

## Lecture 6: Classification Algorithms

**Applied Machine Learning** 

Volodymyr Kuleshov Cornell Tech

#### Part 1: Classification

So far, every supervised learning algorithm that we've seen has been an instance of regression.

Next, let's look at some classification algorithms. First, we will define what classification is.

# Review: Components of A Supervised Machine Learning Problem

At a high level, a supervised machine learning problem has the following structure:

```
\underbrace{\text{Training Dataset}}_{\text{Attributes} + \text{Features}} + \underbrace{\text{Learning Algorithm}}_{\text{Model Class} + \text{Objective} + \text{Optimizer}} \rightarrow \text{Predictive Model}
```

#### Regression vs. Classification

Consider a training dataset  $\mathcal{D} = \{(x^{(1)}, y^{(1)}), (x^{(2)}, y^{(2)}), \dots, (x^{(n)}, y^{(n)})\}.$ 

We distinguish between two types of supervised learning problems depnding on the targets  $y^{(i)}$ .

- 1. **Regression**: The target variable  $y \in \mathcal{Y}$  is continuous:  $\mathcal{Y} \subseteq \mathbb{R}$ .
- 1. Classification: The target variable y is discrete and takes on one of K possible values:  $\mathcal{Y} = \{y_1, y_2, \dots y_K\}$ . Each discrete value corresponds to a *class* that we want to predict.

### **Binary Classification**

An important special case of classification is when the number of classes K=2.

In this case, we have an instance of a binary classification problem.

#### **Classification Dataset: Iris Flowers**

To demonstrate classification algorithms, we are going to use the Iris flower dataset.

It's a classical dataset originally published by <u>R. A. Fisher</u> (<a href="https://en.wikipedia.org/wiki/Ronald Fisher">https://en.wikipedia.org/wiki/Ronald Fisher</a>) in 1936. Nowadays, it's widely used for demonstrating machine learning algorithms.

```
In [1]:
        import numpy as np
        import pandas as pd
        from sklearn import datasets
         import warnings
        warnings.filterwarnings('ignore')
        # Load the Iris dataset
         iris = datasets.load iris(as frame=True)
        print(iris.DESCR)
         .. _iris_dataset:
        Iris plants dataset
        **Data Set Characteristics:**
             :Number of Instances: 150 (50 in each of three classes)
             :Number of Attributes: 4 numeric, predictive attributes and the class
             :Attribute Information:
                - sepal length in cm
                - sepal width in cm
                - petal length in cm
                - petal width in cm
                - class:
                         - Iris-Setosa
                         - Iris-Versicolour
                         - Iris-Virginica
             :Summary Statistics:
                                                   Class Correlation
                                                SD
                             Min Max
                                        Mean
```

sepal length: 4.3 7.9 5.84 0.83 0.7826 sepal width: 2.0 4.4 3.05 0.43 -0.4194petal length: 1.0 6.9 3.76 1.76 0.9490 (high!) petal width: 0.1 2.5 1.20 0.76 0.9565 (high!)

:Missing Attribute Values: None

:Class Distribution: 33.3% for each of 3 classes.

:Creator: R.A. Fisher

:Donor: Michael Marshall (MARSHALL%PLU@io.arc.nasa.gov)

:Date: July, 1988

The famous Iris database, first used by Sir R.A. Fisher. The dataset is taken from Fisher's paper. Note that it's the same as in R, but not as in the UCI Machine Learning Repository, which has two wrong data points.

This is perhaps the best known database to be found in the pattern recognition literature. Fisher's paper is a classic in the field and is referenced frequently to this day. (See Duda & Hart, for example.) The data set contains 3 classes of 50 instances each, where each class refers to a type of iris plant. One class is linearly separable from the other 2; the latter are NOT linearly separable from each other.

#### .. topic:: References

- Fisher, R.A. "The use of multiple measurements in taxonomic problems" Annual Eugenics, 7, Part II, 179-188 (1936); also in "Contributions to Mathematical Statistics" (John Wiley, NY, 1950).
- Duda, R.O., & Hart, P.E. (1973) Pattern Classification and Scene Analysis. (Q327.D83) John Wiley & Sons. ISBN 0-471-22361-1. See page 218.
- Dasarathy, B.V. (1980) "Nosing Around the Neighborhood: A New System Structure and Classification Rule for Recognition in Partially Exposed Environments". IEEE Transactions on Pattern Analysis and Machine Intelligence, Vol. PAMI-2, No. 1, 67-71.
- Gates, G.W. (1972) "The Reduced Nearest Neighbor Rule". IEEE Transactions

