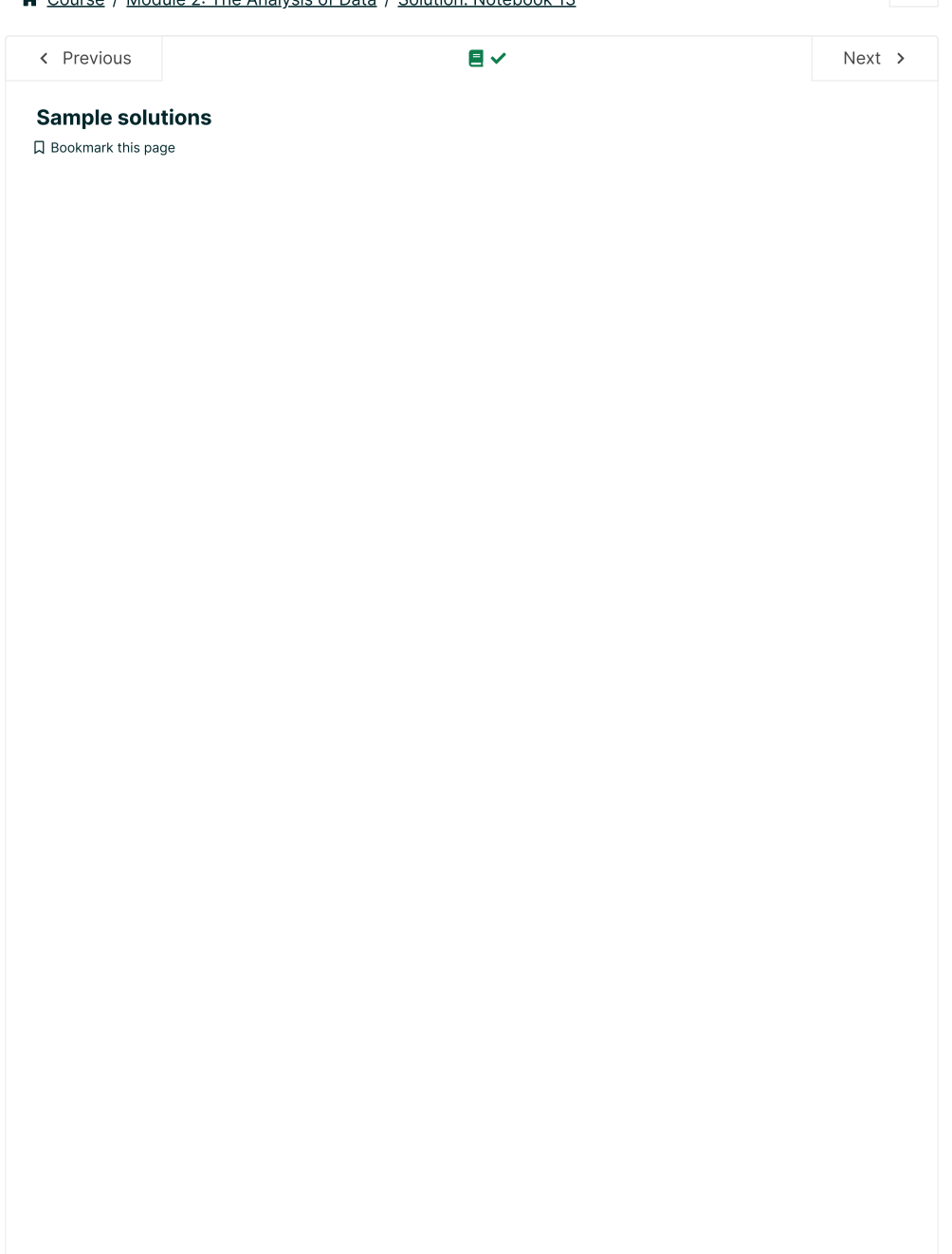
<u>Help</u>

mrajagopal6 v

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# **Logistic regression**

Beyond regression, another important data analysis task is classification, in which you are given a set of labeled data points and you wish to lea the labels. The canonical example of a classification algorithm is *logistic regression*, the topic of this notebook.

Although it's called "regression" it is really a model for classification.

