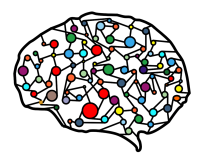
Lecture 14 - Clustering



DSC 40A, Spring 2023

Announcements

- Midterm 1 is Friday during lecture.
- No groupwork tonight. Instead, TA and tutors will host a mock exam and review session today from 7-9pm in FAH 1301.
 - Note the room change (same building).
 - You'll take the midterm from Winter 2022, when I last taught this class.
 - Bring paper. Formula sheet and exam questions provided.

Midterm 1 is Friday during lecture

- Formula sheet will be provided for you. No other notes.
- No calculators. This implies no crazy calculations.
- Assigned seats will be posted on Campuswire.
- We will not answer questions during the exam. State your assumptions if anything is unclear.
- ► The exam will include long-answer homework-style questions, as well as short-answer questions such as multiple choice or filling in a numerical answer.
- ► The exam covers Homeworks 1 through 4, which includes Monday's lecture, but not clustering.

Midterm study strategy

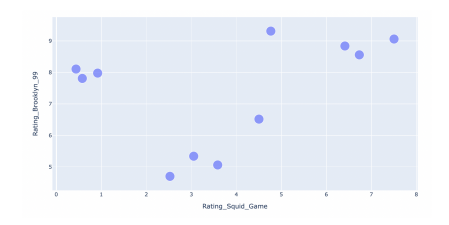
- Review the written solutions to previous homeworks and groupworks.
- Identify which concepts are still iffy. Re-watch podcasts, post on Campuswire, come to office hours, use resources on course website.
- Work through past exams on course website.
- Study in groups.
- Summarize key facts and formulas.

Agenda

- ► The clustering problem.
- k-Means Clustering algorithm.
- Why does k-Means work?
- Practical considerations.

The clustering problem

Question: how might we "cluster" these points into groups?



Problem statement: clustering

Goal: Given a list of n data points, stored as vectors in \mathbb{R}^d , $\vec{x}_1, \vec{x}_2, ..., \vec{x}_n$, and a positive integer k, place the data points into k groups of nearby points.

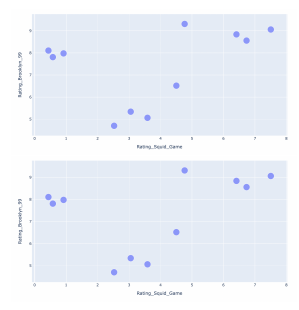
- These groups are called "clusters".
- Think about groups as colors.
 - i.e., the goal of clustering is to assign each point a color, such that points of the same color are close to one another.
- Note, unlike with regression, there is no "right answer" that we are trying to predict there is no y!
 - Clustering is an unsupervised method.

How do we define a group?

One solution: pick k cluster centers, i.e. centroids:

$$\vec{\mu}_1, \vec{\mu}_2, ..., \vec{\mu}_k$$
 in \mathbb{R}^d

- ► These *k* centroids define the *k* groups.
- Each data point "belongs" to the group corresponding to the nearest centroid.
- ► This reduces our problem from being "find the best group for each data point" to being "find the best locations for the centroids".



How do we pick the centroids?

- Let's come up with an **cost function**, *C*, which describes how good a set of centroids is.
 - Cost functions are a generalization of empirical risk functions
- One possible cost function:

$$C(\mu_1, \mu_2, ..., \mu_k)$$
 = total squared distance of each data point \vec{x}_i to its closest centroid μ_i

- This C has a special name, inertia.
- Lower values of C lead to "better" clusterings.
 - ▶ **Goal:** Find the centroids $\mu_1, \mu_2, ..., \mu_k$ that minimize C.

Discussion Question

Suppose we have *n* data points, $\vec{x}_1, \vec{x}_2, ..., \vec{x}_n$, each of which are in \mathbb{R}^d .

Suppose we want to cluster our dataset into k clusters.

How many ways can we assign points to clusters?