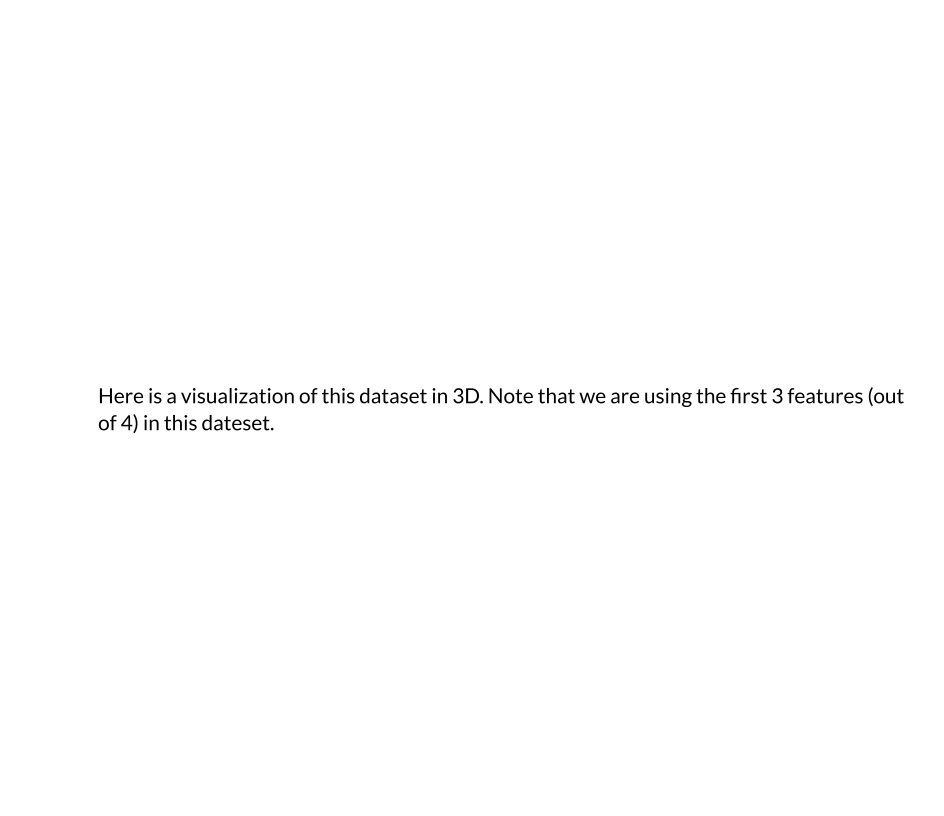
on intormation ineory, may 1912, 431-433.

- See also: 1988 MLC Proceedings, 54-64. Cheeseman et al"s AUTOCLASS II conceptual clustering system finds 3 classes in the data.
- Many, many more ...

In [2]: | # print part of the dataset iris\_X, iris\_y = iris.data, iris.target pd.concat([iris\_X, iris\_y], axis=1).head()

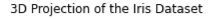
#### Out[2]:

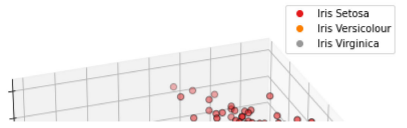
	sepal length (cm)	sepal width (cm)	petal length (cm)	petal width (cm)	target
0	5.1	3.5	1.4	0.2	0
1	4.9	3.0	1.4	0.2	0
2	4.7	3.2	1.3	0.2	0
3	4.6	3.1	1.5	0.2	0
4	5.0	3.6	1.4	0.2	0

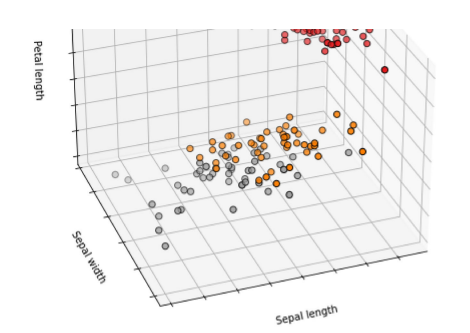


```
In [3]:
        # Code from: https://scikit-learn.org/stable/auto_examples/datasets/plot_iris_data
         set.html
        %matplotlib inline
         import matplotlib.pyplot as plt
        from mpl_toolkits.mplot3d import Axes3D
        from sklearn.decomposition import PCA
        # let's visualize this dataset
        fig = plt.figure(1, figsize=(8, 6))
        ax = Axes3D(fig, elev=-150, azim=110)
         # X_reduced = PCA(n_components=3).fit_transform(iris_X)
        X_reduced = iris_X.to_numpy()[:,:3]
         ax.set_title("3D Projection of the Iris Dataset")
         ax.w_xaxis.set_ticklabels([])
         ax.w_yaxis.set_ticklabels([])
         ax.set_xlabel("Sepal length")
         ax.set_ylabel("Sepal width")
        ax.set_zlabel("Petal length")
        ax.w zaxis.set ticklabels([])
         p1 = ax.scatter(X_reduced[:, 0], X_reduced[:, 1], X_reduced[:, 2], c=iris_y,
                    cmap=plt.cm.Set1, edgecolor='k', s=40)
         plt.legend(handles=p1.legend_elements()[0], labels=['Iris Setosa', 'Iris Versicolo
        ur', 'Iris Virginica'])
```

#### Out[3]: <matplotlib.legend.Legend at 0x129076cc0>







#### **Understanding Classification**

How is clasification different from regression?

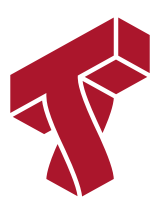
- In regression, we try to fit a curve through the set of targets  $y^{(i)}$ .
- In classification, classes define a partition of the feature space, and our goal is to find the boundaries that separate these regions.
- Outputs of classification models have a simple probabilistic interpretation: they are probabilities that a data point belongs to a given class.

Let's visualize our Iris dataset to see this. Note that we are using the first 2 features in this dateset.

Let's train a classification algorithm on this data.

