DSC 190 Machine Learning: Representations

Lecture 5 | Part 1

Choosing RBF Locations

Recap

We map data to a new representation by first choosing **basis functions**.

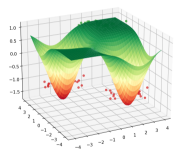
- Radial Basis Functions (RBFs), such as Gaussians, are a popular choice.
- Requires choosing center for each basis function.

Today's Lecture

How do we choose basis function centers automatically?

Prediction Function

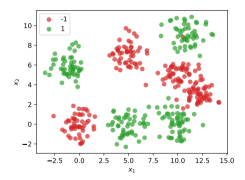
Our prediction function H is a surface that is made up of Gaussian "bumps".



$$H(\vec{x}) = w_0 + w_1 e^{-\|\vec{x} - \vec{\mu}_1\|^2 / \sigma^2} + w_2 e^{-\|\vec{x} - \vec{\mu}_2\|^2 / \sigma^2}$$

Choosing Centers

- Place the centers where the value of the prediction function should be controlled.
- Intuitively: place centers where the data is.



Approaches

- 1. Every data point as a center
- 2. Randomly choose centers
- 3. Clustering

Clustering

- Group data points into clusters.
- Cluster centers are good places for RBFs.
- We'll use k-means clustering to pick k centers.

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Lecture 5 | Part 2

k-means Clustering

Digression...

- Let's forget about RBFs for a minute...
- ▶ What is **clustering**?

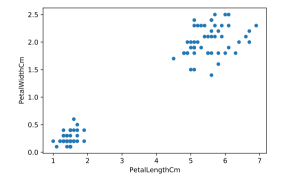
Clustering

- Clustering is a machine learning task whose goal is to find group structure in data.
- ► Why?
 - Exploratory data analysis.
 - Representation learning (today).

Example

- We gather measurements x⁽ⁱ⁾ of a bunch of flowers.
 - Petal length and petal width
- Question: how many species are there?
- **Goal**: **cluster** the similar flowers into groups.

Example



Supervised v. Unsupervised

Clustering is an example of an unsupervised learning task.

Supervised Learning

We tell the machine the "right answer".
 There is a ground truth.

• Data set: $\{(\vec{x}^{(i)}, y_i)\}$.

• **Goal**: learn relationship between features $\vec{x}^{(i)}$ and labels y_i .

Examples: classification, regression.

Unsupervised Learning

We don't tell the machine the "right answer".
 In fact, there might not be one!

• Data set: $\vec{x}^{(i)}$ (usually no test set)

Goal: learn the structure of the data itself.
 To discover something, for compression, to use as a feature later.

Example: clustering

Ground Truth

- If we don't have labels, we can't measure accuracy.
- Sometimes, labels don't exist.
- Example: cluster customers into types by previous purchases.

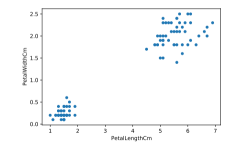
Clustering Approaches

There are many approaches to clustering.

- One of the most popular is k-means clustering.
- It turns clustering into an optimization problem.

Clustering as Optimization

Goal: compress each clustering into a single point while minimizing information loss.

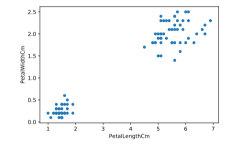


K-Means Objective

- ▶ **Given**: data, $\{\vec{x}^{(i)}\} \in \mathbb{R}^d$ and a parameter k.
- Find: k cluster centers $\vec{\mu}^{(1)}, ..., \vec{\mu}^{(k)}$ so that the average squared distance from a data point to nearest cluster center is small.
- ► The k-means objective function:

$$\mathsf{Cost}(\vec{\mu}^{(1)}, \dots, \vec{\mu}^{(k)}) = \frac{1}{n} \sum_{i=1}^{n} \min_{j \in \{1, \dots, k\}} \|\vec{x}^{(i)} - \vec{\mu}^{(j)}\|^2$$

Clustering as Optimization



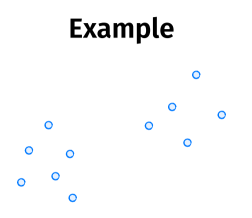
$$\mathsf{Cost}(\vec{\mu}^{(1)}, \dots, \vec{\mu}^{(k)}) = \frac{1}{n} \sum_{i=1}^{n} \min_{j \in \{1, \dots, k\}} \|\vec{x}^{(i)} - \vec{\mu}^{(j)}\|^2$$

Optimization

- **Goal**: find $\vec{\mu}^{(1)}, ..., \vec{\mu}^{(k)}$ minimizing *k*-means objective function.
- Problem: this is NP-Hard.
- We use a heuristic instead of solving exactly.