Here, you'll consider binary classification. Each data point belongs to one of c=2 possible classes. By convention, we will denote these class and "1." However, the ideas can be generalized to the multiclass case, i.e., c > 2, with labels  $\{0, 1, \dots, c - 1\}$ .

You'll also want to review from earlier notebooks the concept of gradient ascent/descent (or "steepest ascent/descent"), when optimizing a sca a vector variable.

## **Part 0: Introduction**

This part of the notebook introduces you to the classification problem through a "geometric interpretation."

#### Setup

```
In [1]: import pandas as pd
        import seaborn as sns
        import matplotlib.pyplot as plt
        import numpy as np
        from IPython.display import display, Math
        %matplotlib inline
        import matplotlib as mpl
        mpl.rc("savefig", dpi=100) # Adjust for higher-resolution figures
```

A note about slicing columns from a Numpy matrix. If you want to extract a column i from a Numpy matrix A and keep it as a column vecto use the slicing notation, A[:, i:i+1]. Not doing so can lead to subtle bugs. To see why, compare the following slices.

```
In [2]: A = np.array ([[1, 2, 3],
                        [4, 5, 6],
                        [7, 8, 9]
                       ], dtype=float)
        print ("A[:, :] ==\n", A)
        print ("\na0 := A[:, 0] ==\n", A[:, 0])
        print ("\na1 := A[:, 2:3] == \n", A[:, 2:3])
        print ("\nAdd columns 0 and 2?")
        a0 = A[:, 0]
        a1 = A[:, 2:3]
        print (a0 + a1)
        A[:, :] ==
         [[1. 2. 3.]
         [4. 5. 6.]
         [7. 8. 9.]]
        a0 := A[:, 0] ==
         [1. 4. 7.]
        a1 := A[:, 2:3] ==
         [[3.]
         [6.]
         [9.]]
        Add columns 0 and 2?
        [[ 4. 7. 10.]
         [ 7. 10. 13.]
         [10. 13. 16.]]
```

Aside: Broadcasting in Numpy. What is happening in the operation, a0 + a1, shown above? When the shapes of two objects do not match, to figure out if there is a natural way to make them compatible. See this supplemental notebook (./mo\_numpy\_mo\_problems.ipynb) for information Numpy's "broadcasting rule," along with other Numpy tips.

#### Example data: Rock lobsters!

As a concrete example of a classification task, consider the results of the following experiment (http://www.stat.ufl.edu/~winner/data/lobster\_si

Some marine biologists started with a bunch of lobsters of varying sizes (size being a proxy for the stage of a lobster's development). They then exposed these lobsters to a variety of predators. Finally, the outcome that they measured is whether the lobsters survived or not.

The data is a set of points, one point per lobster, where there is a single predictor (the lobster's size) and the response is whether the lobsters s "1") or died (label "0").

For the original paper, see this link (http://downeastinstitute.org/assets/files/Published%20papers/Wilkinson%20et%20al%202015-1.p For what we can only guess is what marine biologists do in their labs, see this image (http://i.imgur.com/dQDKgys.jpg) (or this possibly safe-for-work alternative (http://web.archive.org/web/20120628012654/http://www.traemcneely.com/wp-content/uploads/2012/04/wpi Lobster-Fights-e1335308484734.jpeg)).

Start by downloading this data.

```
In [3]: import requests
        import os
        import hashlib
        import io
        def on_vocareum():
            return os.path.exists('.voc')
        def download(file, local_dir="", url_base=None, checksum=None):
            local_file = "{}{}".format(local_dir, file)
            if not os.path.exists(local_file):
                if url base is None:
                    url_base = "https://cse6040.gatech.edu/datasets/"
                url = "{}{}".format(url_base, file)
                print("Downloading: {} ...".format(url))
                r = requests.get(url)
                with open(local_file, 'wb') as f:
                    f.write(r.content)
            if checksum is not None:
                with io.open(local_file, 'rb') as f:
                    body = f.read()
                    body checksum = hashlib.md5(body).hexdigest()
                    assert body checksum == checksum, \
                         "Downloaded file '{}' has incorrect checksum: '{}' instead of '{}'".format(local
                                                                                                     checl
            print("'{}' is ready!".format(file))
        if on vocareum():
            URL BASE = "https://cse6040.gatech.edu/datasets/rock-lobster/"
            DATA_PATH = "../resource/asnlib/publicdata/"
        else:
            URL BASE = "https://github.com/cse6040/labs-fa17/raw/master/datasets/rock-lobster/"
            DATA PATH = ""
        datasets = {'lobster_survive.dat.txt': '12fc1c22ed9b4d7bf04bf7e0fec996b7',
                     logreg points train.csv': '25bbca6105bae047ac4d62ee8b76c841',
                     'log likelihood soln.npz': '5a9e17d56937855727afa6db1cd83306',
                     'grad_log_likelihood_soln.npz': 'a67c00bfa95929e12d423105d8412026',
                     'hess_log_likelihood_soln.npz': 'b46443fbf0577423b084122503125887'}
        for filename, checksum in datasets.items():
            download(filename, local dir=DATA PATH, url base=URL BASE, checksum=checksum)
        print("\n(All data appears to be ready.)")
        'grad_log_likelihood_soln.npz' is ready!
        'hess log likelihood soln.npz' is ready!
        'log_likelihood_soln.npz' is ready!
        'lobster_survive.dat.txt' is ready!
        'logreg_points_train.csv' is ready!
        (All data appears to be ready.)
```