Below, we see the regions predicted to be associated with the blue and non-blue classes and the line between them in the decision boundary.

```
In [5]: | from sklearn.linear_model import LogisticRegression
        logreg = LogisticRegression(C=1e5)
        # Create an instance of Logistic Regression Classifier and fit the data.
        X = iris_X.to_numpy()[:,:2]
        # rename class two to class one
         iris_y2 = iris_y.copy()
         iris_y2[iris_y2==2] = 1
        Y = iris_y2
        logreg.fit(X, Y)
        xx, yy = np.meshgrid(np.arange(4, 8.2, .02), np.arange(1.8, 4.5, .02))
        Z = logreg.predict(np.c_[xx.ravel(), yy.ravel()])
        # Put the result into a color plot
        Z = Z.reshape(xx.shape)
        plt.pcolormesh(xx, yy, Z, cmap=plt.cm.Paired)
        # Plot also the training points
         plt.scatter(X[:, 0], X[:, 1], c=Y, edgecolors='k', cmap=plt.cm.Paired, s=50)
        plt.xlabel('Sepal length')
         plt.ylabel('Sepal width')
        plt.show()
```



### Part 2: Nearest Neighbors

Previously, we have seen what defines a classification problem. Let's now look at our first classification algorithm.

#### **Review: Classification**

Consider a training dataset  $\mathcal{D} = \{(x^{(1)}, y^{(1)}), (x^{(2)}, y^{(2)}), \dots, (x^{(n)}, y^{(n)})\}.$ 

We distinguish between two types of supervised learning problems depnding on the targets  $y^{(i)}$ .

- 1. **Regression**: The target variable  $y \in \mathcal{Y}$  is continuous:  $\mathcal{Y} \subseteq \mathbb{R}$ .
- 2. **Classification**: The target variable y is discrete and takes on one of K possible values:  $\mathcal{Y} = \{y_1, y_2, \dots y_K\}$ . Each discrete value corresponds to a *class* that we want to predict.

### A Simple Classification Algorithm: Nearest Neighbors

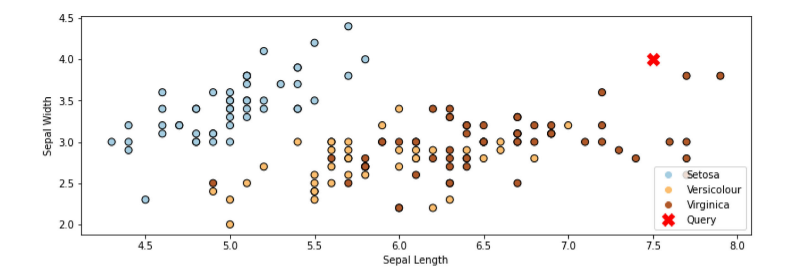
Suppose we are given a training dataset  $\mathcal{D}=\{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\ldots,(x^{(n)},y^{(n)})\}$ . At inference time, we receive a query point x' and we want to predict its label y'.

A really simple but suprisingly effective way of returning  $y^\prime$  is the *nearest neighbors* approach.

- Given a query datapoint x', find the training example (x,y) in  $\mathcal D$  that's closest to x', in the sense that x is "nearest" to x'
- Return y, the label of the "nearest neighbor" x.

In the example below on the Iris dataset, the red cross denotes the query  $x^\prime$ . The closest class to it is "Virginica". (We're only using the first two features in the dataset for simplicity.)

Out[6]: <matplotlib.legend.Legend at 0x12982b4e0>



#### **Choosing a Distance Function**

How do we select the point x that is the closest to the query point x'? There are many options:

- The Euclidean distance  $||x-x'||_2 = \sqrt{\sum_{j=1}^d |x_j-x_j'|^2}$  is a popular choice.
- The Minkowski distance  $||x-x'||_p=(\sum_{j=1}^d|x_j-x_j'|^p)^{1/p}$  generalizes the Euclidean, L1 and other distances.
- Discrete-valued inputs can be examined via the Hamming distance  $|\{j: x_j \neq x_j'\}|$  and other distances.

Let's apply Nearest Neighbors to the above dataset using the Euclidean distance (or equiavalently, Minkowski with p=2)

```
In [23]: | # https://scikit-learn.org/stable/auto_examples/neighbors/plot_classification.html
         from sklearn import neighbors
         from matplotlib.colors import ListedColormap
         # Train a Nearest Neighbors Model
         clf = neighbors.KNeighborsClassifier(n_neighbors=1, metric='minkowski', p=2)
         clf.fit(iris_X.iloc[:,:2], iris_y)
         # Create color maps
         cmap_light = ListedColormap(['orange', 'cyan', 'cornflowerblue'])
         cmap_bold = ListedColormap(['darkorange', 'c', 'darkblue'])
         # Plot the decision boundary. For that, we will assign a color to each
          # point in the mesh [x_min, x_max]x[y_min, y_max].
         x_{min}, x_{max} = iris_X.iloc[:, 0].min() - 1, <math>iris_X.iloc[:, 0].max() + 1
         y_{min}, y_{max} = iris_X.iloc[:, 1].min() - 1, iris_X.iloc[:, 1].max() + 1
         xx, yy = np.meshgrid(np.arange(x_min, x_max, 0.02),
                               np.arange(y_min, y_max, 0.02))
         Z = clf.predict(np.c_[xx.ravel(), yy.ravel()])
         # Put the result into a color plot
         Z = Z.reshape(xx.shape)
         plt.figure()
         plt.pcolormesh(xx, yy, Z, cmap=cmap_light)
         # Plot also the training points
         plt.scatter(iris_X.iloc[:, 0], iris_X.iloc[:, 1], c=iris_y, cmap=cmap_bold,
                      edgecolor='k', s=60)
         plt.xlim(xx.min(), xx.max())
         plt.ylim(yy.min(), yy.max())
          plt.xlabel('Sepal Length')
          plt.ylabel('Sepal Width')
```

In the above example, the regions of the 2D space that are assigned to each class are highly irregular. In areas where the two classes overlap, the decision of the boundary flips between the classes, depending on which point is closest to it.

