

Machine Learning

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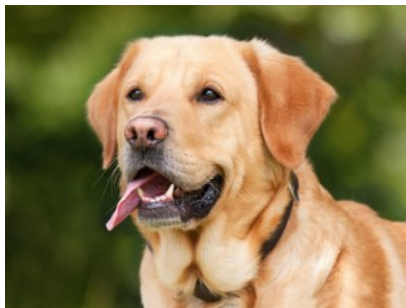
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Outline

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Why machine learning?

- **Question:** How do you write precise instructions (e.g. a computer program) to distinguish a dog from a cat?



(a) A dog.



(b) A cat.

Figure: A dog and a cat. Even a two-years old can tell which is which.

Recognizing cats and dogs

- Dogs have longer snouts.
- Cats have pointy ears.
- What is an ear anyway? Something that has roughly the shape of a triangle I guess.

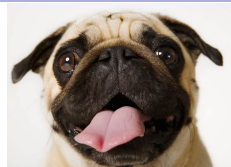


Figure: Short snout.



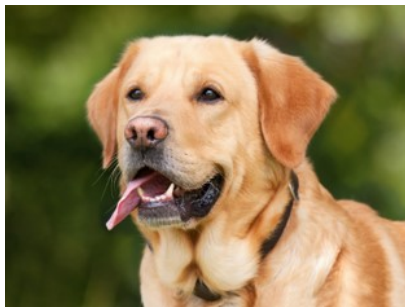
Figure: Pointy boy.



Figure: Nice ear.

Why machine learning?

- **Question:** How do you write precise instructions (e.g. a computer program) to distinguish a dog from a cat?



Mostly: you can't

Others have tried. They have failed.

What is machine learning?

- **Arthur Samuel (1959):** *"Machine Learning is the field of study that gives the computer the ability to learn without being explicitly programmed."*
- **Tom Mitchell (1998):** *"A computer program is said to learn from experience E with respect to some class of tasks T and performance measure P , if its performance at tasks in T , as measured by P , improves with experience E ."*

What is it used for?

- Machine learning is a must for:
 - ▶ Image recognition
 - ▶ Object detection
 - ▶ Speech recognition
 - ▶ ...
- And now also for:
 - ▶ Machine translation
 - ▶ Board games A.I. (go, chess...)
 - ▶ Robotics
 - ▶ ...
- And of course:
 - ▶ Conversational AI (ChatGPT, ...)
 - ▶ Image and video generation
 - ▶ ...



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Supervised versus unsupervised learning

Supervised learning

Annotations, usually called *labels* or *targets*, are provided with the data.

Unsupervised learning

No labels are provided with the data.

Supervised versus unsupervised learning

Supervised learning

Annotations, usually called *labels* or *targets*, are provided with the data.

Unsupervised learning

No labels are provided with the data.

Other learning paradigms

There also exist other paradigms such as reinforcement learning, semi-supervised learning, self supervised learning...

But in this course, we will mostly focus on supervised and unsupervised learning

Supervised versus unsupervised learning

For a same dataset, we can be either in a supervised or unsupervised learning setting depending on whether we use explicit labels or not.

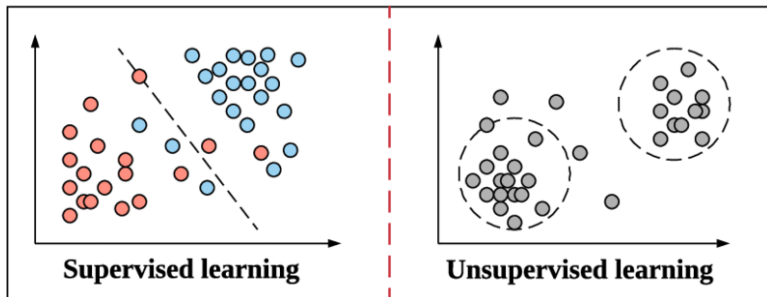


Figure: Left: Supervised learning with labeled points, from **red** and **blue** classes. The task is classification: tell whether a new point is red or blue. Right: Unsupervised learning with **unlabeled** points. The task is to group points into clusters.