- a) $d \cdot k$
- e) $n \cdot k \cdot d$

How do we minimize inertia?

- Problem: there are exponentially many possible clusterings. It would take too long to try them all.
- ► Another Problem: we can't use calculus or algebra to minimize *C*, since to calculate *C* we need to know which points are in which clusters.
- We need another solution.

k-Means Clustering

k-Means Clustering, i.e. Lloyd's Algorithm

Here's an algorithm that attempts to minimize inertia:

- 1. Pick a value of k and randomly initialize k centroids.
- 2. Keep the centroids fixed, and update the groups.
 - Assign each point to the nearest centroid.
- 3. Keep the groups fixed, and update the centroids.
 - Move each centroid to the center of its group.

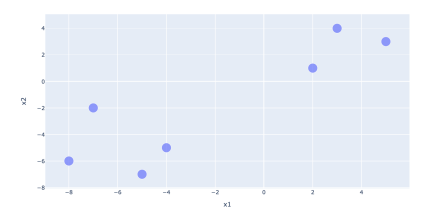
4. Repeat steps 2 and 3 until the centroids stop changing.

Example

See the following site for an interactive visualization of k-Means Clustering: https://tinyurl.com/40akmeans

An example by hand

Suppose we choose the initial centroids $\mu_1 = \begin{bmatrix} 2 \\ 1 \end{bmatrix}$ and $\mu_2 = \begin{bmatrix} 3 \\ 4 \end{bmatrix}$. Where will the centroids move to after one iteration of k-Means Clustering?



Demo

Let's see k-Means Clustering in action. Follow along here.

Why does k-Means work?

What is the goal of k-Means Clustering?

Recall, our goal is to find the centroids $\mu_1, \mu_2, ..., \mu_k$ that minimize inertia:

$$C(\mu_1, \mu_2, ..., \mu_k)$$
 = total squared distance of each data point \vec{x}_i to its closest centroid μ_i

- Let's argue that each step of the k-Means Clustering algorithm reduces inertia.
 - After enough iterations, inertia will be small enough.

Step 1: Pick a value of *k* and randomly initialize *k* centroids.

After initializing our *k* centroids, we have an initial value of inertia. We are going to argue that this only decreases.

Step 2: Keep the centroids fixed, and update the groups by assigning each point to the nearest centroid.

- Assuming the centroids are fixed, for each \vec{x}_i we have a choice which group should it be a part of?
- Mhichever group we choose, inertia will be calculated using the squared distance between \vec{x}_i and that group's centroid.
- Thus, to minimize inertia, we assign each \vec{x}_i to the group corresponding to the closest centroid.

Note that this analysis holds every time we're at Step 2, not just the first time.

Step 3: Keep the groups fixed, and update the centroids by moving each centroid to the center of its group (by averaging coordinates).

▶ Before we justify why this is optimal, let's revisit inertia.

Aside: separating inertia

► Inertia:

$$C(\mu_1, \mu_2, ..., \mu_k)$$
 = total squared distance of each data point \vec{x}_i to its closest centroid μ_i

Note that an equivalent way to write inertia is

$$C(\mu_1, \mu_2, ..., \mu_k) = C(\mu_1) + C(\mu_2) + ... + C(\mu_k)$$
 where $C(\mu_j) = \text{total squared distance of each}$ data point \vec{x}_i in group j to centroid μ_i

What's the point?

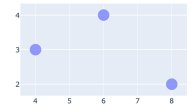
$$C(\mu_1, \mu_2, ..., \mu_k) = C(\mu_1) + C(\mu_2) + ... + C(\mu_k)$$
 where
$$C(\mu_j) = \text{total squared distance of each data point } \vec{x}_i$$
 in group j to centroid μ_j

Step 3: Keep the groups fixed, and update the centroids by moving each centroid to the center of its group (by averaging coordinates).

Let's argue why this minimizes $C(\mu_i)$, for each group j.