#### K-Nearest Neighbors

Intuitively, we expect the true decision boundary to be smooth. Therefore, we average K nearest neighbors at a query point.

- Given a query datapoint x', find the K training examples  $\mathcal{N}=\{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\ldots,(x^{(K)},y^{(K)})\}\subseteq D$  that are closest to x'.
- Return  $y_{\mathcal{N}}$ , the consensus label of the neighborhood  $\mathcal{N}$ .

The consesus  $y_{\mathcal{N}}$  can be determined by voting, weighted average, etc.

Let's look at Nearest Neighbors with a neighborhood of 30. The decision boundary is much smoother than before.

```
In [8]: | # https://scikit-learn.org/stable/auto_examples/neighbors/plot_classification.html
        # Train a Nearest Neighbors Model
        clf = neighbors.KNeighborsClassifier(n_neighbors=30, metric='minkowski', p=2)
        clf.fit(iris_X.iloc[:,:2], iris_y)
        Z = clf.predict(np.c_[xx.ravel(), yy.ravel()])
        # Put the result into a color plot
        Z = Z.reshape(xx.shape)
        plt.figure()
        plt.pcolormesh(xx, yy, Z, cmap=cmap_light)
        # Plot also the training points
        plt.scatter(iris_X.iloc[:, 0], iris_X.iloc[:, 1], c=iris_y, cmap=cmap_bold,
                     edgecolor='k', s=60)
        plt.xlim(xx.min(), xx.max())
        plt.ylim(yy.min(), yy.max())
        plt.xlabel('Sepal Length')
        plt.ylabel('Sepal Width')
```

Out[8]: Text(0, 0.5, 'Sepal Width')

#### **Review: Data Distribution**

We will assume that the dataset is governed by a probability distribution  $\mathbb{P}$ , which we will call the *data distribution*. We will denote this as

$$x,y\sim \mathbb{P}.$$

The training set  $\mathcal{D}=\{(x^{(i)},y^{(i)})\mid i=1,2,\ldots,n\}$  consists of independent and identically distributed (IID) samples from  $\mathbb{P}$ .

#### **KNN Estimates Data Distribution**

Suppose that the output y' of KNN is the average target in the neighborhood  $\mathcal{N}(x')$  around the query x'. Observe that we can write:

$$y' = rac{1}{K} \sum_{(x,y) \in \mathcal{N}(x')} y pprox \mathbb{E}[y \mid x'].$$

- When  $x \approx x'$  and when  $\mathbb P$  is reasonably smooth, each y for  $(x,y) \in \mathcal N(x')$  is approximately a sample from  $\mathbb P(y \mid x')$  (since  $\mathbb P$  doesn't change much around x',  $\mathbb P(y \mid x') \approx \mathbb P(y \mid x)$ ).
- Thus y' is essentially a Monte Carlo estimate of  $\mathbb{E}[y\mid x']$  (the average of K samples from  $\mathbb{P}(y\mid x')$ ).

#### Algorithm: K-Nearest Neighbors

- Type: Supervised learning (regression and classification)
- Model family: Consensus over K training instances.
- Objective function: Euclidean, Minkowski, Hamming, etc.
- **Optimizer**: Non at training. Nearest neighbor search at inference using specialized search algorithms (Hashing, KD-trees).
- Probabilistic interpretation: Directly approximating the density  $P_{\mathrm{data}}(y|x)$ .

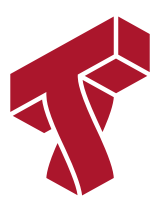
#### **Pros and Cons of KNN**

Pros:

• Can approximate any data distribution arbtrarily well.

#### Cons:

- Need to store entire dataset to make queries, which is computationally prohibitive.
- Number of data needed scale exponentially with dimension ("curse of dimensionality").



#### Part 3: Non-Parametric Models

Nearest neighbors is the first example of an important type of machine learning algorithm called a non-parametric model.

### Review: Supervised Learning Model

We'll say that a model is a function

$$f:\mathcal{X} o\mathcal{Y}$$

that maps inputs  $x \in \mathcal{X}$  to targets  $y \in \mathcal{Y}$ .

Often, models have parameters  $\theta\in\Theta$  living in a set  $\Theta$ . We will then write the model as  $f_{\theta}:\mathcal{X}\to\mathcal{Y}$  to denote that it's parametrized by  $\theta$ .

#### **Review: K-Nearest Neighbors**

Suppose we are given a training dataset

 $\mathcal{D}=\{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\ldots,(x^{(n)},y^{(n)})\}$ . At inference time, we receive a query point x' and we want to predict its label y'.

- Given a query datapoint x', find the K training examples  $\mathcal{N}=\{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\ldots,(x^{(K)},y^{(K)})\}\subseteq D$  that are closest to x'.
- Return  $y_{\mathcal{N}}$ , the consensus label of the neighborhood  $\mathcal{N}$ .

