

Lecture 5 pt.1: Outline

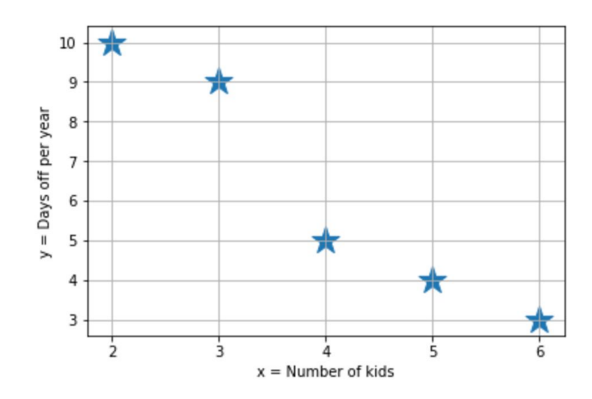
- 1. Linear Regression recap
- 2. Gradient Descent
- 3. Feature scaling
- 4. Intro to Classification
- 5. Logistic Regression
- 6. Example Code

Recap: Linear Regression

Recap: Prediction

Given some data:

X	У
2	10
4	5
3	9
5	4
6	3



Objective: Be able to predict y given new input x

Recap: Simple Linear Regression

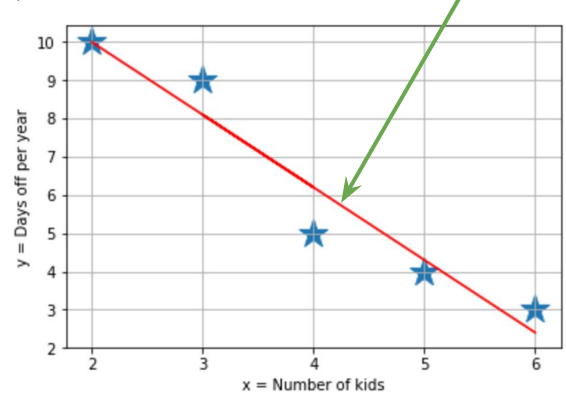
Simple Linear Regression: Prediction hypothesis $h_{\theta}(x)$

$$\hat{y} = f(x, \theta) = h_{\theta}(x) = \theta^T x = \theta_0 + \theta_1 x_1$$

$$x = \begin{bmatrix} 1 \\ x_1 \end{bmatrix}$$
 x is given

Objective: fit the best possible linear function to the training data, l.e. to find the optimal parameters θ

$$\theta = \begin{vmatrix} \theta_0 \\ \theta_1 \end{vmatrix}$$



Recap: Cost function (MSE)

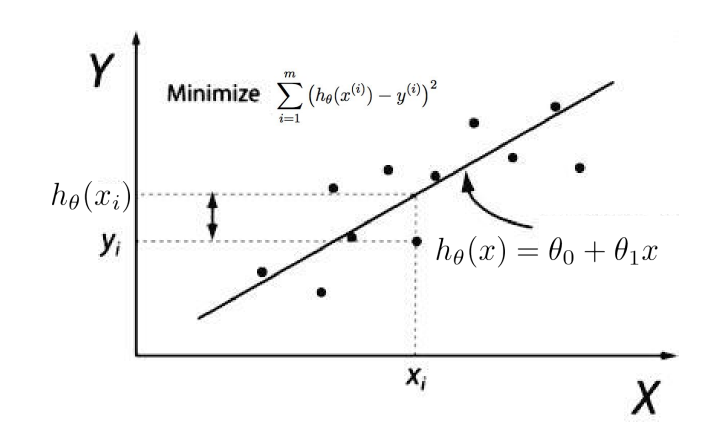
Simple Linear Regression

$$\hat{y} = h_{\theta}(x) = \theta_0 + \theta_1 x_1$$

Cost function:

Measures how good the fit is (MSE)

$$J(\theta) = \frac{1}{2m} \sum_{i=1}^{m} \left(h_{\theta}(x^{(i)}) - y^{(i)} \right)^{2}$$





