Lecture 02

Nearest Neighbor Methods

STAT 451: Intro to Machine Learning, Fall 2020 Sebastian Raschka

http://stat.wisc.edu/~sraschka/teaching/stat451-fs2020/

Lecture 2 (Nearest Neighbors) Topics

- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python



Sebastian Raschka

STAT 451: Intro to ML



Saudi Computer Society, King Saud University

Applied Computing and Informatics

(http://computer.org.sa) www.ksu.edu.sa www.sciencedirect.com



ORIGINAL ARTICLE

Automated web usage data mining and recommendation system using K-Nearest Neighbor (KNN) classification method



D.A. Adeniyi, Z. Wei, Y. Yongquan *

The major problem of many on-line web sites is the presentation of many choices to the client at a time; this usually results to strenuous and time consuming task in finding the right product or information on the site. In this work, we present a study of automatic web usage data mining and recommendation system based on current user behavior through his/her click stream data on the newly developed Really Simple Syndication (RSS) reader website, in order to provide relevant information to the individual without explicitly asking for it. **The K-Nearest-Neighbor (KNN) classification** method has been trained to be used on-line and in Real-Time to identify clients/visitors click stream data, matching it to a particular user group and recommend a tailored browsing option that meet the need of the specific user at a particular time. [...]

Distance Metric Learning for Large Margin Nearest Neighbor Classification

Weinberger, Kilian Q., John Blitzer, and Lawrence K. Saul. "Distance metric learning for large margin nearest neighbor classification." *Advances in Neural Information Processing Systems*. 2006.

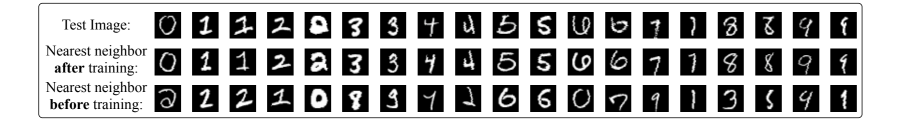
Kilian Q. Weinberger, John Blitzer and Lawrence K. Saul

Department of Computer and Information Science, University of Pennsylvania Levine Hall, 3330 Walnut Street, Philadelphia, PA 19104 {kilianw, blitzer, lsaul}@cis.upenn.edu

Among 3 nearest neighbors after but not before training:

Among 3 nearest neighbors before but not after training:

We show how to learn a Mahanalobis distance metric for knearest neighbor (kNN) classification by semidefinite programming. The metric is trained with the goal that the knearest neighbors always belong to the same class while examples from different classes are separated by a large margin. On seven data sets of varying size and difficulty, we find that metrics trained in this way lead to significant improvements in kNN classification—for example, achieving a test error rate of 1.3% on the MNIST handwritten digits. As in support vector machines (SVMs), the learning problem reduces to a convex optimization based on the hinge loss. Unlike learning in SVMs, however, our framework requires no modification or extension for problems in multiway (as opposed to bi- nary) classification.

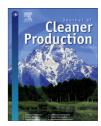




Contents lists available at ScienceDirect

Journal of Cleaner Production

journal homepage: www.elsevier.com/locate/jclepro



Remaining useful life estimation of lithium-ion cells based on *k*-nearest neighbor regression with differential evolution optimization



Yapeng Zhou ^a, Miaohua Huang ^{a, *}, Michael Pecht ^b

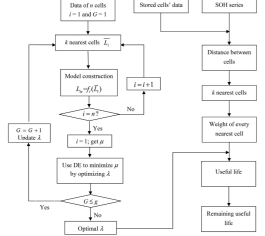


Fig. 1. Flowchart of parameter optimization and RUL estimation.

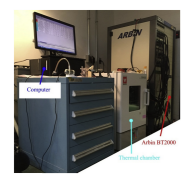


Fig. 2. Cell test bench

different jellyroll configurations. All the cells have a LiCoO₂ cathode and graphite anode, and the electrolyte material contains LiPF₆ EC, and DEC, and the rated voltage is 3.7 V. The cathode and anode layers of groups A and B are wrapped around orthogonal rotation center. Cells were charged with constant current and constant voltage protocol and discharged with constant current to 2.7 V under 24 °C. The detailed specifications and charge/discharge method of these cells are shown in Table 1. As shown in Table 1, cells of group B were discharged with constant current 0.5 C, and a rate of x C is a current equal of multiplying x and the rated capacity. Groups A and B are used to validate the feasibility and online applicability of this method, respectively.

The detailed experiment procedure is as follows:

- Program the charge/discharge with Bits Pro software on computer.
- Connect the cells to the circuit, and put them into the thermal chamber
- Turn on the thermal chamber and set the temperature at 24 °C, and rest 1 h.
- 4. Start the charge/discharge cycling with the Bits Pro softward 5. Terminate the cycling when the SOH reaches 80%.

Note that there is an interval of 5 min between each charge and discharge. The capacity was calculated by integrating the discharge

Remaining useful life estimation is of great importance to customers who use battery-powered products. This paper develops a remaining useful life estimation model based on **k-nearest** neighbor regression by incorporating data from all the cells in a battery pack. A differential evolution technique is employed to optimize the parameters in the estimation model. In this approach, remaining useful life is estimated from a weighted average of the useful life of several nearest cells that share a similar degradation trend to the cell whose remaining useful life needs to be estimated. The developed method obtains a remaining useful life estimation result with average error of 9 cycles, and the best estimation only has an error of 2 cycles. [...]