Examples of **supervised** learning tasks

- Classification: predict a category
 - ▶ Image classification
 - ▶ Fraud detection in financial transactions
- Regression: predict a quantitative value
 - ▶ Stock price prediction
 - ▶ Sales forecast
- Other applications:
 - ▶ Translation from one language to another
 - ▶ Image captioning
 - ▶ Speech recognition

Targets

Can you identify what the labels are for each of these applications?

Examples of **unsupervised** learning tasks

- Clustering: identify groups of similar points in data
 - ▶ Customer segmentation in marketing
 - ▶ Document clustering in text analysis
 - ▶ Gene expression clustering in bioinformatics
- Dimensionality reduction: reduce the number of dimensions of the input data while keeping as much information as possible
 - ▶ Visualization of high dimensional data
 - ▶ Image compression
 - ▶ Representation learning: e.g. learn meaningful representations of users and movies for recommendation systems
- ...

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A first supervised learning algorithm

Let's introduce a first supervised learning method: **K-Nearest Neighbors** or **KNN**.

- KNN is a versatile and intuitive algorithm
- Used for both classification and regression tasks
- Instance-based learning: no explicit training phase
- Relies on the principle: "Similar inputs have similar outputs"

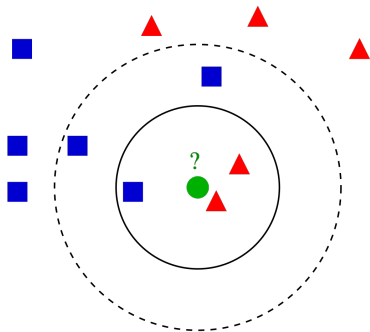


Figure: KNN illustration

How KNN Works

To classify a new, unlabeled data point (green dot in the center):

- Calculate distance to all training points
- Find the K nearest neighbors:
 - ▶ e.g. the 3 shapes inside the solid circle for $K = 3$
 - ▶ or the 5 shapes inside the dotted circle for $K = 5$
- Output the majority among the K neighbors
 - ▶ red triangle for $K = 3$ (2 out of 3)
 - ▶ blue square for $K = 5$ (3 out of 5)

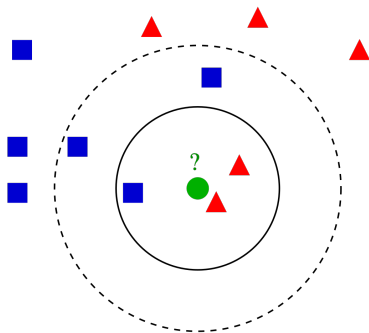


Figure: KNN illustration

Formalization

Assume we have a set of N points $\{\mathbf{x}_1, \dots, \mathbf{x}_N\}$ with corresponding labels $\{y_1, \dots, y_N\}$. The points \mathbf{x}_n have dimension D , such that

$$\forall n \quad \mathbf{x}_n = \begin{pmatrix} x_{n,1} \\ x_{n,2} \\ \vdots \\ x_{n,D} \end{pmatrix} = (x_{n,1}, \dots, x_{n,D})^\top \in \mathbb{R}^D$$

A note on notations

We will use the notations \mathbf{x}_n , y_n , D etc a lot throughout this course.

Assume we are also provided with a distance d , for example the Euclidean distance

$$d(\mathbf{x}_i, \mathbf{x}_j) = \|\mathbf{x}_i - \mathbf{x}_j\|_2$$

Given a new point \mathbf{x} , we want to predict its label y .

Formalization (continued)

To estimate the label \hat{y} of \mathbf{x} , we proceed as follows:

- 1 Calculate distances to all training points:

$$d_n = d(\mathbf{x}, \mathbf{x}_n) \quad \forall n \in \{1, \dots, N\}$$

- 2 Find indices of K nearest neighbors:

$$\mathcal{N}_K(\mathbf{x}) = \text{argsort}_K(d_1, \dots, d_N)$$

where argsort_K returns the indices of the K smallest values.