Recap: Multiple Linear Regression

Multiple Linear Regression: $\hat{y} = h_{\theta}(x) = \theta_0 + \theta_1 x_1 + \theta_2 x_2 + ... + \theta_n x_n = \theta^T X$

$$\theta = \begin{bmatrix} \theta_0 \\ \theta_1 \\ \vdots \\ \vdots \\ \theta_n \end{bmatrix} \text{ is the parameter vector and }$$

$$X = \begin{bmatrix} x_0^{(1)} & x_1^{(1)} & \dots & x_n^{(1)} \\ x_0^{(2)} & x_1^{(2)} & \dots & x_n^{(2)} \\ \vdots & \vdots & \ddots & \vdots \\ x_0^{(m)} & x_1^{(m)} & \dots & x_n^{(m)} \end{bmatrix}$$
is the feature vector and

$$h_{\theta}(X) = \begin{bmatrix} h_{\theta}(x^{(1)}) \\ h_{\theta}(x^{(2)}) \\ \vdots \\ h_{\theta}(x^{(m)}) \end{bmatrix}$$
 is the hypotheses vector

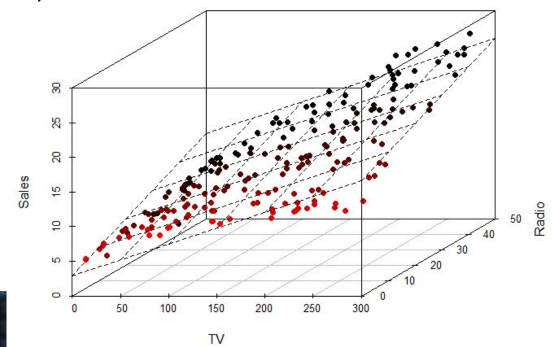
Source: 3.bp.blogspot.com/

Example of multiple linear regression (2 features)

x 1 = TV advertising

x_2 = Radio advertising

y = Sales



Recap: Minimize cost function

Optimal parameters are found when the cost function / the error $J(\theta)$ is minimized

$$J(\theta) = \frac{1}{2m} \sum_{i=1}^{m} \left(h_{\theta}(x^{(i)}) - y^{(i)} \right)^{2}$$

$$J(\theta) = \frac{1}{2m} (X\theta - y)^T (X\theta - y)$$

$$\min_{\theta} J(\theta)$$

$$\frac{\partial J}{\partial \theta} = 2X^T X \theta - 2X^T y = 0$$

Minimize by taking the derivative w.r.t. θ = 0

Normal equation for Linear Regression

Closed form, analytical solution. Finds θ that minimizes $J(\theta)$

$$\theta = (X^T X)^{-1} X^T y$$

Pros:

- Finds optimal answer with one calculation
- Really quick for small data sets

Cons:

- $\mathcal{O}(n^3)$ complexity, **slow**, because of matrix inverse
- $(X^TX)^{-1}$ might not be invertible (can be solved by taking pseduo-inverse)

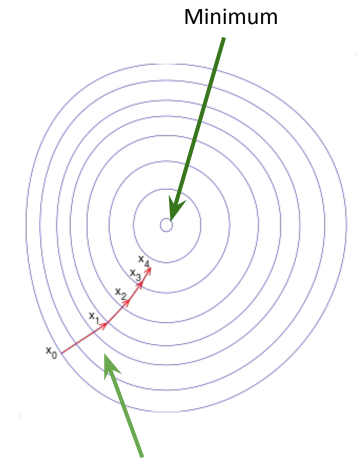
Gradient Descent

Introducing Gradient Descent

WIKIPEDIA:

Gradient descent is a an iterative optimization algorithm for finding the minimum of a function.

To reach minima one takes steps proportional to the negative of the gradient (or approximate gradient) of the function at the current point.



Step sizes & neg. gradient directions

Introducing Gradient Descent

Alternative way of minimizing the cost function:

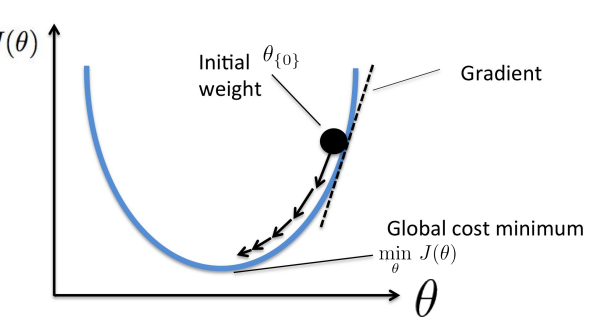
$$J(\theta) = \frac{1}{2m} \sum_{i=1}^{m} \left(h_{\theta}(x^{(i)}) - y^{(i)} \right)^{2}$$

- Gradient Descent minimizes $J(\theta)$ iteratively,
- Will always converge because J(θ) is convex