^a Hubei Key Laboratory of Advanced Technology for Automotive Components, Wuhan University of Technology, Wuhan, 430070, PR China

^b Center for Advanced Life Cycle Engineering, University of Maryland, College Park, MD, 20742, USA

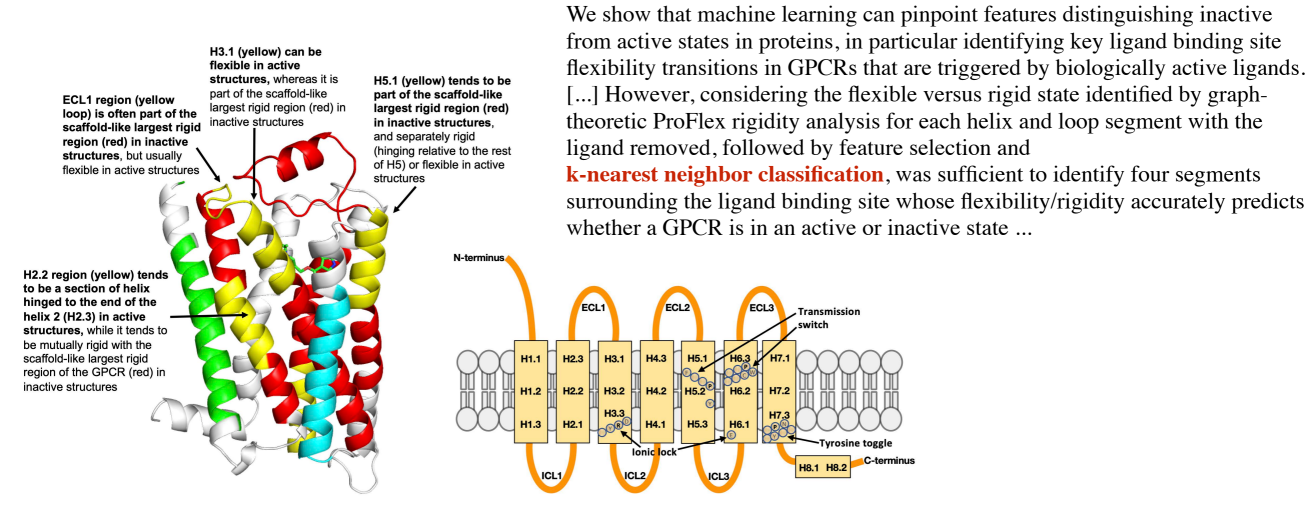




Article

Machine Learning to Identify Flexibility Signatures of Class A GPCR Inhibition

Joseph Bemister-Buffington ¹, Alex J. Wolf ¹, Sebastian Raschka ^{1,2,*} and Leslie A. Kuhn ^{1,3,*}

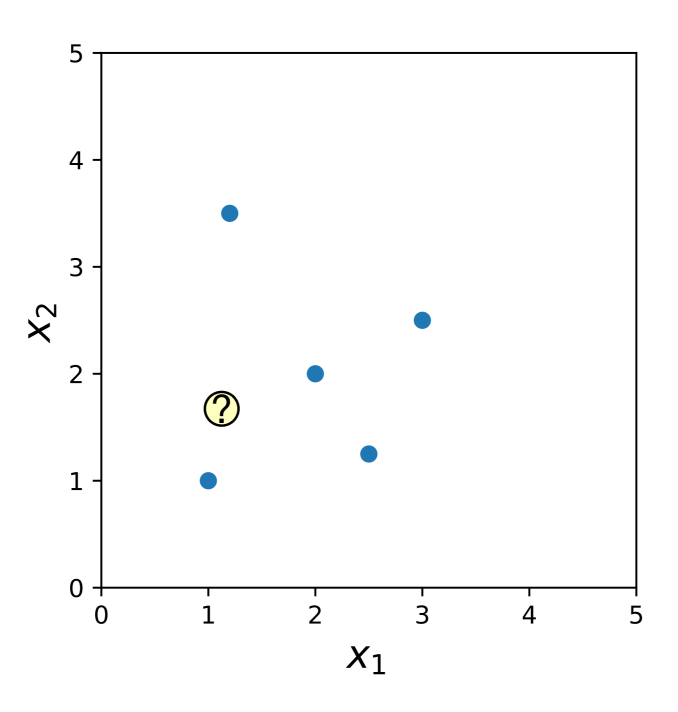


Joe Bemister-Buffington, Alex J. Wolf, Sebastian Raschka, and Leslie A. Kuhn (2020) *Machine Learning to Identify Flexibility Signatures of Class A GPCR Inhibition* Biomolecules 2020, 10, 454.

1-Nearest Neighbor

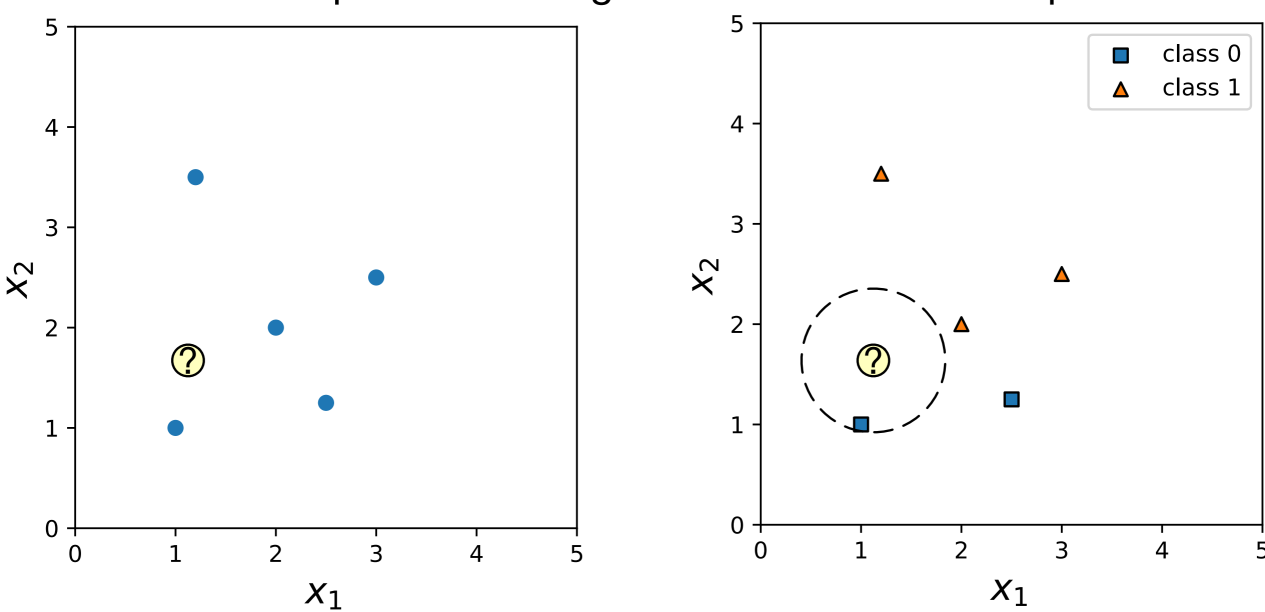
1-Nearest Neighbor

Task: predict the target / label of a new data point



1-Nearest Neighbor

Task: predict the target / label of a new data point



How? Look at most "similar" data point in training set

1-Nearest Neighbor Training Step

$$\langle \mathbf{x}^{[i]}, y^{[i]} \rangle \in \mathcal{D} \quad (|\mathcal{D}| = n)$$

How do we "train" the 1-NN model?