 $C(\mu_j)$ = total squared distance of each data point \vec{x}_i in group j to centroid μ_j

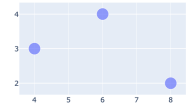
Suppose group j contains the points (4, 3), (6, 4), and (8, 2). Where should we put $\mu_j = \begin{bmatrix} a \\ b \end{bmatrix}$ to minimize $C(\mu_j)$?



 $C(\mu_i)$ = total squared distance of each data point \vec{x}_i in group j to centroid μ_i

Suppose group j contains the points (4, 3), (6, 4), and (8, 2).

Where should we put $\mu_j = \begin{bmatrix} a \\ b \end{bmatrix}$ to minimize $C(\mu_j)$?



Cost and empirical risk

On the previous slide, we saw a function of the form

$$C(\mu_j) = C(a, b) = (4 - a)^2 + (3 - b)^2$$
$$+ (6 - a)^2 + (4 - b)^2$$
$$+ (8 - a)^2 + (2 - b)^2$$

- ightharpoonup C(a, b) can be thought of as the sum of two separate functions, f(a) and g(b).
 - ► $f(a) = (4 a)^2 + (6 a)^2 + (8 a)^2$ computes the total squared distance of each x_1 coordinate to a.
 - From earlier in the course, we know that $a^* = \frac{4+6+8}{3} = 6$ minimizes f(a).

Practical considerations

Initialization

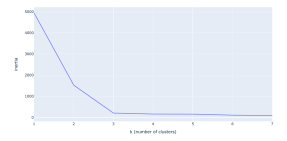
- Depending on our initial centroids, k-Means may "converge" to a clustering that doesn't actually have the lowest possible inertia.
 - In other words, like gradient descent, k-Means can get caught in a **local minimum**.
- Some solutions:
 - Run k-Means several times, each with different randomly chosen initial centroids. Keep track of the inertia of the final result in each attempt. Choose the attempt with the lowest inertia.
 - k-Means++: choose one initial centroid at random, and place other centroids far from all other centroids.

Choosing *k*

- Note that as *k* increases, inertia decreases.
 - Intuitively, as we add more centroids, the distance between each point and its closest centroid will drop.
- But the goal of clustering is to put data points into groups, and having a large number of groups may not be meaningful.
- This suggests a tradeoff between k and inertia.

The "elbow" method

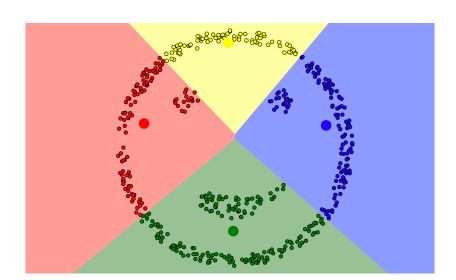
- Strategy: run k-Means Clustering for many choices of k (e.g. k = 1, 2, 3, ..., 8).
- Compute the value of inertia for each resulting set of centroids.
- ▶ Plot a graph of inertia vs k.
- Choose the value of k that appears at an "elbow".



See the notebook for a demo.

Low inertia isn't everything!

- Even if k-Means works as intended and finds the choice of centroids that minimize inertia, the resulting clustering may not look "right" to us humans.
 - Recall, inertia measures the total squared distance to centroids.
 - This metric doesn't always match our intuition.
- Let's look at some examples at https://tinyurl.com/40akmeans.
 - ► Go to "I'll Choose" and "Smiley Face". Good luck!



Other clustering techniques

- k-Means Clustering is just one way to cluster data.
- There are many others, each of which work differently and produce different kinds of results.
- Another common technique: agglomerative clustering.
 - ► High level: start out with each point being in its own cluster. Repeatedly combine clusters until only *k* are left.
- Check out this chart.

Summary

- k-Means Clustering attempts to minimize inertia.
 - We showed that it minimizes inertia at each step, but it's possible that it converges to a local minimum.
 - Different initial centroids can lead to different clusterings.
- To choose *k*, the number of clusters, we can use the elbow method.
- Next time: switching gears to probability and combinatorics.