- Start with / initialize θ_0, θ_1 . E.g. $(\theta_0, \theta_1) = (0, 0)$
- Keep changing θ_0, θ_1 to reduce $J(\theta_0, \theta_1)$,

Illustration of Gradient Descent

for one parameter θ



Source: https://sebastianraschka.com



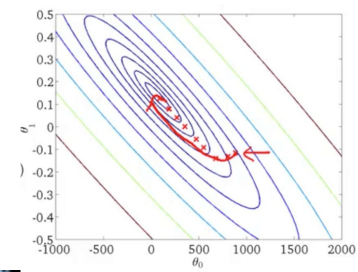
Gradient Descent Algorithm: Linear Regression

- 1. Calculate the gradient $\frac{\partial}{\partial \theta_i} J(\theta)$ for all j
- 2. Form the **update rule** for every parameter:

$$\theta_{j,iter+1} := \theta_{j,iter} - \alpha \frac{\partial}{\partial \theta_j} J(\theta) = \theta_{j,iter} - \alpha / m \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)}) x_j^{(i)}$$

- 3. Choose a step size/ learning rate α (often between 10^-6 and 10^2 -- not too big, then divergence).
- 4. Update all the parameters $\theta_1..\theta_n$ at once (this is called "batch Gradient descent)
- 5. Stop when the error has converged.

$$J_{train}(\theta) = \frac{1}{2m} \sum_{i=1}^{m} (h_{\theta}(x^{(i)}) - y^{(i)})^2$$
 Repeat {
$$\theta_j := \theta_j - \alpha \frac{1}{m} \sum_{i=1}^{m} (h_{\theta}(x^{(i)}) - y^{(i)}) x_j^{(i)}$$
 (for every $j = 0, \dots, n$)



Gradient Descent Tips

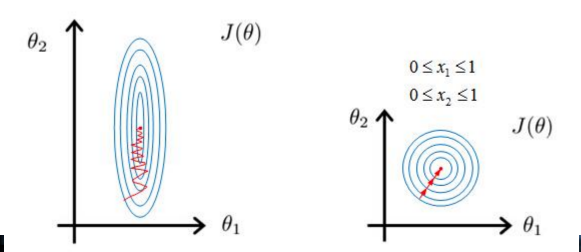
Featuring Scaling

Gradient Descent will be more likely to converge and be faster if the features are scaled, ie $-1 \le x_i \le 1$

For all features:

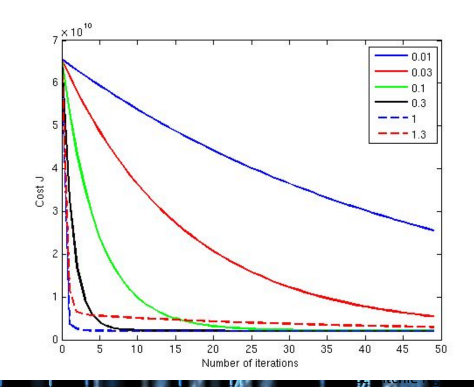
- Subtract mean
- Divide by st.dev.

$$x_i \leftarrow \frac{x_i - \mu(x_i)}{\sigma(x_i)}$$



Monitor convergence

Plot the error function for every iteration / batch parameter update. Check that the error becomes smaller and that we have chosen a suitable learning rate.



Gradient Descent Pros / Cons

Pros

- Will always converge to minima if learning rate is chosen correctly
- Fast (time complexity is $\mathcal{O}(n)$)

Cons

- We have to choose a learning rate α and initialize the parameters
- Often takes A LOT of iterations to reach global minima of the objective function



Classification

Regression vs. Classification

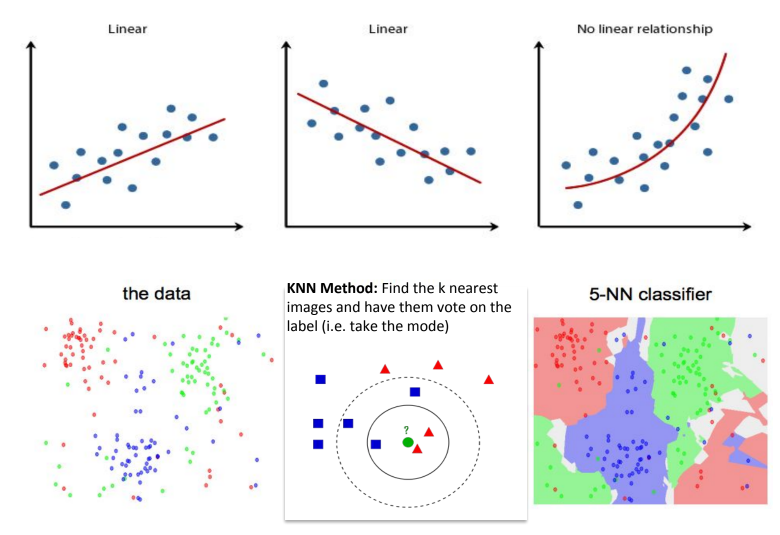
Regression:

- Continuous output y
- Quantitative approach
- Linear or Non-linear

Classification:

- Discrete output y
- Qualitative approach
- Linear or Non-linear

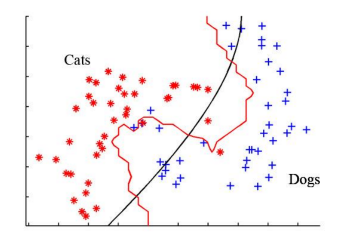
Ex. KNN, K-means Logitstic, SVM, ..



Examples of classification

Examples

- Weather: Sunny / Rainy
- Spam Detection
- Image Classification: Cats VS Dogs
- Image Classification: Recognizing Digits







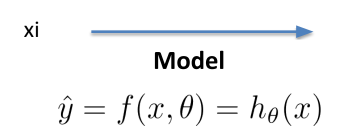


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Our Goal: To classify items

i.e. find the best hypothesis model $h_{ heta}(x)$ that maps x to y





We have this:

(X,Y): (x1,y1), (x2, y2) .. (xn,yn)

- xi is a vector (or even matrix) for each data element
- **Example:** xi = [12 15 22] = [height, weight, color]
- For a picture: $x_i = [32 \times 32 \times 3]$: array of numbers

Binary classification (cat vs dog):

y = 1 if picture is dog

 $y \in \{0, 1\}$

Y = 0 if picture is cat

Multi-class classication:

$$y_{i} = [y_{i,1}, y_{i,2}, ...y_{i,k}]$$

$$y_{i} = [+1, 0, ... 0]*$$

 $y \in \{0, 1, 2..k\}$

Y(i,0) = 1 if picture is a dog

Y(i,1) = 1 if picture is a cat

Y(i,2) = 1 if picture is a elephant etc.

^{*} Sometimes: y ; = [+1,-1,..-1] -1 or 1 instead of 0 or 1

Our Goal: To classify items.

We have this: (X,Y)



Model:
$$h_{\theta}(x)$$

Actual Results:

$$y_{i} = [y_{i,1}, y_{i,2}, ...y_{i,k}]$$

 $y_{i} = [+1, 0, ... 0]$

Machine Learning Steps to train a classifier model

1. Choose our model: $h_{\theta}(x)$ = our estimate of Y

Linear model: $f(x,\theta) = h_{\theta}(x) = \theta x_i = \theta_0 + \theta_1 x_{i,1} + \theta_2 x_{i,2} \dots$

 θ is list of parameters.

Results in a different estimated value of Yi for each sample point x_i

- 2. Define a loss function $(J(\theta))$ = which is a function $(Y_atual, vs Y_estimated)$
- 3. Optimize across the parameter space (θ) to minimize the loss function (to some small threshold)

Logistic Regression

Example Classification with Logistic Regression

Data: students study for an exam

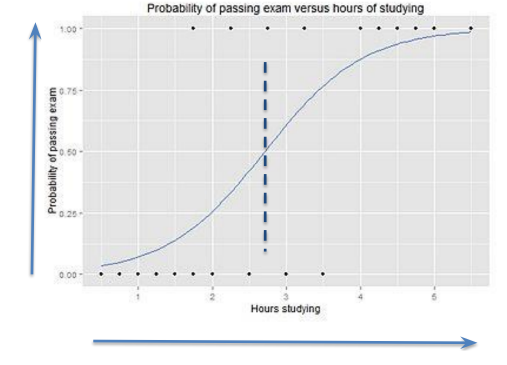
(x= hours studied, y = pass/not pass)

Hours	0.50	0.75	1.00	1.25	1.50	1.75	1.75	2.00	2.25	2.50	2.75	3.00	3.25	3.50	4.00	4.25	4.50	4.75	5.00	5.50
Pass	0	0	0	0	0	0	1	0	1	0	1	0	1	0	1	1	1	1	1	1

y, binary output

0 = fail

1 = pass



Problem:

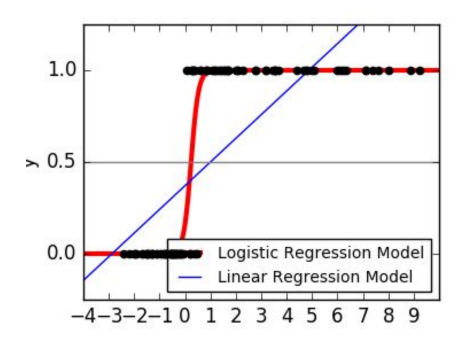
Student studies x hours
We want to predict will the student pass?

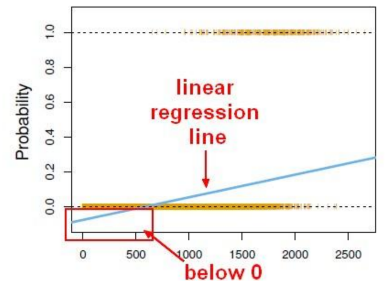
We use this curve to predict the probability that the student would pass given x hours of study

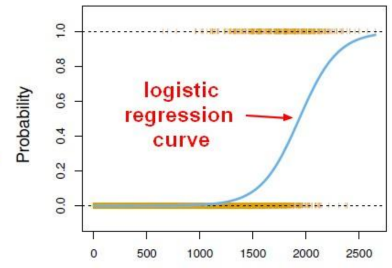
If Prob >= 0.5, we classify student will pass the exam. I.e. y=1

If Prob < 0.5, classify student will fail i.e. y=0

Example: Logistic Regression We use Logistic Regression because a line is not a good estimator for binary results (classification)





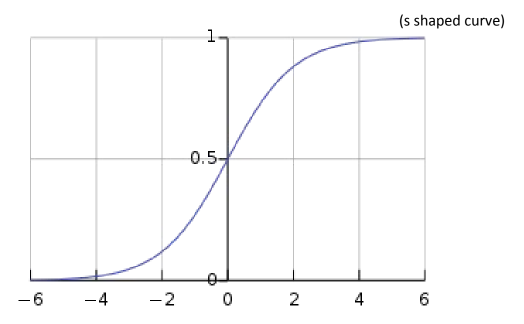


The logistic / sigmoid function is a better fit for binary results

The sigmoid function is:

$$\mathsf{z(t)} = rac{e^t}{e^t + 1} = rac{1}{1 + e^{-t}} \qquad egin{array}{c} \mathsf{Large} \ \mathsf{t} imes \mathsf{1} \\ \mathsf{Small} \ \mathsf{t} o \mathsf{0} \end{array}$$

This function only takes values between 0 and 1 for all real numbers (like a probability)



And if t has this form:

$$t = \theta_0 + x_1 \theta_1$$
 (a line)

$$z(t) = z(\theta^T x) = \frac{1}{1 + e^{-\theta_0 + \theta_1 x_1}} = h_{\theta}(x)$$

If θ_1 is small \rightarrow slow rise If θ_1 is large \rightarrow fast rise

- Think of $f(x,\theta)$ as probability y = 1 given any x
- Prob (y=1) = $\frac{1}{2}$ when $e^{-(\theta_{0} + x_{1}, 1\theta_{1})} = 1$, ie $\theta_{0} + x_{1}\theta_{1} = 0$
- Choose all θ to get best fit
- Still need a **loss function** $J(\theta)$, then solve for best θ

Decision Boundary

The decision boundary separates our predicted categories from one another, in the feature space.

If we have two inputs, x_1 and x_2 , the decision boundary is the line when the predicted probability for either y=0 or y=1 equals 50%

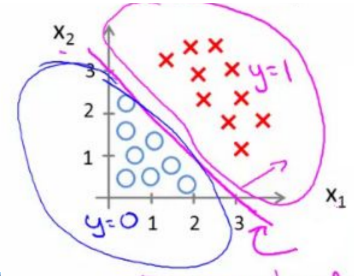
$$h_{\theta}(x) = z(\theta^T X) = \frac{1}{1 + e^{-(\theta_0 + \theta_1 x_1 + \theta_2 x_2)}} = 0.5$$

$$\Leftrightarrow$$

$$\theta_0 + \theta_1 x_1 + \theta_2 x_2 = 0$$

Example

$$\theta_{0}$$
 = -3 θ_{1} = 1 θ_{2} = 1 Then $x_{1} + x_{2} - 3 \geq 0$ will predict y=1 and vice versa (see example below)