1-Nearest Neighbor Training Step

$$\langle \mathbf{x}^{[i]}, y^{[i]} \rangle \in \mathcal{D} \quad (|\mathcal{D}| = n)$$

To train the 1-NN model, we simply "remember" the training dataset

1-Nearest Neighbor Prediction Step

Given:
$$\langle \mathbf{x}^{[i]}, y^{[i]} \rangle \in \mathcal{D} \quad (|\mathcal{D}| = n)$$

 $\langle \mathbf{x}^{[q]}, ??? \rangle$

Predict: $f(\mathbf{x}^{[q]})$

Algorithm:

closest_point := None

 $closest_distance := \infty$

query point

- for i = 1, ..., n:
 - \circ current_distance := $d(\mathbf{x}^{[i]}, \mathbf{x}^{[q]})$
 - if current_distance < closest_distance:
 - closest_distance := current_distance
 - closest_point := $\mathbf{x}^{[i]}$
- return *f*(closest_point)

Commonly used: Euclidean Distance (L2)

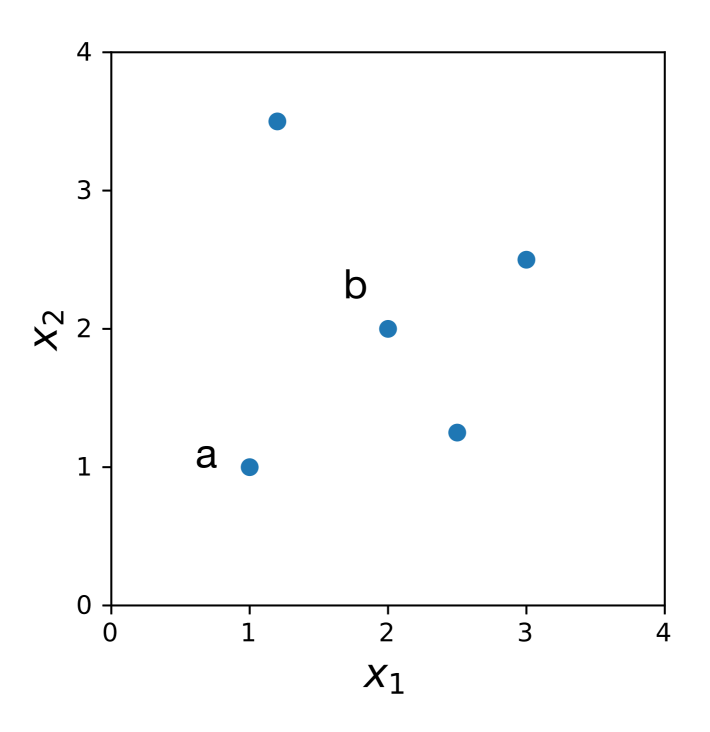
$$d(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \sqrt{\sum_{j=1}^{m} \left(x_j^{[a]} - x_j^{[b]}\right)^2}$$

Lecture 2 (Nearest Neighbors) Topics

- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python

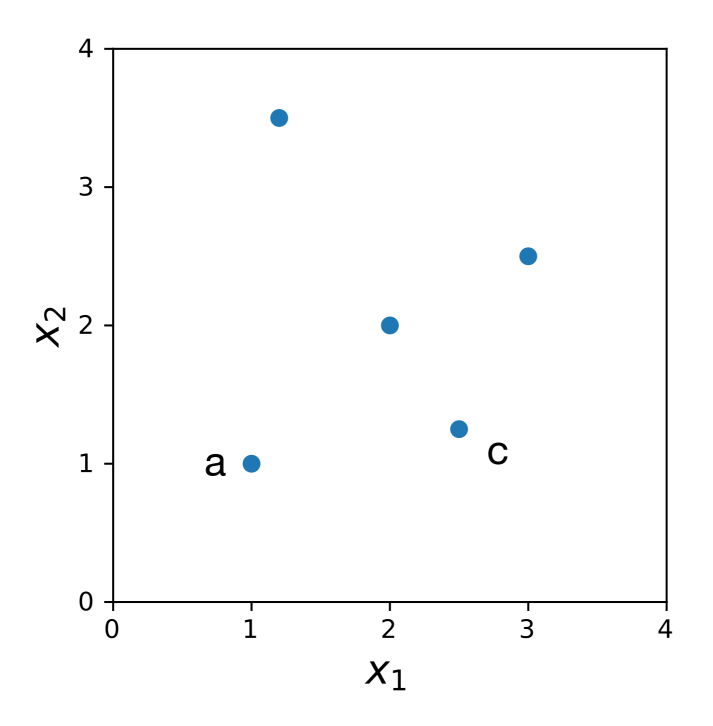
Nearest Neighbor Decision Boundary

Decision Boundary Between (a) and (b)



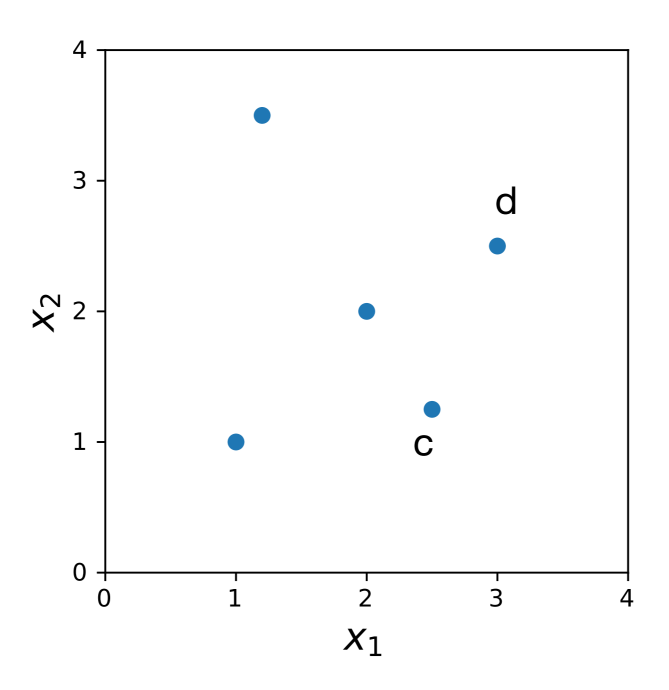
How does it look like?