Here is a plot of the raw data, which was taken from this source (http://www.stat.ufl.edu/~winner/data/lobster\_survive.dat).

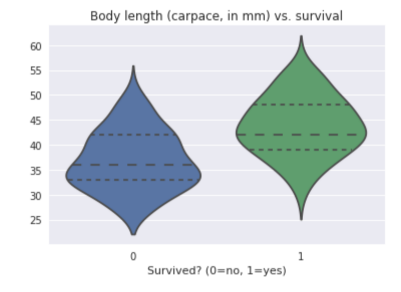
```
In [4]: df lobsters = pd.read table('{}lobster survive.dat.txt'.format(DATA PATH),
                                    sep=r'\s+', names=['CarapaceLen', 'Survived'])
```

```
display(df_lobsters.head())
print("...")
display(df_lobsters.tail())
```

		_
	CarapaceLen	Survived
0	27	0
1	27	0
2	27	0
3	27	0
4	27	0

	CarapaceLen	Survived
154	54	1
155	54	1
156	54	1
157	54	1
158	57	1

```
In [5]: ax = sns.violinplot(x="Survived", y="CarapaceLen",
                            data=df_lobsters, inner="quart")
        ax.set(xlabel="Survived? (0=no, 1=yes)",
               ylabel="",
               title="Body length (carpace, in mm) vs. survival");
```



Although the classes are distinct in the aggregate, where the median carapace (outer shell) length is around 36 mm for the lobsters that died ar those that survived, they are not cleanly separable.

#### **Notation**

To develop some intuition and a classification algorithm, let's formulate the general problem and apply it to synthetic data sets.

Let the data consist of m observations of d continuously-valued predictors. In addition, for each data observation we observe a binary label wh either 0 or 1.

Just like our convention in the linear regression case, represent each observation, or data point, by an augumented vector,  $\hat{x}_i^T$ ,

$$\hat{x}_i^T \equiv (x_{i,0} \quad x_{i,1} \quad \cdots \quad x_{i,d-1} \quad 1).$$

That is, the point is the d coordinates augmented by an initial dummy coordinate whose value is 1. This convention is similar to what we did in regression.

We can also stack these points as rows of a matrix, X, again, just as we did in regression:

$$X \equiv \begin{pmatrix} \hat{x}_0^T \\ \hat{x}_1^T \\ \vdots \\ \hat{x}_{m-1}^T \end{pmatrix} = \begin{pmatrix} x_{0,1} & x_{0,2} & \cdots & x_{0,d} & 1 \\ x_{1,1} & x_{1,2} & \cdots & x_{1,d} & 1 \\ \vdots & \vdots & & & 1 \\ x_{m-1,1} & x_{m-1,2} & \cdots & x_{m-1,d} & 1 \end{pmatrix}.$$

We will take the labels to be a binary vector,  $\mathbf{y}^T \equiv \left(y_0, y_1, \dots, y_{m-1}\right)^T$ .