The consesus  $y_{\mathcal{N}}$  can be determined by voting, weighted average, etc.

#### **Non-Parametric Models**

Nearest neighbors is an example of a *non-parametric* model. Parametric vs. non-parametric are is a key distinguishing characteristic for machine learning models.

A parametric model  $f_{\theta}(x): \mathcal{X} imes \Theta o \mathcal{Y}$  is defined by a finite set of parameters  $\theta \in \Theta$  whose dimensionality is constant with respect to the dataset. Linear models of the form  $f_{\theta}(x) = \theta^{\top} x$ 

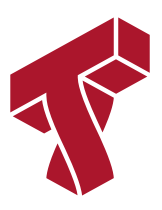
are an example of a parametric model.

In a non-parametric model, the function $f$ uses the entire training datase proccessed version of it) to make predictions, as in $K$ -Nearest Neighbor the complexity of the model increases with dataset size.	

c models have the adv they are also computa		

# Algorithm: K-Nearest Neighbors

- Type: Supervised learning (regression and classification)
- ullet Model family: Non-parametric: Consensus over K training instances.
- Objective function: Euclidean, Minkowski, Hamming, etc.
- **Optimizer**: None at training. Nearest neighbor search at inference using specialized search algorithms (Hashing, KD-trees).
- Probabilistic interpretation: Directly approximating the density  $P_{\mathrm{data}}(y|x)$ .



# Part 4: Logistic Regression

Next, we are going to see a simple parametric classification algorithm that addresses many of these limitations of Nearest Neighbors.

#### **Review: Classification**

Consider a training dataset  $\mathcal{D} = \{(x^{(1)}, y^{(1)}), (x^{(2)}, y^{(2)}), \dots, (x^{(n)}, y^{(n)})\}.$ 

We distinguish between two types of supervised learning problems depnding on the targets  $y^{(i)}$ .

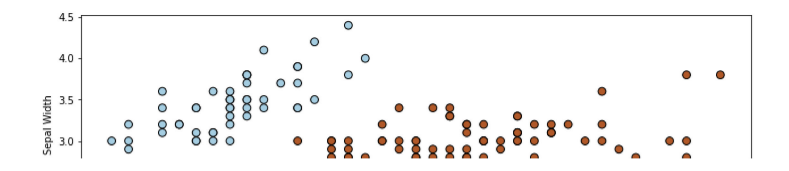
- 1. **Regression**: The target variable  $y \in \mathcal{Y}$  is continuous:  $\mathcal{Y} \subseteq \mathbb{R}$ .
- 2. **Classification**: The target variable y is discrete and takes on one of K possible values:  $\mathcal{Y} = \{y_1, y_2, \dots y_K\}$ . Each discrete value corresponds to a *class* that we want to predict.

# Binary Classification and the Iris Dataset

We are going to start by looking at binary (two-class) classification.

To keep things simple, we will use the Iris dataset. We will be predicting the difference between class 0 (Iris Setosa) and the other two classes.

Out[9]: <matplotlib.legend.Legend at 0x10ac3f278>



## **Review: Least Squares**

Recall that the linear regression algorithm fits a linear model of the form

$$f(x) = \sum_{j=0}^d heta_j \cdot x_j = heta^ op x.$$

It minimizes the mean squared error (MSE)

$$J( heta) = rac{1}{2n} \sum_{i=1}^n (y^{(i)} - heta^ op x^{(i)})^2$$

on a dataset  $\{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\dots,(x^{(n)},y^{(n)})\}.$ 

We could also use the above model for classification problem for which  $\mathcal{Y}=\{0,1\}.$ 

```
In [10]:
         # https://scikit-learn.org/stable/auto_examples/linear_model/plot_iris_logistic.ht
         m7.
          import warnings
         warnings.filterwarnings("ignore")
         from sklearn.linear_model import LinearRegression
         logreg = LinearRegression()
         # Create an instance of Logistic Regression Classifier and fit the data.
         X = iris_X.to_numpy()[:,:2]
         Y = iris_y2
         logreg.fit(X, Y)
         # Plot the decision boundary. For that, we will assign a color to each
         # point in the mesh [x_min, x_max]x[y_min, y_max].
         x_{min}, x_{max} = X[:, 0].min() - .5, X[:, 0].max() + .5
         y_{min}, y_{max} = X[:, 1].min() - .5, X[:, 1].max() + .5
         h = .02 # step size in the mesh
         xx, yy = np.meshgrid(np.arange(x_min, x_max, h), np.arange(y_min, y_max, h))
         Z = logreg.predict(np.c_[xx.ravel(), yy.ravel()])
         Z[Z>0.5] = 1
         Z[Z<0.5] = 0
         # Put the result into a color plot
         Z = Z.reshape(xx.shape)
         plt.pcolormesh(xx, yy, Z, cmap=plt.cm.Paired)
         # Plot also the training points
          plt.scatter(X[:, 0], X[:, 1], c=Y, edgecolors='k', cmap=plt.cm.Paired, s=60)
         plt.xlabel('Sepal length')
         plt.ylabel('Sepal width')
         plt.show()
```

Least squares returns an acceptable decision boundary on this dataset. However, it is problematic for a few reasons.

- There is nothing to prevent outputs larger than one or smaller than zero, which is conceptually wrong
- We also don't have optimal performance: at least one point is misclassified, and others are too close to the decision boundary.