- 3 Predict the most common class among neighbors:

$$\hat{y} = \text{mode}\{y_i : i \in \mathcal{N}_K(\mathbf{x})\}$$

Variants of KNN

- We can also use KNN for regression: predict the average of neighbor values:

$$\hat{y} = \frac{1}{K} \sum_{i \in \mathcal{N}_K(\mathbf{x})} y_i$$

- In some variants, the neighbors can be weighted:

$$\hat{y} = \frac{\sum_{i \in \mathcal{N}_K(\mathbf{x})} w_i y_i}{\sum_{i \in \mathcal{N}_K(\mathbf{x})} w_i}$$

with a weigh function $w(\cdot)$ usually decreasing with the distance d (e.g., $w_n = \frac{1}{d_n}$ or $w_n = e^{-d_n}$)

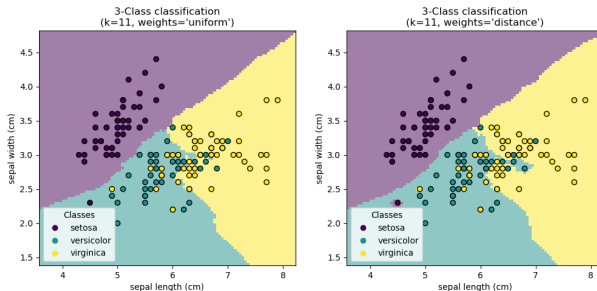
- Other variant: Fixed Radius NN. Instead of K neighbors, we use all points within a fixed radius R . Define the neighborhood as:

$$\mathcal{N}_R(\mathbf{x}) = \{i : d(\mathbf{x}, \mathbf{x}_i) \leq R\}$$

Then apply similar formulas as in standard or weighted KNN.

Another KNN example

Border contours from Scikit learn.



Limits of KNN

- Are we done with (supervised) machine learning? Have we found the perfect classification and regression algorithm?

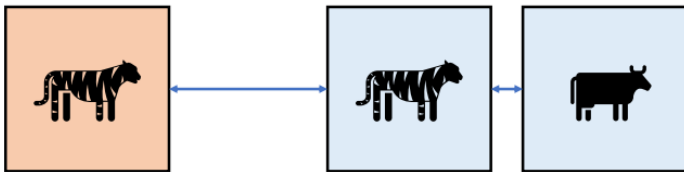


Figure: Measuring pixel distances. The center image is closer to the image on the right than the one on the left. Is it a cow?

- (Un)fortunately, it looks like **no, we are not**.

The curse of dimensionality

In high-dimensional spaces, the distances between points become less meaningful.

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Unsupervised learning

We will now shortly introduce a first unsupervised learning algorithm:
K-Means.

K-Means is a *clustering* algorithm.

Clustering

Clustering is the task of dividing a set of objects into groups, or **clusters**, in such a way that **objects in the same cluster are more similar** to each other than to those in other clusters.

A word of caution

- How can we group these vehicles?
 - ▶ Motor / no motor?
 - ▶ Touches the ground / does not touch the ground?
 - ▶ Single passenger / multiple passengers?



Clustering can be subjective

There can be a part of subjectivity in how we define groups or the idea of “close to each other”

Back to Euclidean space

- Let's focus on a simple, 2-dimensional, Euclidean example for now.

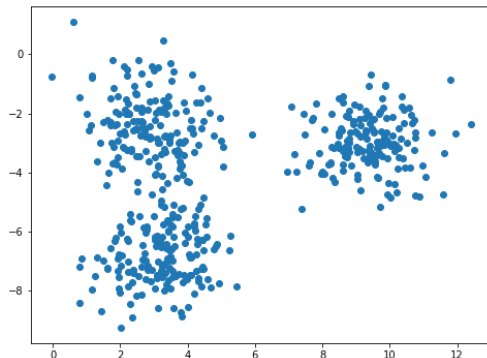


Figure: How many clusters are there?

- How could we define what would constitute “good” clusters here?

What makes “good” clusters?