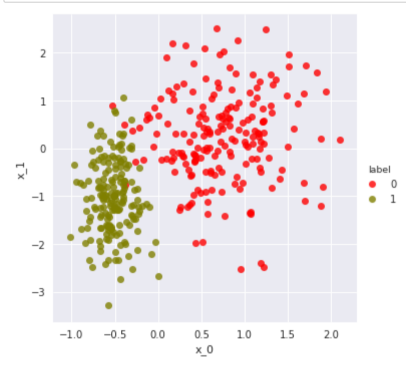
Example: A synthetic training set. We've pre-generated a synethetic data set consisting of labeled data points. Let's download and inspect it table and then visually.

```
In [6]: | df = pd.read_csv('{}logreg_points_train.csv'.format(DATA_PATH))
        display(df.head())
        print("...")
        display(df.tail())
```

	x_0	x_1	label
0	-0.234443	-1.075960	1
1	0.730359	-0.918093	0
2	1.432270	-0.439449	0
3	0.026733	1.050300	0
4	1.879650	0.207743	0

x_0	x_1	label
1.314300	0.746001	0
-0.759737	-0.042944	1
0.683560	-0.047791	0
0.774747	0.743837	0
0.899119	1.576390	0
	1.314300 -0.759737 0.683560 0.774747	1.314300 0.746001 -0.759737 -0.042944 0.683560 -0.047791 0.774747 0.743837

```
In [7]: def make_scatter_plot(df, x="x_0", y="x_1", hue="label",
                              palette={0: "red", 1: "olive"},
                              size=5):
            sns.lmplot(x=x, y=y, hue=hue, data=df, palette=palette,
                       fit_reg=False)
        mpl.rc("savefig", dpi=120) # Adjust for higher-resolution figures
        make scatter plot(df)
```



Next, let's extract the coordinates as a Numpy matrix of points and the labels as a Numpy column vector labels. Mathematically, the point corresponds to X and the labels vector corresponds to y.

```
In [8]: points = np.insert(df.as_matrix (['x_0', 'x_1']), 2, 1.0, axis=1)
       labels = df.as_matrix(['label'])
       print ("First and last 5 points:\n", '='*23, '\n', points[:5], '\n', points[-5:], '\n')
       print ("First and last 5 labels:\n", '='*23, '\n', labels[:5], '\n...\n', labels[-5:], '\n')
       First and last 5 points:
        _____
        [[-0.234443 -1.07596
                             1.
                                      ]
        [ 0.730359 -0.918093 1.
        [ 1.43227 -0.439449
                             1.
                                     ]
        [ 0.0267327 1.0503
                             1.
                                     ]
        1.
                                     ]]
        [[ 1.3143
                    0.746001
                             1.
                                      ]
        [-0.759737 -0.0429435]
                                     ]
        [ 0.68356
                  -0.0477909
                            1.
                                     ]
                   0.743837
        [ 0.774747
                             1.
                                     ]
        [ 0.899119
                   1.57639
                                     ]]
       First and last 5 labels:
```

[[1] [0] [0] [0] [0]] [[0]] [1] [0] [0] [0]]

## Linear discriminants and the heaviside function

Suppose you think that the boundary between the two clusters may be represented by a line. For the synthetic data example above, I hope you such a model is not a terrible one.

A linear boundary is also known as a *linear discriminant*. Any point x on this line may be described by  $\theta^T x$ , where  $\theta$  is a vector of coefficients:

$$\theta \equiv \begin{pmatrix} \theta_0 \\ \theta_1 \\ \vdots \\ \theta_d \end{pmatrix}.$$

For example, suppose our observations have two predictors each (d=2). Let the corresponding data point be  $x^T \equiv (x_0, x_1, x_2 = 1.0)$ . Then, means that

$$\theta^{T}x = 0 = \theta_{0}x_{0} + \theta_{1}x_{1} + \theta_{2}$$

$$\Rightarrow x_{1} = -\frac{\theta_{2}}{\theta_{1}} - \frac{\theta_{0}}{\theta_{1}}x_{0}.$$

So that describes points on the line. However, given any point x in the d-dimensional space that is not on the line,  $\theta^T x$  still produces a value: the be positive on one side of the line ( $\theta^T x > 0$ ) or negative on the other ( $\theta^T x < 0$ ).

In other words, you can use the linear discriminant function,  $\theta^T x$ , to generate a label for each point x: just reinterpret its sign!