Decision Boundary Between (a) and (c)

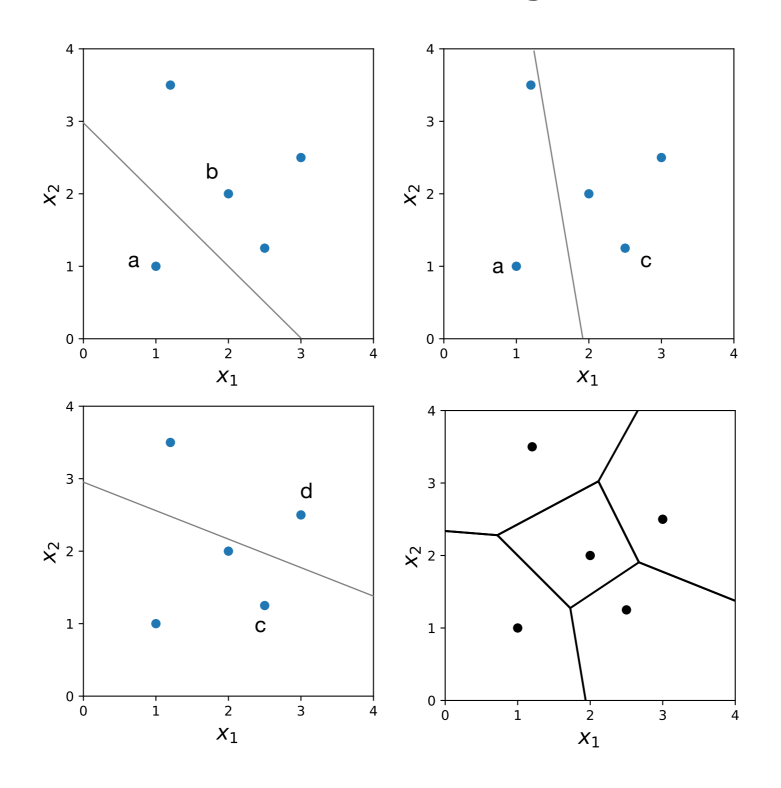


How does it look like?

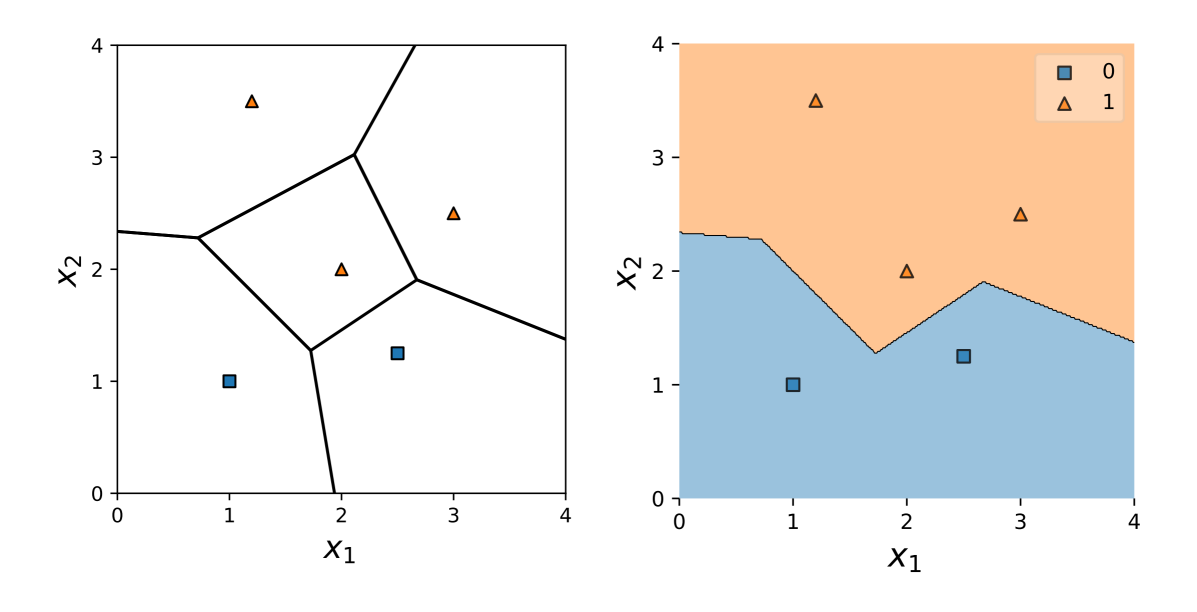
Decision Boundary Between (a) and (c)