Lloyd's Algorithm for K-Means

- ▶ Initialize centers, $\vec{\mu}^{(1)}$, ..., $\vec{\mu}^{(k)}$ somehow.
- Repeat until convergence:
 - Assign each point $\overline{\vec{x}}^{(i)}$ to closest center
 - Update each $\vec{\mu}^{(i)}$ to be mean of points assigned to it

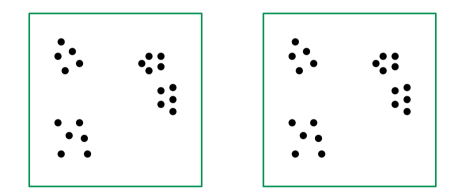


Theory

Each iteration reduces cost.

- ► This guarantees convergence to a **local** min.
- Initialization is very important.

Example



Initialization Strategies

- Basic Approach: Pick k data points at random.
- Better Approach: **k-means++**:
 - Pick first center at random from data.
 - Let $C = {\vec{\mu}^{(1)}}$ (centers chosen so far)
 - Repeat k 1 more times:
 - Pick random data point x according to distribution

$$\mathbb{P}(\vec{x}) \propto \min_{\vec{\mu} \in C} \|\vec{x} - \mu\|^2$$

Add x to C

Picking k

How do we know how many clusters the data contains?

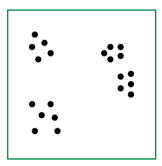
Exercise

How does the **minimum** of the *k*-means objective function change as *k* is increased (that is, as we allow more clusters)? **Hint**: What is the minimum when k = n (we have one cluster per data point).

Plot of K-Means Objective

Observation

- Increasing k always decreases objective function
- But increasing k beyond "true" number of clusters has diminishing returns.



Picking k

► The **elbow method**:

- Run k-means repeatedly with increasing values of k
- Plot the value of the objective as a function of k
- Find an **elbow** in the plot

Applications of K-Means

Discovery

Vector Quantization

Find a finite set of representatives of a large (possibly infinite) set.

Example

- Cluster animal descriptions.
- ▶ 50 animals: grizzly bear, dalmatian, rabbit, pig, ...
- 85 attributes: long neck, tail, walks, swims, ...
- ▶ 50 data points in \mathbb{R}^{85} . Run *k*-means with *k* = 10

Results

🕕 zebra

- 2 spider monkey, gorilla, chimpanzee
- 3 tiger, leopard, wolf, bobcat, lion
- 4 hippopotamus, elephant, rhinoceros
- 6 killer whale, blue whale, humpback whale, seal, walrus, dolphin
- 6 giant panda
- skunk, mole, hamster, squirrel, rabbit, bat, rat, weasel, mouse, raccoon
- B antelope, horse, moose, ox, sheep, giraffe, buffalo, deer, pig, cow
- 9 beaver, otter
- grizzly bear, dalmatian, persian cat, german shepherd, siamese cat, fox, chihuahua, polar bear, collie

zebra

- 2 spider monkey, gorilla, chimpanzee
- 3 tiger, leopard, fox, wolf, bobcat, lion
- hippopotamus, elephant, rhinoceros, buffalo, pig
- (5) killer whale, blue whale, humpback whale, seal, otter, walrus, dolphin
- dalmatian, persian cat, german shepherd, siamese cat, chihuahua, giant panda, collie
- beaver, skunk, mole, squirrel, bat, rat, weasel, mouse, raccoon
- (8) antelope, horse, moose, ox, sheep, giraffe, deer, cow
- I hamster, rabbit
- 🕕 grizzly bear, polar bear

K-Means

- Perhaps the most popular clustering algorithm.
- **Fast, easy to understand.**
- Assumes spherical clusters.

Example

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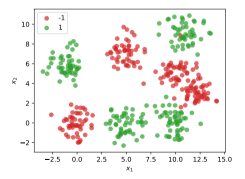
Lecture 5 | Part 3

K-Means for Finding RBF Centers

Idea

Use k-means centers as RBF centers.

▶ Typically "over-cluster" by setting *k* to be large.