If you want "0" and "1" labels, the heaviside function, H(y), will convert a positive y to the label "1" and all other values to "0."

$$H(y) \equiv \begin{cases} 1 & \text{if } y > 0 \\ 0 & \text{if } y \le 0 \end{cases}.$$

**Exercise 0** (2 points). Given the a  $m \times (d+1)$  matrix of augmented points (i.e., the X matrix) and a column vector  $\theta$  of length d+1, implement compute the value of the linear discriminant at each point. That is, the function should return a (column) vector y where the  $y_i = \theta^T x_i$ .

```
In [9]: def lin discr (X, theta):
               ### BEGIN SOLUTION
              return X.dot(theta)
               ### END SOLUTION
In [10]: # Test cell: `lin_discr__check`
          import random
          theta_test = [random.random() for _ in range (3)]
          x0_test = [random.random() for _ in range (2)]
          x1_test = [(-theta_test[2] - theta_test[0]*x0) / theta_test[1] for x0 in x0_test]
          X_{\text{test}} = \text{np.array} ([[x0*2 \text{ for } x0 \text{ in } x0_{\text{test}}] + [x0*0.5 \text{ for } x0 \text{ in } x0_{\text{test}}],
                                x1 test + x1 test,
                                [1.0, 1.0, 1.0, 1.0],]).T
          print(X_test, "\n")
          LD_test = lin_discr(X_test, np.array([theta_test]).T)
          print (LD test)
          assert (LD_test[:2] > 0).all ()
           assert (LD_test[2:] < 0).all ()
          print("\n(Passed.)")
          [[ 1.05035323 -0.52024135 1.
           [ 1.49196858 -0.59315241 1.
                                                   ]
           [ 0.26258831 -0.52024135 1.
                                                   ]
           [ 0.37299215 -0.59315241 1.
                                                   ]]
          [[ 0.13320893]
           [ 0.1892159 ]
           [-0.06660446]
           [-0.09460795]]
          (Passed.)
```

**Exercise 1** (2 points). Implement the *heaviside function*, H(y). Your function should allow for an arbitrary *matrix* of input values and should app heaviside function to each element. In the returned matrix, the elements should have a floating-point type.

Example, the code snippet

There are several possible approaches that lead to one-line solutions. One uses only logical and arithmetic operators, which you will re implemented as elementwise operations for Numpy arrays. Another uses Numpy's <a href="mailto:sign()">sign()</a>
(<a href="http://docs.scipy.org/doc/numpy/reference/generated/numpy.sign.html">http://docs.scipy.org/doc/numpy/reference/generated/numpy.sign.html</a>) function.

```
In [11]: def heaviside(Y):
              ### BEGIN SOLUTION
              return 1.0*(Y > 0.0)
              # Alternative solution:
              \#return (np.sign(Y) > 0) * 1.0
              ### END SOLUTION
In [12]: # Test cell: `heaviside check`
          Y_{\text{test}} = \text{np.array}([[-2.3, 1.2, 7.],
                              [0.0, -np.inf, np.inf]])
         H_Y_test = heaviside(Y_test)
         print("Y:\n", Y_test)
         print("\nH(Y):\n", H_Y_test)
          assert (H_Y_test.astype(int) == np.array([[0, 1, 1], [0, 0, 1]])).all ()
         print ("\n(Passed.)")
         Y:
          [[-2.3 \ 1.2 \ 7.]
          [ 0. -inf inf]]
         H(Y):
          [[0. 1. 1.]
          [0. 0. 1.]]
          (Passed.)
```