Decision Boundary of 1-NN

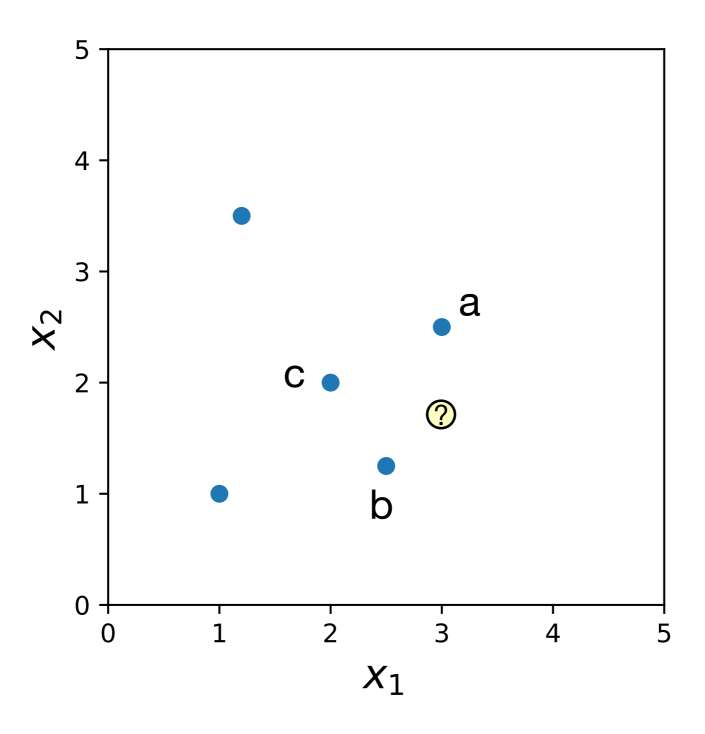


Decision Boundary of 1-NN

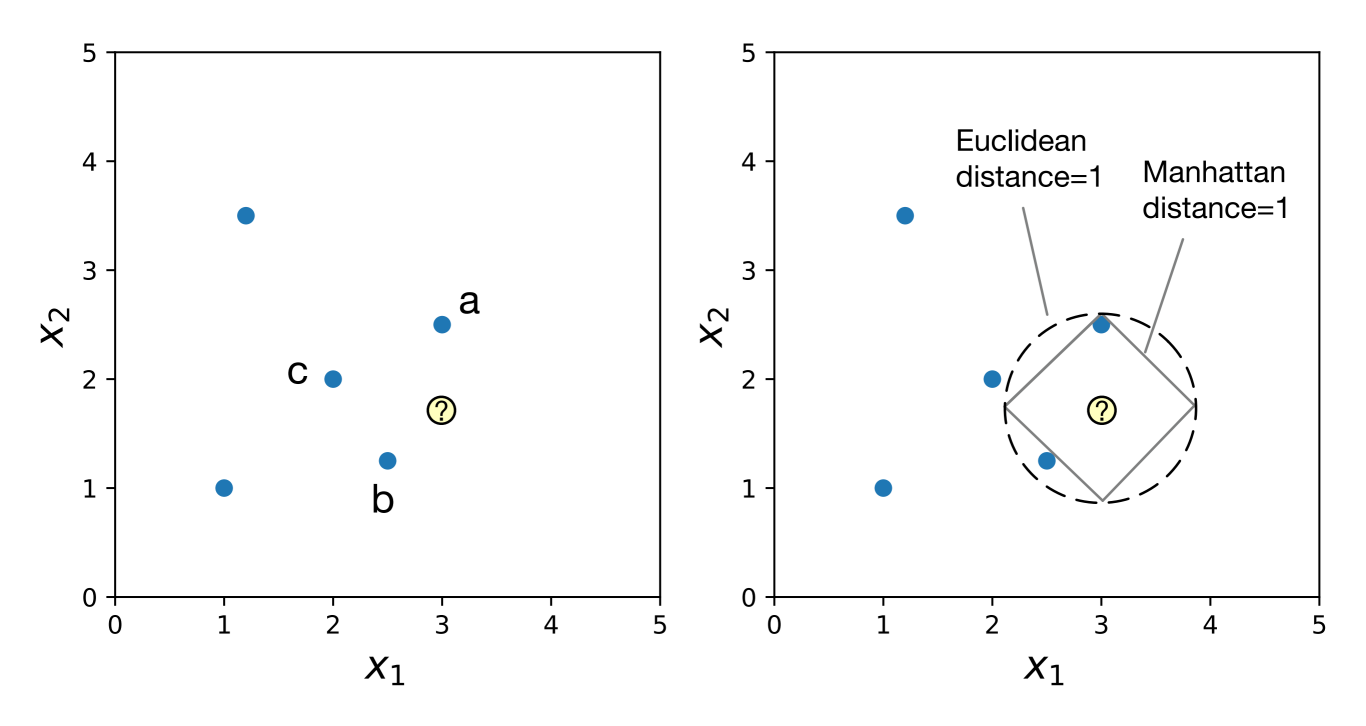


Which Point is Closest to ?





Depends on the Distance Measure!



Some Common Continuous Distance Measures

Euclidean

Manhattan

Minkowski:
$$d(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \left[\sum_{j=1}^{m} \left(\left|x_j^{[a]} - x_j^{[b]}\right|\right)^p\right]^{\overline{p}}$$

Mahalanobis

Cosine similarity

- - -

Some Discrete Distance Measures

Hamming distance:
$$d(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \sum_{j=1}^{m} \left| x_j^{[a]} - x_j^{[b]} \right|$$
 where $x_j \in \{0, 1\}$

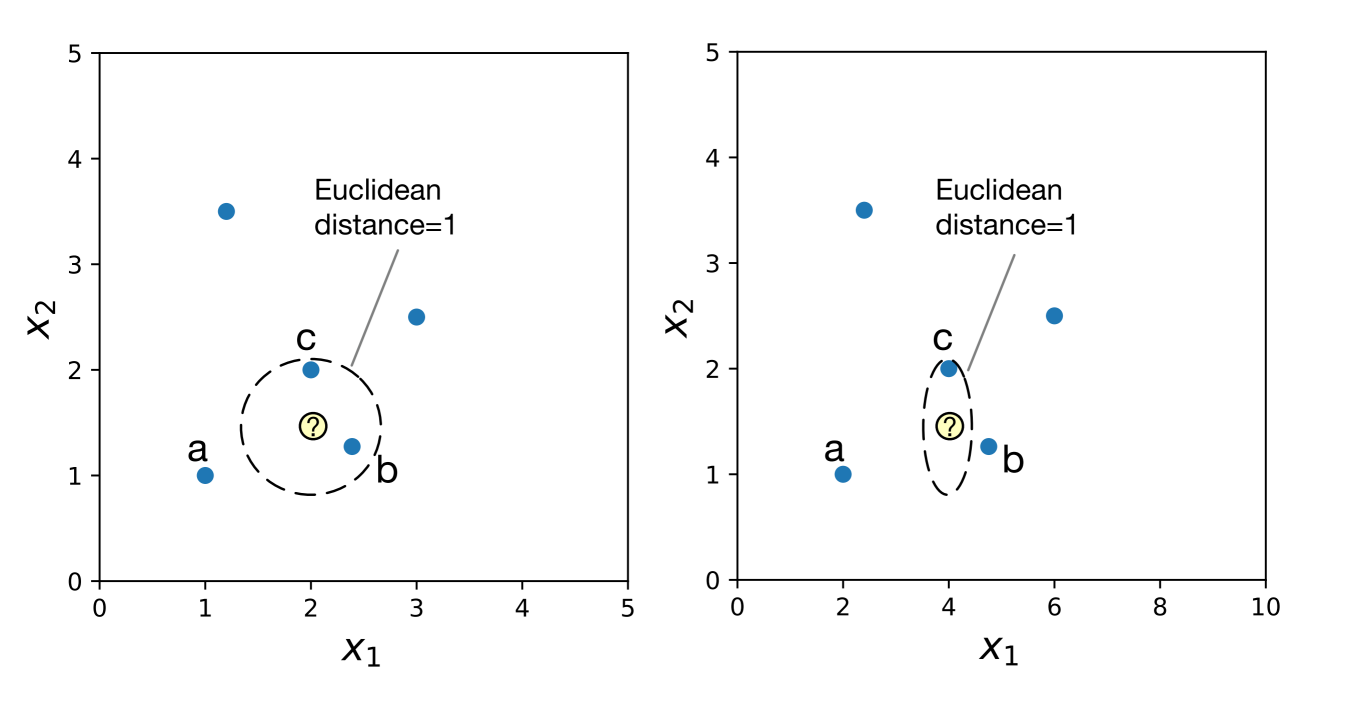
Jaccard/Tanimoto similarity:

$$J(A,B) = \frac{|A \cap B|}{|A \cup B|} = \frac{|A \cap B|}{|A| + |B| - |A \cap B|}$$

Dice:
$$D(A, B) = \frac{2|A \cap B|}{|A| + |B|}$$

- - -

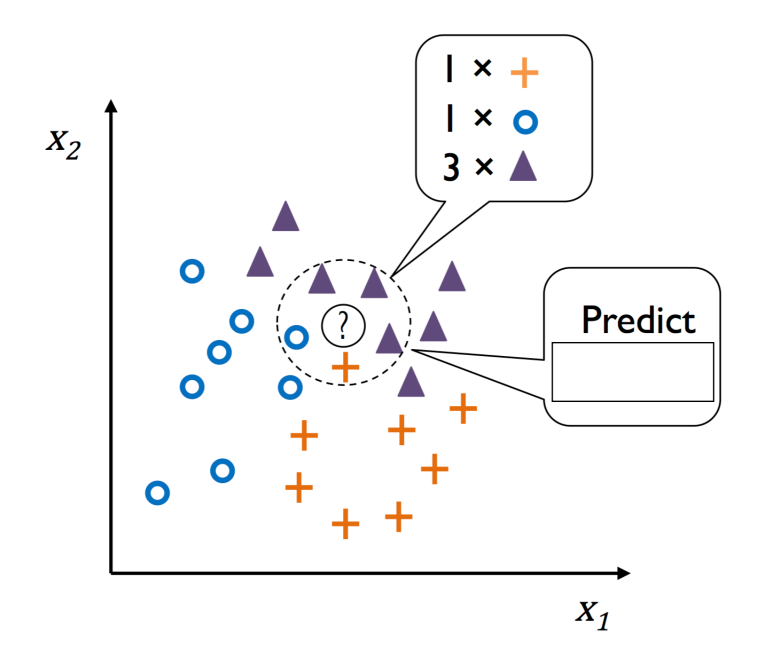
Feature Scaling

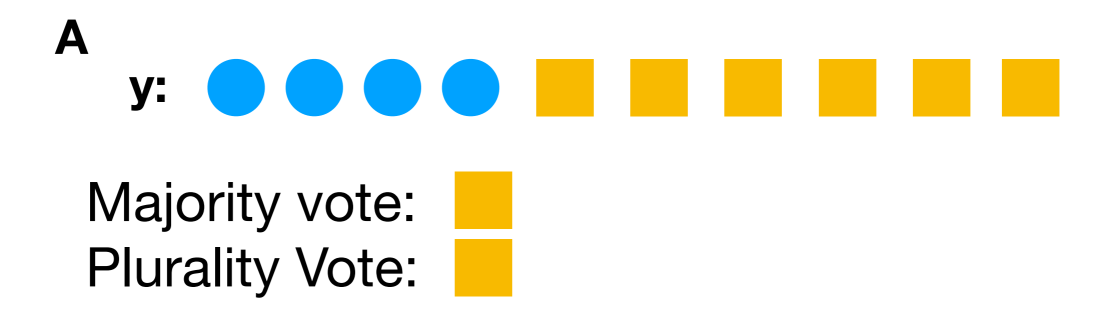


Lecture 2 (Nearest Neighbors) Topics

- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python

k-Nearest Neighbors







Majority vote: None

Plurality Vote:

kNN for Classification

$$\mathcal{D}_k = \{ \langle \mathbf{x}^{[1]}, f(\mathbf{x}^{[1]}) \rangle, \dots, \langle \mathbf{x}^{[k]}, f(\mathbf{x}^{[k]}) \rangle \} \qquad \mathcal{D}_k \subseteq \mathcal{D}$$

$$h(\mathbf{x}^{[q]}) = arg \max_{y \in \{1,...,t\}} \sum_{i=1}^{K} \delta(y, f(\mathbf{x}^{[i]}))$$
$$\delta(a, b) = \begin{cases} 1, & \text{if } a = b, \\ 0, & \text{if } a \neq b. \end{cases}$$

$$h(\mathbf{x}^{[t]}) = \mathsf{mode}(\{f(\mathbf{x}^{[1]}), ..., f(\mathbf{x}^{[k]})\})$$

kNN for Regression

$$\mathcal{D}_k = \{ \langle \mathbf{x}^{[1]}, f(\mathbf{x}^{[1]}) \rangle, \dots, \langle \mathbf{x}^{[k]}, f(\mathbf{x}^{[k]}) \rangle \} \qquad \mathcal{D}_k \subseteq \mathcal{D}$$