## The Logistic Function

To address this problem, we will look at a different hypothesis class. We will choose models of the form:

$$f(x) = \sigma( heta^ op x) = rac{1}{1 + \exp(- heta^ op x)},$$

where

$$\sigma(z) = rac{1}{1 + \exp(-z)}$$

is known as the sigmoid or logistic function.

The logistic function  $\sigma:\mathbb{R} o [0,1]$  "squeezes" points from the real line into [0,1].

# The Logistic Function: Properties

The sigmoid function is defined as

$$\sigma(z) = rac{1}{1 + \exp(-z)}.$$

A few observations:

- The tends to 1 as  $z \to \infty$  and tends to 0 as  $z \to -\infty$ .
- Thus, models of the form  $\sigma(\theta^{\top}x)$  output values between 0 and 1, which is suitable for binary classification.
- It is easy to show that the derivative of  $\sigma(z)$  has a simple form:  $\frac{d\sigma}{dz}=\sigma(z)(1-\sigma(z)).$

Let's implement our model using the sigmoid function.

```
In [12]: def f(X, theta):
    """The sigmoid model we are trying to fit.

Parameters:
    theta (np.array): d-dimensional vector of parameters
    X (np.array): (n,d)-dimensional data matrix

Returns:
    y_pred (np.array): n-dimensional vector of predicted targets
    """
    return sigmoid(X.dot(theta))
```

## A Probabilistic Approach to Classification

We can take this probabilistic perspective to derive a new algorithm for binary classification.

We will start by using our logistic model to parametrize a probability distribution as follows:

$$egin{aligned} p(y=1|x; heta) &= \sigma( heta^ op x) \ p(y=0|x; heta) &= 1 - \sigma( heta^ op x). \end{aligned}$$

A probability over  $y \in \{0,1\}$  of the form P(y=1)=p is called Bernoulli.

Note that we can write this more compactly as

$$p(y|x; heta) = \sigma( heta^ op x)^y \cdot (1 - \sigma( heta^ op x))^{1-y}$$

#### Review: Conditional Maximum Likelihood

A general approach of optimizing conditional models of the form  $P_{ heta}(y|x)$  is by minimizing expected KL divergence with respect to the data distribution:

$$\min_{ heta} \mathbb{E}_{x \sim \mathbb{P}_{ ext{data}}} \left[ D(P_{ ext{data}}(y|x) \mid\mid P_{ heta}(y|x)) 
ight].$$

With a bit of math, we can show that the maximum likelihood objective becomes

$$\max_{ heta} \mathbb{E}_{x,y \sim \mathbb{P}_{ ext{data}}} \log P_{ heta}(y|x).$$

This is the principle of conditional maximum likelihood.

## Applying Maximum Lilkelihood

Following the principle of maximum likelihood, we want to optimize the following objective defined over a training dataset

$$egin{align} \mathcal{D} &= \{(x^{(1)},y^{(1)}),(x^{(2)},y^{(2)}),\dots,(x^{(n)},y^{(n)})\}.\ &L( heta) = \prod_{i=1}^n p(y^{(i)}\mid x^{(i)}; heta)\ &= \prod_{i=1}^n \sigma( heta^ op x^{(i)})^{y^{(i)}} \cdot (1-\sigma( heta^ op x^{(i)}))^{1-y^{(i)}}. \end{aligned}$$

This log of this objective is also often called the log-loss, or cross-entropy.

This model and objective function define *logistic regression*, one of the most widely used classification algorithms (the name "regression" is an unfortunate misnomer!).

Let's implement the likelihood objective.

```
In [13]: def log_likelihood(theta, X, y):
    """The cost function, J(theta0, theta1) describing the goodness of fit.

    We added the 1e-6 term in order to avoid overflow (inf and -inf).

    Parameters:
    theta (np.array): d-dimensional vector of parameters
    X (np.array): (n,d)-dimensional design matrix
    y (np.array): n-dimensional vector of targets
    """
    return (y*np.log(f(X, theta) + 1e-6) + (1-y)*np.log(1-f(X, theta) + 1e-6)).mea
    n()
```

#### **Review: Gradient Descent**

If we want to optimize  $J(\theta)$ , we start with an initial guess  $\theta_0$  for the parameters and repeat the following update:

$$heta_i := heta_{i-1} - lpha \cdot 
abla_{ heta} J( heta_{i-1}).$$

As code, this method may look as follows:

```
theta, theta_prev = random_initialization()
while norm(theta - theta_prev) > convergence_threshold:
    theta_prev = theta
    theta = theta_prev - step_size * gradient(theta_prev)
```

# Derivatives of the Log-Likelihood

Let's work out the gradient for our log likelihood objective:

$$egin{aligned} rac{\partial \log L( heta)}{\partial heta_j} &= rac{\partial}{\partial heta_j} \log igg( \sigma( heta^ op x) \cdot (1 - \sigma( heta^ op x))^{1-y} igg) \ &= igg( y \cdot rac{1}{\sigma( heta^ op x)} - (1 - y) rac{1}{1 - \sigma( heta^ op x)} igg) rac{\partial}{\partial heta_j} \sigma( heta^ op x) \ &= igg( y \cdot rac{1}{\sigma( heta^ op x)} - (1 - y) rac{1}{1 - \sigma( heta^ op x)} igg) \sigma( heta^ op x) (1 - \sigma( heta^ op x)) rac{\partial}{\partial heta_j} heta^ op x \ &= ig( y \cdot (1 - \sigma( heta^ op x)) - (1 - y) \sigma( heta^ op x) igg) x_j \ &= ig( y - f_ heta(x) ig) x_j. \end{aligned}$$

## Gradient of the Log-Likelihood

Using the above expression, we obtain the following gradient:

$$abla_{ heta}J( heta)=(y-f_{ heta}(x))\cdot\mathbf{x}$$
 .