For the next exercise, we'll need the following functions.

```
In [13]: def heaviside int(Y):
              """Evaluates the heaviside function, but returns integer values."""
             return heaviside(Y).astype(dtype=int)
         def gen_lin_discr_labels(points, theta, fun=heaviside_int):
             Given a set of points and the coefficients of a linear
              discriminant, this function returns a set of labels for
              the points with respect to this discriminant.
             score = lin_discr(points, theta)
             labels = fun(score)
             return labels
         def plot lin discr(theta, df, x="x 0", y="x 1", hue="label",
                             palette={0: "red", 1: "olive"}, size=5,
                             linewidth=2):
             lm = sns.lmplot(x=x, y=y, hue=hue, data=df, palette=palette,
                              size=size, fit_reg=False)
             x \min, x \max = df[x].min(), df[x].max()
             y \min, y \max = df[y].min(), df[y].max()
             x1_{min} = (-theta[2][0] - theta[0][0]*x_{min}) / theta[1][0]
             x1_{max} = (-theta[2][0] - theta[0][0]*x_{max}) / theta[1][0]
             plt.plot([x_min, x_max], [x1_min, x1_max], linewidth=linewidth)
             def expand interval(x limits, percent=10.0):
                  x_min, x_max = x_limits[0], x_limits[1]
                  if x min < 0:
                     x_min *= 1.0 + 1e-2*percent
                  else:
                      x min *= 1.0 - 1e-2*percent
                  if x max > 0:
                     x_max *= 1.0 + 1e-2*percent
                  else:
                      x max *= 1.0 + 1e-2*percent
                  return (x min. x max)
```

```
LCCULM (A_MIM, A_MGA
    x_view = expand_interval((x_min, x_max))
    y_view = expand_interval((y_min, y_max))
    lm.axes[0,0].set_xlim(x_view[0], x_view[1])
    lm.axes[0,0].set_ylim(y_view[0], y_view[1])
def mark_matches(a, b, exact=False):
    Given two Numpy arrays of {0, 1} labels, returns a new boolean
    array indicating at which locations the input arrays have the
    same label (i.e., the corresponding entry is True).
    This function can consider "inexact" matches. That is, if `exact`
    is False, then the function will assume the {0, 1} labels may be
    regarded as the same up to a swapping of the labels. This feature
    allows
      a == [0, 0, 1, 1, 0, 1, 1]
     b == [1, 1, 0, 0, 1, 0, 0]
    to be regarded as equal. (That is, use `exact=False` when you
    only care about "relative" labeling.)
    assert a.shape == b.shape
    a_int = a.astype(dtype=int)
    b_int = b.astype(dtype=int)
    all_axes = tuple(range(len(a.shape)))
    assert ((a_int == 0) | (a_int == 1)).all()
    assert ((b_int == 0) | (b_int == 1)).all()
    exact_matches = (a_int == b_int)
    if exact:
        return exact matches
    assert exact == False
    num exact matches = np.sum(exact matches)
    if (2*num exact matches) >= np.prod(a.shape):
        return exact matches
    return exact_matches == False # Invert
def count matches(a, b, exact=False):
    Given two sets of {0, 1} labels, returns the number of mismatches.
    This function can consider "inexact" matches. That is, if `exact'
    is False, then the function will assume the {0, 1} labels may be
    regarded as similar up to a swapping of the labels. This feature
    allows
      a == [0, 0, 1, 1, 0, 1, 1]
      b == [1, 1, 0, 0, 1, 0, 0]
    to be regarded as equal. (That is, use `exact=False` when you
    only care about "relative" labeling.)
   matches = mark_matches(a, b, exact=exact)
    return int(matches.sum())
```

**Exercise 2** (2 points). For the synthetic data you loaded above, try by hand to find a value for  $\theta$  such that  $H(\theta^T x)$  "best" separates the two clu this  $\theta$  in a variable named my\_theta, which should be a Numpy column vector. That is, define my\_theta here using a line like:

```
my\_theta = np\_col\_vec([3., 0., -1.])
```

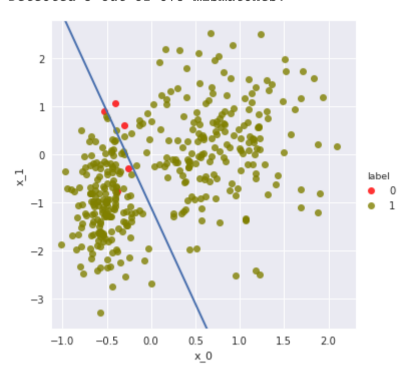
where np col vec is defined below and the list of values are your best guesses at discriminating coefficients. The test code will check that your best guesses at discriminating coefficients. makes no more than ten misclassifications.

