu_k$ .

If you have a cluster centroid with **0 points** assigned to it, you can randomly **re-initialize** that centroid to a new point. You can also simply **eliminate** that cluster group.

After a number of iterations the algorithm will converge, where new iterations do not affect the clusters.

Note on non-separated clusters: some datasets have no real inner separation or natural structure. K-means can still evenly segment your data into K subsets, so can still be useful in this case.

# **Optimization Objective**

Recall some of the parameters we used in our algorithm:

 $c^{(i)} = \text{index of cluster } (1,2,...,K) \text{ to which example } x^{(i)} \text{ is currently assigned}$ 

$$\mu_k=$$
 cluster centroid  $k$   $(\mu_k\in\mathbb{R}^n)$ 

 $\mu_{c^{(i)}}=$  cluster centroid of cluster to which example  $x^{(i)}$  has been assigned

Using these variables we can define our **cost function**:

$$J(c^{(i)},\dots,c^{(m)},\mu_1,\dots,\mu_K) = rac{1}{m}\sum_{i=1}^m ig|ig|x^{(i)} - \mu_{c^{(i)}}ig|ig|^2$$

Our optimization objective is to minimize all our parameters using the above cost function:

$$min_{c,\mu} J(c,\mu)$$

That is, we are finding all the values in sets c, representing all our clusters, and  $\mu$ , representing all our centroids, that will minimize **the average of the distances** of every training example to its corresponding cluster centroid.

The above cost function is often called the **distortion** of the training examples.

In the cluster assignment step, our goal is to:

Minimize 
$$J(\ldots)$$
 with  $c^{(1)},\ldots,c^{(m)}$  (holding  $\mu_1,\ldots,\mu_K$  fixed)

In the **move centroid** step, our goal is to:

Minimize 
$$J(\ldots)$$
 with  $\mu_1, \ldots, \mu_K$ 

With k-means, it is not possible for the cost function to sometimes increase. It should always descend.

### **Random Initialization**

There's one particular recommended method for randomly initializing your cluster centroids.

- 1. Have K < m. That is, make sure the number of your clusters is less than the number of your training examples.
- 2. Randomly pick K training examples. (Not mentioned in the lecture, but also be sure the selected examples are unique).
- 3. Set  $\mu_1, \ldots, \mu_k$  equal to these K examples.

K-means can get stuck in local optima. To decrease the chance of this happening, you can run the algorithm on many different random initializations. In cases where K < 10 it is strongly recommended to run a loop of random initializations.

```
for i = 1 to 100:
randomly initialize k-means
run k-means to get 'c' and 'm'
compute the cost function (distortion) J(c,m)
pick the clustering that gave us the lowest cost
```

# **Choosing the Number of Clusters**

Choosing K can be quite arbitrary and ambiguous.

The elbow method: plot the cost J and the number of clusters K. The cost function should reduce as we increase the number of clusters, and then flatten out. Choose K at the point where the cost function starts to flatten out.

However, fairly often, the curve is very gradual, so there's no clear elbow.

Note: J will always decrease as K is increased. The one exception is if k-means gets stuck at a bad local optimum.

Another way to choose K is to observe how well k-means performs on a **downstream purpose**. In other words, you choose K that proves to be most useful for some goal you're trying to achieve from using these clusters.

## **Bonus: Discussion of the drawbacks of K-Means**

From StackExchange (http://stats.stackexchange.com/questions/133656/how-to-understand-the-drawbacks-of-k-means) This links to a discussion that shows various situations in which K-means gives totally correct but unexpected results.

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