- Intuitively, we want points in the same cluster to be:
 - ▶ Close to each other
 - ▶ Far from points in other clusters
- How can we do that? For example, we could:
 - ▶ Define a representative point for each cluster
 - ▶ Assign points to the closest representative
- This way we can measure “closeness” using distance to the representative
- A natural choice for the representative is the center of the cluster

Key ideas

- Use cluster centers as representatives
- Assign points to the nearest center
- Minimize the total distance between points and their assigned centers

Formalization

Given N unlabeled points $\{\mathbf{x}_1, \dots, \mathbf{x}_N\}$, $\mathbf{x}_n \in \mathbb{R}^D$, we want to:

- Find K representatives $\{\mathbf{c}_k\}_{k \in [1, K]} = \{\mathbf{c}_1, \dots, \mathbf{c}_K\}$, $\mathbf{c}_k \in \mathbb{R}^D$
- Find an assignment a that assigns each point \mathbf{x}_n to one cluster \mathbf{c}_k :
 - ▶ $a : \mathbb{R}^D \rightarrow [1, K]$
 - ▶ define $a_n := a(\mathbf{x}_n)$ the assignment of point $\mathbf{x}_n \forall n$

Objective:

$$\begin{array}{ll} \text{minimize} & \sum_{n=1}^N \|\mathbf{x}_n - \mathbf{c}_{a_n}\|_2^2 \\ \text{w.r.t. } & \{a_n\}_n, \{\mathbf{c}_k\}_k \end{array}$$

- That is, we want to find centroids and assignments such that each point \mathbf{x}_n is as close as possible to its centroid \mathbf{c}_{a_n}

Formalization

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- That is, we want to find centroids and assignments such that each point \mathbf{x}_n is as close as possible to its centroid \mathbf{c}_{a_n}

One tiny issue

This problem is NP-hard

K-means: a greedy heuristic approach

Objective (reminder):

$$\underset{\text{w.r.t. } \{a_n\}_n, \{\mathbf{c}_k\}_k}{\text{minimize}} \sum_{n=1}^N \|\mathbf{x}_n - \mathbf{c}_{a_n}\|_2^2$$

The K-means algorithm is a greedy heuristic for this objective, that alternates between two steps:

- ① Optimize assignments $\{a_n\}_n$ given fixed representatives, also called *centroids* $\{\mathbf{c}_k\}_k$
- ② Optimizing centroid locations $\{\mathbf{c}_k\}_k$ given fixed assignments $\{a_n\}_n$

This approach is computationally efficient, and reasonably intuitive.

K-means algorithm: formalization

- Initialize centroids $\{\mathbf{c}_1, \dots, \mathbf{c}_K\}$ randomly
- Assign each point \mathbf{x}_n to the nearest centroid \mathbf{c}_k

$$a(\mathbf{x}_n) = \operatorname{argmin}_{k \in [1, K]} \|\mathbf{x}_n - \mathbf{c}_k\|_2^2$$

- Update clusters as the mean of the assigned points:

$$\mathbf{c}_k = \frac{1}{|\mathcal{C}_k|} \sum_{\mathbf{x} \in \mathcal{C}_k} \mathbf{x}$$

(Mean of points in cluster \mathbf{c}_k)

$$\mathcal{C}_k = \{\mathbf{x} | a(\mathbf{x}) = k\}$$

(Set of points \mathbf{x} assigned to cluster \mathbf{c}_k)

Repeat

Repeat the last 2 steps until the assignments a_n stop changing.

K-Means: illustration

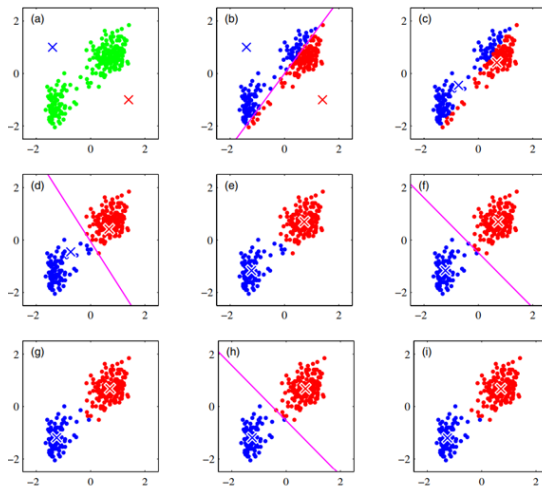


Figure: (a) Random initialization of the 2 centroids. (b) Assignment step: points are assigned to the nearest centroid. (c) Update step: centroid are updated to the mean of their assigned points. (d-h) Repeat. (i) Algorithm has converged. From Bishop.