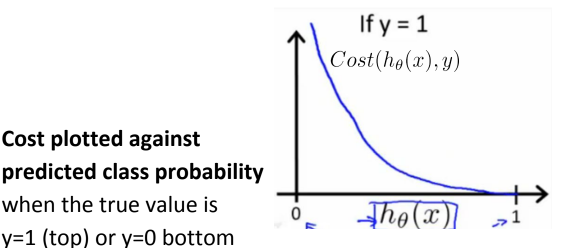
Derivation of the logistic cost function

Cost plotted against

Intuition:

- How do we get the optimal decision boundary?
- Output y can only take on two values (0 or 1)
- We want a cost function that penalizes when our prediction $h_{\theta}(x)$ is wrong

when the true value is y=1 (top) or y=0 bottom

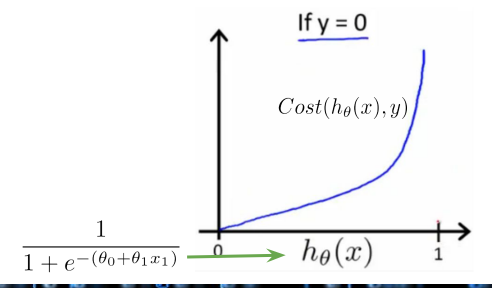


Our total overall cost is $J(\theta)$

$$J(\theta) = \frac{1}{m} \sum_{i=1}^{m} \operatorname{Cost}(h_{\theta}(x^{(i)}), y^{(i)})$$

$$\operatorname{Cost}(h_{\theta}(x), y) = -\log(h_{\theta}(x)) \quad \text{if } y = 1$$

$$\operatorname{Cost}(h_{\theta}(x), y) = -\log(1 - h_{\theta}(x)) \quad \text{if } y = 0$$



Logistic cost function

Cross Entropy =

Note: Loss Function on the former slide can be added to form cross entropy.

We choose this cost function, because it can be derived from the Maximum Likelihood estimation of the parameters.



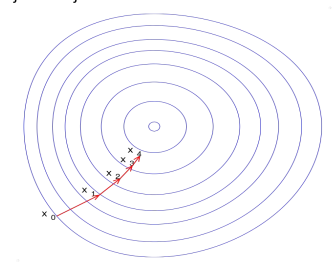
Gradient Descent & Logistic Regression

 $J(\theta)$ = is a cost a function of our estimate $h_{\theta}(x)$ and the true y.

Try to find optimal θ (first initialize with some random value)

Take small steps in the direction where $J(\theta)$ is decreasing

<u>Update rule:</u> $\theta_{i+1} = \theta_i - [(\text{step size } \alpha) \times - \text{ gradient of } J(\theta)]$



Formal update rule

looks exactly like Linear Regression, but note that $h_{\theta}(x)$ has changed)

$$J(\theta) = \frac{-1}{m} \left[\sum_{i=1}^{m} y^{(i)} \log h_{\theta}(x^{(i)}) + (1 - y^{(i)}) \log (1 - h_{\theta}(x^{(i)})) \right]$$

Repeat {
$$\theta_{j} := -\alpha \frac{\partial}{\partial \theta_{j}} J(\theta)$$
 }

Repeat {
$$\theta_j := \theta_j - \alpha \sum_{i=1}^m (h_{\theta}(x^{(i)}) - y^{(i)}) x_j^{(i)}$$





Multi-class Logistic Regression: One-vs-all y∈{0,1...k}

Sigmoid function:

$$z(t) = rac{e^t}{e^t + 1} = rac{1}{1 + e^{-t}}$$

Large t -> 1, Small t -> 0

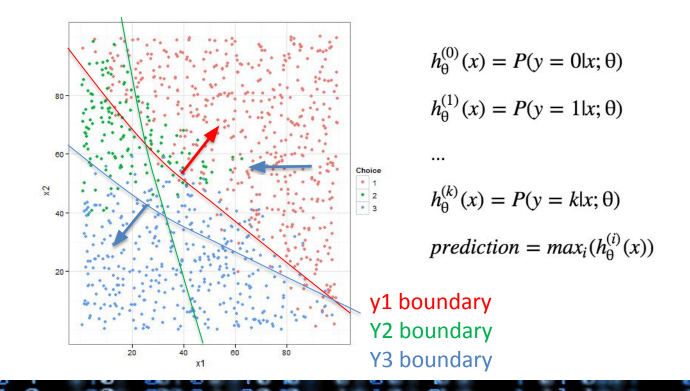
And if t has this form $t = \theta x_i$ (in matrix form) = $\theta_1 x_{i,1} + \theta_2 x_{i,2}$...