- This expression looks very similar to the gradient of the mean squared error.
- ullet However, it is different, because the model  $f_{ heta}$  is different.

Let's implement the gradient.

```
In [14]: def loglik_gradient(theta, X, y):
    """The cost function, J(theta0, theta1) describing the goodness of fit.

Parameters:
    theta (np.array): d-dimensional vector of parameters
    X (np.array): (n,d)-dimensional design matrix
    y (np.array): n-dimensional vector of targets

Returns:
    grad (np.array): d-dimensional gradient of the MSE
    """
    return np.mean((y - f(X, theta)) * X.T, axis=1)
```

## **Gradient Descent for Logistic Regression**

Putting this together, we obtain a complete learning algorithm, logistic regression.

```
theta, theta_prev = random_initialization()
while abs(J(theta) - J(theta_prev)) > conv_threshold:
    theta_prev = theta
    theta = theta_prev - step_size * (f(x, theta)-y) * x
```

Let's implement this algorithm.

```
In [15]: | threshold = 5e-5
         step_size = 1e-1
         theta, theta_prev = np.zeros((3,)), np.ones((3,))
         opt_pts = [theta]
         opt_grads = []
          iter = 0
         iris_X['one'] = 1
         X_train = iris_X.iloc[:,[0,1,-1]].to_numpy()
         y_train = iris_y2.to_numpy()
         while np.linalg.norm(theta - theta_prev) > threshold:
         # while True:
              if iter % 50000 == 0:
                  print('Iteration %d. Log-likelihood: %.6f' % (iter, log_likelihood(theta,
         X_train, y_train)))
             theta_prev = theta
             gradient = loglik_gradient(theta, X_train, y_train)
             theta = theta_prev + step_size * gradient
              opt_pts += [theta]
              opt_grads += [gradient]
              iter += 1
```

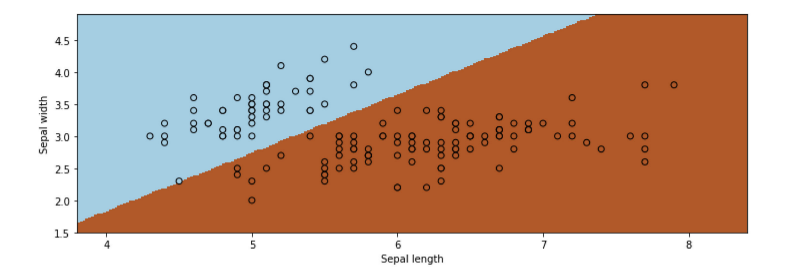
Iteration 0. Log-likelihood: -0.693145 Iteration 50000. Log-likelihood: -0.021506 Iteration 100000. Log-likelihood: -0.015329 Iteration 150000. Log-likelihood: -0.012062 Iteration 200000. Log-likelihood: -0.010076 Let's now visualize the result.

```
In [16]: xx, yy = np.meshgrid(np.arange(x_min, x_max, .02), np.arange(y_min, y_max, .02))
    Z = f(np.c_[xx.ravel(), yy.ravel(), np.ones(xx.ravel().shape)], theta)
    Z[Z<0.5] = 0
    Z[Z \geq 0.5] = 1

# Put the result into a color plot
    Z = Z.reshape(xx.shape)
    plt.pcolormesh(xx, yy, Z, cmap=plt.cm.Paired)

# Plot also the training points
    plt.scatter(X[:, 0], X[:, 1], c=Y, edgecolors='k', cmap=plt.cm.Paired)
    plt.xlabel('Sepal length')
    plt.ylabel('Sepal width')

plt.show()</pre>
```



## Observations About Logistic Regression

• Logistic regression finds a linear decision boundary, is the set of points for which P(y=1|x)=P(y=0|x), or equivalently:

$$0 = \log rac{P(y=1|x)}{P(y=0|x)} = \log rac{rac{1}{1+\exp(- heta^ op x)}}{1-rac{1}{1+\exp(- heta^ op x)}} = heta^ op x$$

The claim holds because  $0=\theta^{ op}x$  is a linear function.

• Unlike least squares, we don't have a closed form solution for  $\theta$ , but we can still apply gradient descent.

# Algorithm: Logistic Regression

- **Type**: Supervised learning (binary classification)
- Model family: Linear decision boundaries.
- Objective function: Cross-entropy, a special case of log-likelihood.
- Optimizer: Gradient descent.
- Probabilistic interpretation: Parametrized Bernoulli distribution.



#### Part 5: Multi-Class Classification

Finally, let's look at an extension of logistic regression to an arbitrary number of classes.

## Review: Logistic Regression

Logistic regression fits models of the form:

$$f(x) = \sigma( heta^ op x) = rac{1}{1 + \exp(- heta^ op x)},$$

where

$$\sigma(z) = rac{1}{1 + \exp(-z)}$$

is known as the sigmoid or logistic function.

#### **Multi-Class Classification**

Linear regression only applies to binary classification problems. What if we have an arbitrary number of classes K?