$$h(\mathbf{x}^{[t]}) = \frac{1}{k} \sum_{i=1}^{k} f(\mathbf{x}^{[i]})$$

Lecture 2 (Nearest Neighbors) Topics

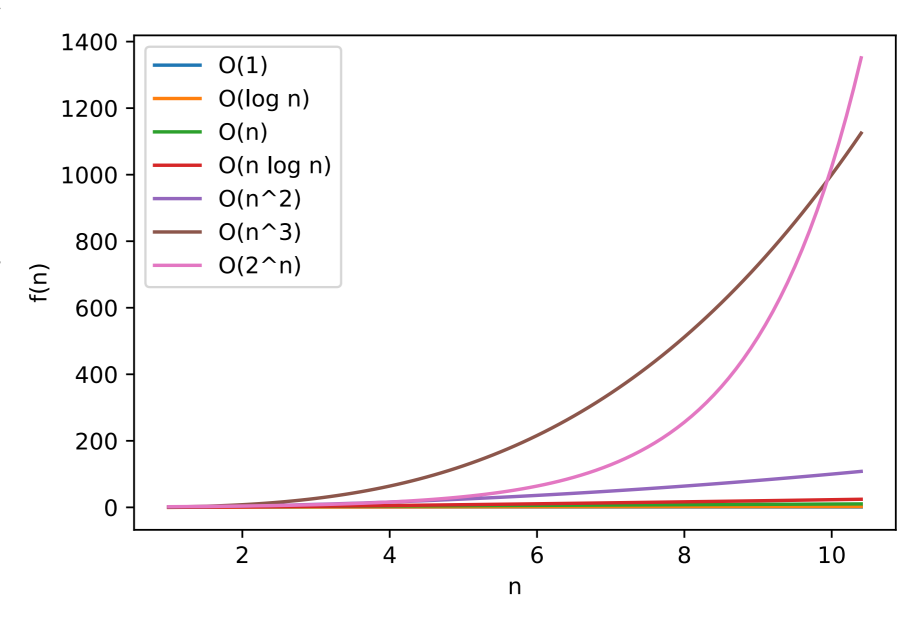
- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python

Big-O

f(n)	Name
1	Constant
$\log n$	Logarithmic
n	Linear
$n \log n$	Log Linear
n^2	Quadratic
n^3	Cubic
n^c	Higher-level polynomial
2^n	Exponential

Big-O

f(n)	Name
1	Constant
$\log n$	Logarithmic
n	Linear
$n \log n$	Log Linear
n^2	Quadratic
n^3	Cubic
n^c	Higher-level polynomial
$\frac{2^n}{n}$	Exponential



Big-O Example 1

$$f(x) = 14x^2 - 10x + 25$$

$$\mathcal{O}($$

Big-O Example 2

$$f(x) = (2x + 8)\log_2(x + 9)$$

 $\mathcal{O}($

Big-O Example 2

$$f(x) = (2x + 8)\log_2(x + 9)$$

Why don't we have to distinguish between different logarithms?

Big-O Example 3

```
A = [[1, 2, 3],
    [2, 3, 4]]
B = [[5, 8],
   [6, 9],
     [7, 10]]
def matrixmultiply (A, B):
    C = [[0 \text{ for row in range(len(A))}]
          for col in range(len(B[0]))]
    for row a in range(len(A)):
        for col b in range(len(B[0])):
            for col a in range(len(A[0])):
                C[row a][col b] += \
                     A[row a][col a] * B[col a][col b]
    return C
matrixmultiply(A, B)
```

```
Out[16]:
[[38, 56], [56, 83]]
```

Big O of kNN

40

Naive Nearest Neighbor Search

Variant A

$$\mathcal{D}_k := \{\}$$
 while $|\mathcal{D}_k| < k$:



- ullet closest_distance := ∞
- for i = 1, ..., n, $\forall i \notin \mathcal{D}_k$:
 - current_distance := $d(\mathbf{x}^{[i]}, \mathbf{x}^{[q]})$
 - if current_distance < closest_distance:</pre>
 - * closest_distance := current_distance
 - * closest_point $:= \mathbf{x}^{[i]}$
- ullet add closest_point to \mathcal{D}_k

Naive Nearest Neighbor Search

Variant B

$$\mathcal{D}_k := \mathcal{D}$$
 while $|\mathcal{D}_k| > k$:

- $largest_distance := 0$
- for $i = 1, ..., n \quad \forall i \in \mathcal{D}_k$:
 - current_distance $:=d(\mathbf{x}^{[i]},\mathbf{x}^{[q]})$
 - if current_distance > largest_distance:
 - * largest_distance := current_distance
 - * farthest_point $:=\mathbf{x}^{[i]}$
 - ullet remove farthest_point from \mathcal{D}_k

 $\mathcal{O}($

Naive Nearest Neighbor Search

Using a priority queue

O(_____)

Lecture 2 (Nearest Neighbors) Topics

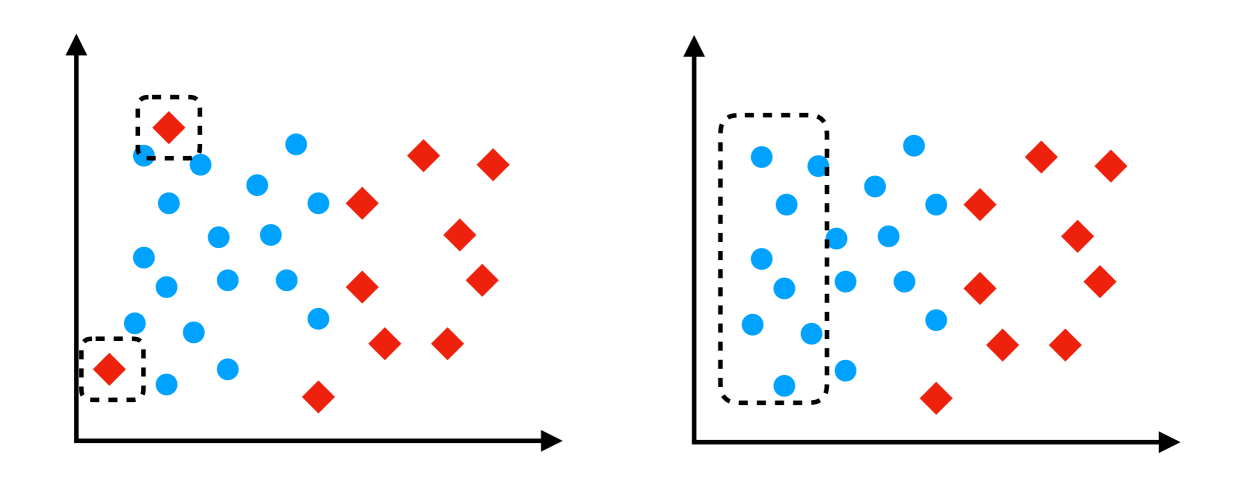
- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python