$$z(t) = z(\theta^T x) = h_{\theta}(x) = \frac{1}{1 + e^{-\theta^T x}} = \frac{1}{1 + e^{(-\theta_0 + \theta_1 x_1 + \theta_2 x_2 + \dots)}}$$

- Easily extends to multiple features (x1, x2, x3..)
- And multiple parameter weights

One-vs-all

- Take i:th class (against all other grouped into an alternative class),
 create decision boundary and calculate probability
- Choose the class that had the highest probability against all others.



End of Section

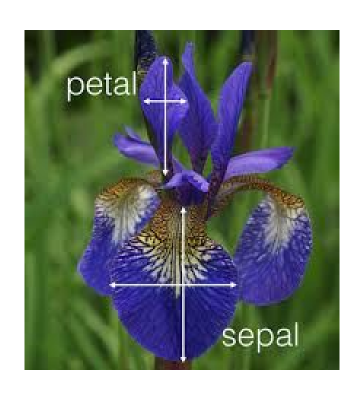
References

- The material presented in this lecture references lecture material draws on the materials the following courses:
- UC Berkeley CS 294-129 (Designing, Visualizing, and Understanding Deep Neural Networks):
 https://bcourses.berkeley.edu/courses/1453965/pages/cs294-129-designing-visualizing-and-understanding-deep-neural-networks
- Stanford CS231n (Convolutional Neural Networks for Visual Recognition): http://cs231n.stanford.edu/
- Stanford CS229 (Machine Learning) & Andrew Ng's Machine Learning at Coursera: http://cs229.stanford.edu/ & https://www.coursera.org/learn/machine-learning

Example Code: Logistic Regression in Scikit-learn

Data

Example Code Sample with Logistic Regression Classifier



Input data

X: Attribute Information:

- sepal length in cm
- sepal width in cm
- petal length in cm
- petal width in cm

Y:

0 = 'setosa',

1 = 'versicolor',

2 = 'virginica'

print type (X) print X[0:5] <type 'numpy.ndarray'> [[5.1 3.5 1.4 0.2] [4.9 3. 1.4 0.2] [4.7 3.2 1.3 0.2] [4.6 3.1 1.5 0.2] [5. 3.6 1.4 0.2]] print Y[0:5] $[0\ 0\ 0\ 0\ 0]$

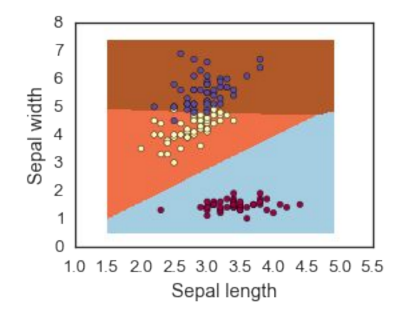
Code Samples with SciKit Learn

```
# 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
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()])
# numpy.ravel: Return a contiguous flattened array.
```