- The simplest approach that can be used with any machine learning algorithm is the "one vs. all" approach. We train one classifer for each class to distinguish one class from all the others.
- This works, but loses a valid probabilistic interpretation and is not very elegant.
- Alternatively, we may fit a probabilistic model that outputs multi-class probabilities.

Let's load a fully multiclass version of the Iris dataset.

#### The Softmax Function

The logistic function  $\sigma:\mathbb{R} o [0,1]$  be seen as mapping input an input  $ec z\in\mathbb{R}$  to a probability.

Its multi-class extension  $\vec{\sigma}:\mathbb{R}^K o [0,1]^K$ : maps a K-dimensional input  $z\in\mathbb{R}$  to a K-dimensional vector of probabilities.

Each componnent of  $\vec{\sigma}(\vec{z})$  is defined as

$$\sigma(ec{z})_k = rac{\exp(z_k)}{\sum_{l=1}^K \exp(z_l)}.$$

We call this the softmax function.

When K=2, this looks as follows:

$$\sigma(ec{z})_1 = rac{\exp(z_1)}{\exp(z_1) + \exp(z_2)}.$$

We can assume that  $\exp(z_1)=1$  because multiplying the numerator and denominator doesn't change any of the probabilities (so we can just divide by  $\exp(z_1)$ ). Thus we obtain:  $\sigma(\vec{z})_1=rac{1}{1+\exp(z_2)}.$ 

$$\sigma(ec{z})_1 = rac{1}{1+\exp(z_2)}.$$

This is essentially our sigmoid function. Hence softmax generalizes the sigmoid function.

#### The Softmax Model

We can use the softmax function to define a K-class classification model.

In the binary classification setting, we mapped weights  $\theta$  and features x into a probability as follows:

$$\sigma( heta^ op x) = rac{1}{1 + \exp(- heta^ op x)},$$

In the multi-class setting, we define a model  $f: \mathcal{X} \to [0,1]^K$  that outputs the probability of class k based on the features x and class-specific weights  $\theta_k$ :

$$\sigma( heta_k^ op x)_k = rac{\exp( heta_k^ op x)}{\sum_{l=1}^K \exp( heta_l^ op x)}.$$

Its parameter space lies in  $\Theta^K$ , where  $\Theta=\mathbb{R}^d$  is the parameter space of logistic regression.

You may have noticed that this model is slightly over-parametrized: multiplying every  $\theta_k$  by a constant results in an equivalent model. For this reason, it is often assumed that one of the class weights  $\theta_l=0$ .

#### **Softmax Regression**

We again take a probabilistic perspective to derive a K-class classification algorithm based on this model.

We will start by using our softmax model to parametrize a probability distribution as follows:

$$p(y=k|x; heta) = ec{\sigma}( heta^ op x)_k$$

This is called a categorial distribution, and it generalizes the Bernoulli.

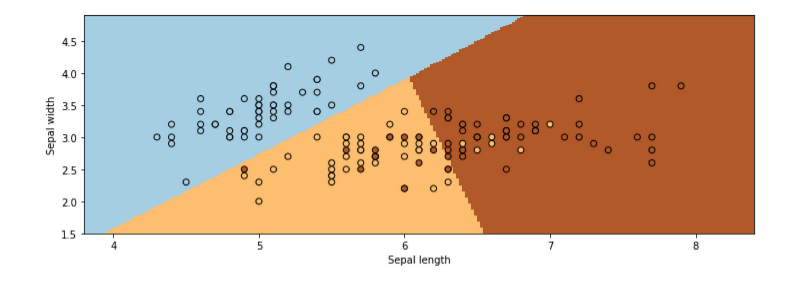
Following the principle of maximum likelihood, we want to optimize the following objective defined over a training dataset

$$\mathcal{D} = \{(x^{(1)}, y^{(1)}), (x^{(2)}, y^{(2)}), \dots, (x^{(n)}, y^{(n)})\}. \ L( heta) = \prod_{i=1}^n p(y^{(i)} \mid x^{(i)}; heta) = \prod_{i=1}^n ec{\sigma}( heta^ op x^{(i)})_{y^{(i)}}$$

This model and objective function define *softmax regression*. (The term "regression" here is again a misnomer.)

Let's now apply softmax regression to the Iris dataset by using the implementation from  ${\tt sklearn}$  .

```
In [19]:
         # https://scikit-learn.org/stable/auto_examples/linear_model/plot_iris_logistic.ht
         ml
         from sklearn.linear_model import LogisticRegression
         logreg = LogisticRegression(C=1e5, multi_class='multinomial')
         # Create an instance of Softmax and fit the data.
         logreg.fit(X, iris_y)
         Z = logreg.predict(np.c_[xx.ravel(), yy.ravel()])
         # Put the result into a color plot
         Z = Z.reshape(xx.shape)
         plt.pcolormesh(xx, yy, Z, cmap=plt.cm.Paired)
         # Plot also the training points
         plt.scatter(X[:, 0], X[:, 1], c=iris_y, edgecolors='k', cmap=plt.cm.Paired)
         plt.xlabel('Sepal length')
         plt.ylabel('Sepal width')
         plt.show()
```



# Algorithm: Softmax Regression

- **Type**: Supervised learning (classification)
- Model family: Linear decision boundaries.
- Objective function: Softmax loss, a special case of log-likelihood.
- Optimizer: Gradient descent.
- Probabilistic interpretation: Parametrized categorical distribution.