Data Structures

Dimensionality Reduction

Editing / "Pruning"

Editing / "Pruning"



Prototypes

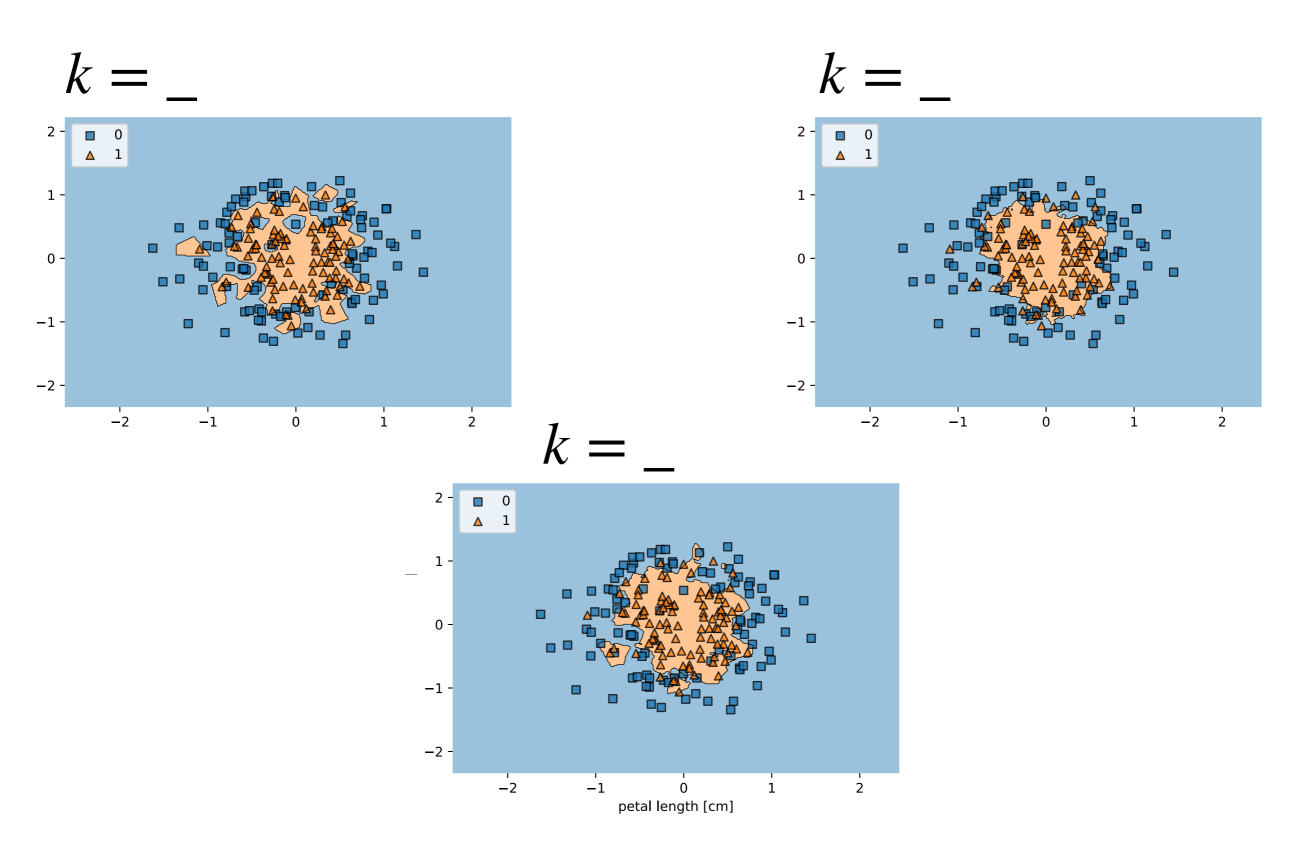
Improving Predictive Performance

Hyperparameters

Hyperparameters

- Value of k
- Scaling of the feature axes
- Distance measure
- Weighting of the distance measure

$k \in \{1,3,7\}$



Feature-Weighting via Euclidean Distance

$$d_{w}(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \sqrt{\sum_{j=1}^{m} w_{j} \left(x_{j}^{[a]} - x_{j}^{[b]}\right)^{2}}$$

As a dot product:

$$\mathbf{c} = \mathbf{x}^{[a]} - \mathbf{x}^{[b]}, \quad (\mathbf{c}, \mathbf{x}^{[a]}, \mathbf{x}^{[b]} \in \mathbb{R}^m)$$
$$d(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \sqrt{\mathbf{c}^{\mathsf{T}} \mathbf{c}}$$
$$d_w(\mathbf{x}^{[a]}, \mathbf{x}^{[b]}) = \mathbf{c}^{\mathsf{T}} \mathbf{W} \mathbf{c},$$

$$\mathbf{W} \in \mathbb{R}^{m \times m} = \mathbf{diag}(w_1, w_2, \dots, w_m)$$

Distance-weighted kNN

$$h(\mathbf{x}^{[t]}) = arg \max_{j \in \{1, ..., p\}} \sum_{i=1}^{k} w^{[i]} \delta(j, f(\mathbf{x}^{[i]}))$$

$$w^{[i]} = \frac{1}{d(\mathbf{x}^{[i]}, \mathbf{x}^{[t]})^2}$$

Small constant to avoid zero division or set $h(\mathbf{x}) = f(\mathbf{x})$

Lecture 2 (Nearest Neighbors) Topics

- 1. Intro to nearest neighbor models
- 2. Nearest neighbor decision boundary
- 3. K-nearest neighbors
- 4. Big-O & k-nearest neighbors runtime complexity
- 5. Improving & k-nearest neighbors: modifications and hyperparameters
- 6. K-nearest neighbors in Python

kNN in Python

DEMO