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Questions about K-Means

Will this algorithm always converge?



Does it always produce the same solution?



Does it find the optimal solution?



Questions about K-Means

Will this algorithm always converge?

- This is the hardest of the 3 questions. Answer in a few slides.

Does it always produce the same solution?

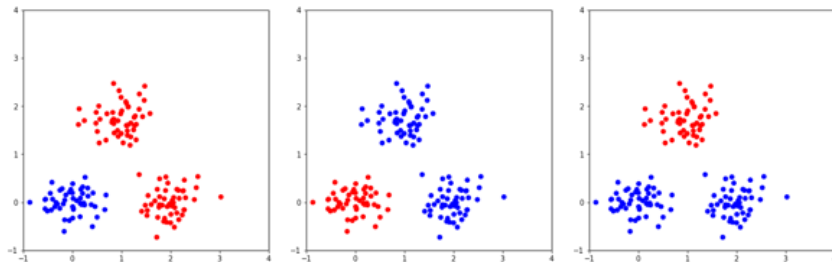
- **Hint:** Think about the initialization step.

Does it find the optimal solution?

- **Hint:** Consider the answer to the previous question.

Does K-Means always produce the same, optimal solution?

- No, different initializations can lead to different results



K-Means in practice

It is good practice to run the algorithm several times

Question time!

- Which clusters will we find if we apply K-Means with $K = 2$?

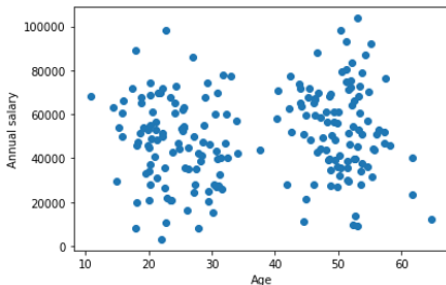
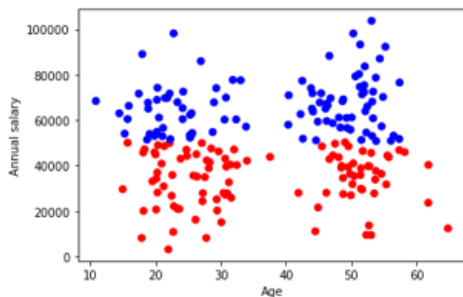


Figure: Some points forming 2 clusters.

(Trick) question time! Answer time



- Why? Because the scale of each variable (age and salary) is very different.

K-Means in practice

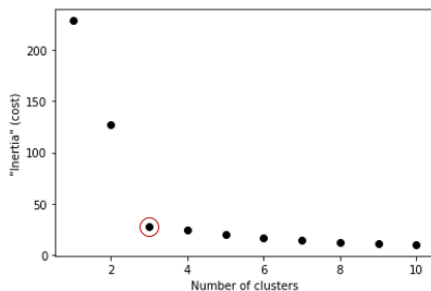
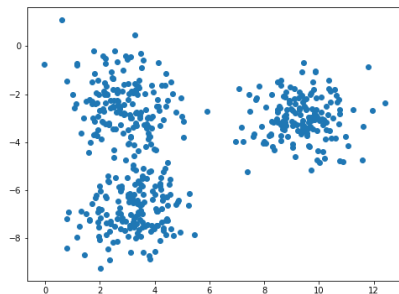
Standardization: it is often useful to “standardize” variables:

$$x_{\text{stand}} = \frac{x - \mu}{\sigma}$$

where μ is the mean and σ the standard deviation of all x .

How to choose K: elbow method

- K is easy to estimate in 2 dimensions. Not so much when $D = 200$.
- The (*theoretically* optimal) cost $\sum_{n=1}^N \|\mathbf{x}_n - \mathbf{c}_{a(\mathbf{x}_n)}\|_2^2$, sometimes also called the *inertia*, decreases with the number of clusters K .