Workflow

- "Over-cluster" with k-means to find centers
- Create new features using k RBFs
- Fit a least squares classifier

The Data

	×1	x2
0	0.496714	-0.138264
1	0.647689	1.523030
2	-0.234153	-0.234137
3	1.579213	0.767435
4	-0.469474	0.542560
395	10.429618	0.207688
396	10.271579	-1.276749
397	8.918943	1.053153
398	9.960445	0.681501
399	10.028318	0.029756

400 rows × 2 columns

Step 1) k-means

```
>>> import sklearn.cluster
```

```
>>> # let's start with 20 clusters
```

>>> kmeans = sklearn.cluster.KMeans(n_clusters=20)

```
>>> kmeans.fit(data)
```

```
>>> cluster_centers = kmeans.cluster_centers_
```

```
>>> cluster_centers.shape
```

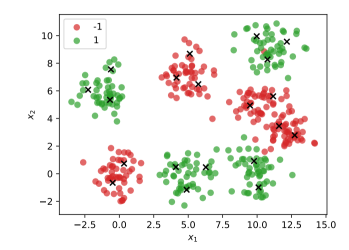
(20, 2)

```
>>> cluster_centers
```

```
array([[ 4.10556507, 0.48176175],
        [ 9.48493465, 4.93921129],
        [-0.67384089, 5.34791854],
        [12.73048608, 2.78757872],
        [12.16178039, 9.56285306],
        [ 4.14585321, 6.95969919],
```

• • •

Cluster Centers



Step 2) Create New Features

• We've found k = 20 cluster centers, $\vec{\mu}^{(1)}, \dots, \vec{\mu}^{(20)}$.

Center a Gaussian RBF at each.

► Take σ = 3 for now.

• We have k = 20 basis functions $\rightarrow 20$ newfeatures for every data point \vec{x} .

Creating the Features

phis = [make_phi(center, 3) for center in cluster_centers]

Example

```
>>> phi_0 = phis[0]
>>> phi_0(np.array([[1, 2], [4, 5], [6, 7]]))
array([0.26507796, 0.10336246, 0.00597847])
```

Applying the Basis Functions

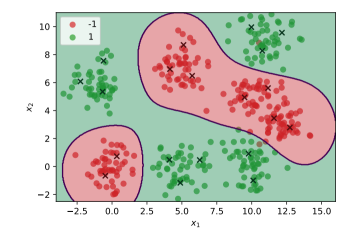
>>> new_features = np.column_stack([phi(data) for phi in phis])
>>> new_features.shape
(400, 20)

Step 3) Fitting the classifier

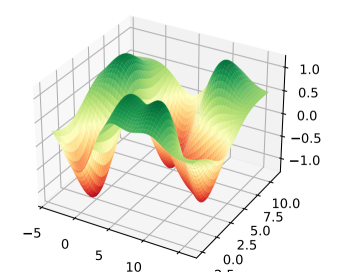
Fitting the Classifier

```
>>> X = augment(new_features)
>>> w = np.linalg.lstsq(X, y)[0]
>>> predictions = np.sign(X @ w)
>>> # training accuracy
>>> (predictions == y).mean()
0.995
```

Decision Boundary



Prediction Surface



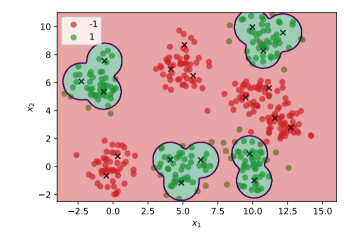
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What if...

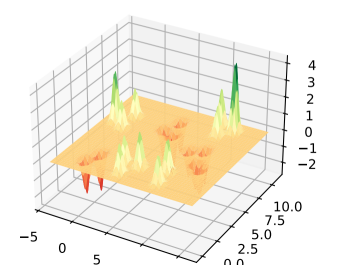
• What if we make σ smaller?

• Set
$$\sigma = \frac{1}{2}$$

Small Sigma



Small Sigma



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Model Complexity

- RBF network complexity is determined by:
 Number of basis functions (more = more complex)
 RBF width parameter (smaller = more complex)
- Choose via cross validation
- More complex = greater danger of overfitting

Representation Learning

- This class is about "representation learning"
- This is the first time we've actually learned a representation.
- Previously: chose basis functions by hand.
- Today: we used k-means to learn good basis functions using the data.