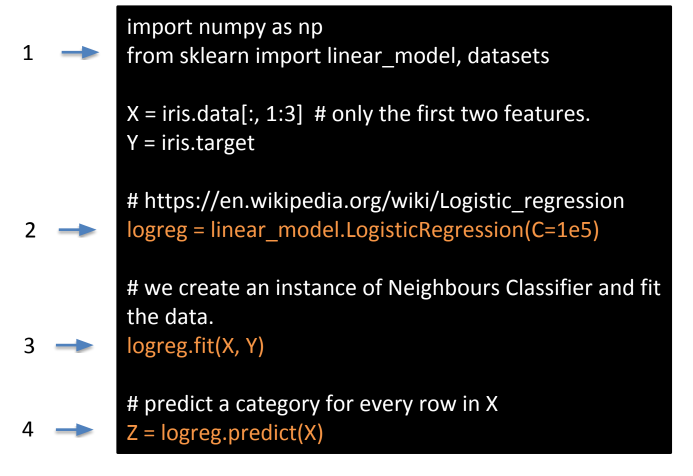
```
xx is a matrix of all the first values
      xx shape is (171, 231)
                                                                                 1.5
      yy shape is (171, 231)
                                                                       3.8
                                                                                                            3.8 ....
                                                                                              3.8
                                                                                 3.8
      np.c returns shape (39501, 2)
                                                                                                            4.0 ....
                                                                                 4.0
                                                                                              4.0
      [[ 3.8 1.5 ]
                                                                                4.2
                                                                                              4.2
                                                                                                            4.2....
      [3.82 1.5]
      [ 3.84 1.5 ] ...]
                                                                        8.4
      Z shape is (39501,)
                                                                             yy is matric of only the second values
                                                                                               yy -> X[:,1]
             1.5
                           yy -> X[:,1]
                                                    4.9
                                                                       3.8
    3.8
                                                                                              1.7
                                                                                                           1.9 ....
                           3.8, 1.7
                                        3.8, 1.9
                                                                   xx-> X[:,0]
             3.8, 1.5
                                                                                                           1.9 ....
                                                                                              1.7
xx -> X[:,0]
                                                                                                           1.9....
                                                                                              1.7
                                                                                1.5
                           4.0, 1.7
                                        4.0, 1.9....
             4.0, 1.5
             4.2, 1.5
                           4.2, 1.7
                                        4.2, 1.9....
                                                                        8.4
     8.4
```

Plotting the Results

```
# Put the result into a color plot
Z = Z.reshape(xx.shape)
plt.figure(1, figsize=(4, 3))
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=get_cmap("Spectral"))
plt.xlabel('Sepal length')
plt.ylabel('Sepal width')
#plt.xlim(xx.min(), xx.max())
#plt.ylim(yy.min(), yy.max())
#plt.xticks(())
#plt.yticks(())
plt.show()
```



Example Code Sample with Logistic Regression Classifier



Class sklearn.linear_model. LogisticRegression

(penalty='l2', dual=False, tol=0.0001, C=1.0, fit_intercept=True, intercept_scaling=1, class_weight=None, random_state=None, solver='liblinear', max_iter=100, multi_class='ovr', verbose=0, warm_start=False, n_jobs=1)

http://scikit-learn.org/stable/modules/generatedsklearn.linear_model.LogisticRegression.html

^{*} Z[2] will be the predicted number for row X[2]

Methods for LogisticRegression

Methods

decision_function(X)	Predict confidence scores for samples.
densify()	Convert coefficient matrix to dense array format.
<pre>fit(X, y[, sample_weight])</pre>	Fit the model according to the given training data.
${\tt fit_transform}(X[,y])$	Fit to data, then transform it.
<pre>get_params([deep])</pre>	Get parameters for this estimator.
predict(X)	Predict class labels for samples in X.
<pre>predict_log_proba(X)</pre>	Log of probability estimates.
predict_proba(X)	Probability estimates.
<pre>score(X, y[, sample_weight])</pre>	Returns the mean accuracy on the given test data and labels.
<pre>set_params(**params)</pre>	Set the parameters of this estimator.
sparsify()	Convert coefficient matrix to sparse format.
<pre>transform(*args, **kwargs)</pre>	DEPRECATED: Support to use estimators as feature selectors will be removed in version 0.19.



fit(X, y, sample_weight=None)

[source]

Fit the model according to the given training data.

Parameters: X: {array-like, sparse matrix}, shape (n_samples, n_features)

Training vector, where n_samples is the number of samples and n_features is the number of features.

y: array-like, shape (n_samples,)

Target vector relative to X.

sample_weight : array-like, shape (n_samples,) optional

Array of weights that are assigned to individual samples. If not provided, then each sample is given unit weight.

New in version 0.17: sample_weight support to LogisticRegression.

Returns:

self : object

Returns self.

predict(X)

[source]

Predict class labels for samples in X.

Parameters: X: {array-like, sparse matrix}, shape = [n_samples, n_features]

Samples.

Returns:

C: array, shape = [n_samples]

Predicted class label per sample.

Fit and predict from ScikitLearn

Regularization

Why: To avoid over-fitting

How: You penalize your loss function by adding a multiple of an L1 (LASSO) or an L2 (Ridge) norm of your weights vector w

Your new loss function = $L(X,Y) + \lambda N(w)$

Tuning the regularization term \lambda: Cross-validation:

- divide your training data,
- train your model for a fixed value of λ and test it on the remaining subsets
- repeat this procedure while varying λ . Then you select the best λ that minimizes your loss function.



Shrinkage Methods II: An example