- We can select a value corresponding to a sharp decrease, $K = 3$ in this example (“elbow method”)

How to choose K, alternative method: silhouette score

- For point \mathbf{x}_n , we define u_n the average distance to points of the corresponding cluster \mathcal{C}_{a_n} :

$$u_n = \frac{1}{|\mathcal{C}_{a_n}| - 1} \sum_{\substack{\mathbf{x}_j \in \mathcal{C}_{a_n} \\ \mathbf{x}_n \neq \mathbf{x}_j}} d(\mathbf{x}_n, \mathbf{x}_j)$$

- We define v_n as $v_n = \min_{k \neq a_n} \frac{1}{|\mathcal{C}_k|} \sum_{\mathbf{x}_j \in \mathcal{C}_k} d(\mathbf{x}_n, \mathbf{x}_j)$

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the average distance to points in the closest different cluster \mathcal{C}_k , $k \neq a_n$

- The *silhouette score* s_n of point \mathbf{x}_n is

$$s_n = \frac{v_n - u_n}{\max\{u_n, v_n\}} \quad (\text{and } s_n = 0 \text{ if } |\mathcal{C}_{a_n}| = 1). \quad -1 \leq s_n \leq 1 \forall n$$

- An average silhouette score $S = \frac{1}{N} \sum s_n$ close to 1 corresponds to “good” clusters

How to choose K, alternative method: silhouette score

- For point \mathbf{x}_n , we define u_n the average distance to points of the corresponding cluster \mathcal{C}_{a_n} :

Warning

Don't worry if you don't understand everything. I am actually not a big fan of the silhouette score.

- We define v_n as $v_n = \min_{k \neq a_n} \frac{1}{|\mathcal{C}_k|} \sum_{\mathbf{x}_j \in \mathcal{C}_k} d(\mathbf{x}_n, \mathbf{x}_j)$
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K-Means in practice

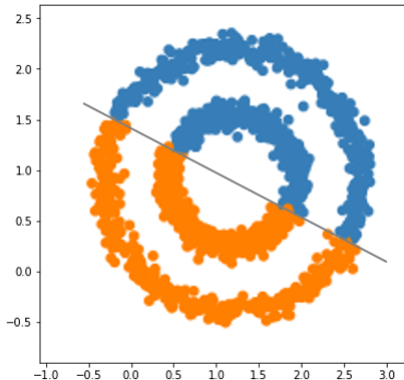
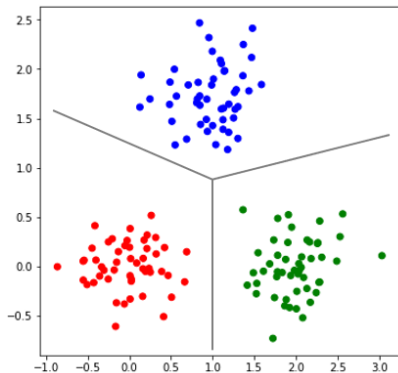
Why? Because it makes it too easy to simply choose the K that gives the "best" silhouette score $S(K)$. This can make us forget that the choice of K is always somehow arbitrary. In the elbow at least, this part is clear.

$$s_n = \frac{u_n}{\max\{u_n, v_n\}} \quad (\text{and } s_n = 0 \text{ if } |\mathcal{C}_{a_n}| = 1). \quad 1 \geq s_n \geq 0$$

- An average silhouette score $S = \frac{1}{N} \sum s_n$ close to 1 corresponds to "good" clusters

Limits of K-means

- K-means can be considered “linear” in that cluster boundaries are linear



(From the scikit-learn documentation)

Setting up a Python Environment with Conda

- 1 Create a new Conda environment:

```
conda create -n mlclass python=3.10
```

- 2 Activate the environment:

```
conda activate mlclass
```

- 3 Install scikit-learn using pip:

```
pip install scikit-learn
```

Alternatively, you may use Google Collab:

<https://colab.research.google.com/>