Hint: We found a set of coefficients that commits just 5 errors for the 375 input points

```
In [14]: def np col vec (list values):
              """Returns a Numpy column vector for the given list of scalar values."""
             return np.array ([list_values]).T
         # Redefine `my_theta` as instructed above to reduce the number of mismatches:
         my_theta = np_col_vec([-1., 3., 0.]) # 123 mismatches
         ### BEGIN SOLUTION
         my_{theta} = np_{col_{vec}([-6.5, -1., -1.35])} # 5 mismatches
         my_{theta} = np_{col_{vec}([-2., -0.5, -0.55])} # 5 mismatches
         ### END SOLUTION
In [15]: # Here are the labels generated by your discriminant:
         my_labels = gen_lin_discr_labels(points, my_theta)
         # Here is a visual check:
         num mismatches = len(labels) - count matches(labels, my labels)
         print ("Detected", num mismatches, "out of", len(labels), "mismatches.")
```

```
df matches = df.copy ()
df_matches['label'] = mark_matches (my_labels, labels).astype (dtype=int)
mpl.rc("savefig", dpi=100) # Adjust for higher-resolution figures
plot_lin_discr (my_theta, df_matches)
assert num mismatches <= 10</pre>
```

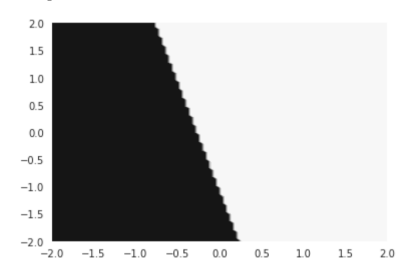
Detected 5 out of 375 mismatches.



How the heaviside divides the space. The heaviside function,  $H(\theta^T x)$ , enforces a sharp boundary between classes around the  $\theta^T x = 0$  line code produces a contour plot (https://matplotlib.org/api/ as gen/matplotlib.axes.Axes.contourf.html) to show this effect: there will be a sharp of between 0 and 1 values, with one set of values shown as a solid dark area and the remaining as a solid light-colored area.

```
In [16]: x0 = np.linspace(-2., +2., 100)
         x1 = np.linspace(-2., +2., 100)
         x0_grid, x1_grid = np.meshgrid(x0, x1)
         h_grid = heaviside(my_theta[2] + my_theta[0]*x0_grid + my_theta[1]*x1_grid)
         plt.contourf(x0, x1, h grid)
```

Out[16]: <matplotlib.contour.QuadContourSet at 0x7f0ec25ad978>



# Part 1: The logistic (or sigmoid) function as an alternative discriminant

As the lobsters example suggests, real data are not likely to be cleanly separable, especially when the number of features we have at our dispo small.

Since the labels are 0 or 1, you could look for a way to interpret labels as probabilities rather than as hard (0 or 1) labels. One such function is the function, also referred to as the logit or sigmoid (https://en.wikipedia.org/wiki/Sigmoid function) function.

$$G(y) \equiv \frac{1}{1 + e^{-y}}$$

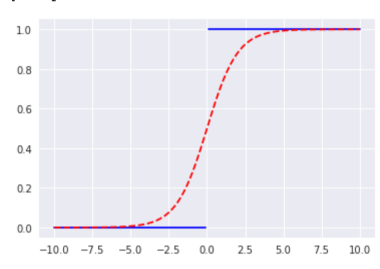
The logistic function takes any value in the range  $(-\infty, +\infty)$  and produces a value in the range (0, 1). Thus, given a value y, we can interpret conditional probability that the label is 1 given y, i.e.,  $G(y) \equiv \Pr[\text{label is } 1 \mid y]$ .

**Exercise 3** (2 points). Implement the logistic function. Inspect the resulting plot of G(y) in 1-D and then the contour plot of  $G(\theta^T x)$ . Your function accept a Numpy matrix of values, Y, and apply the sigmoid elementwise.

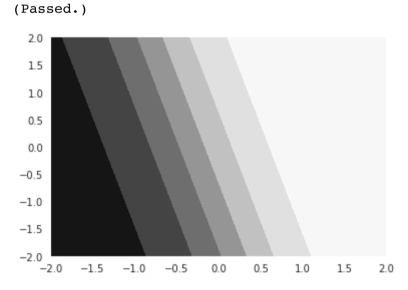
```
In [17]: def logistic(Y):
             ### BEGIN SOLUTION
             return 1.0 / (1.0 + np.exp (-Y))
             ### END SOLUTION
         # Plot your function for a 1-D input.
```

```
y_values = np.linspace(-10, 10, 100)
mpl.rc("savefig", dpi=120) # Adjust for higher-resolution figures
sns.set_style("darkgrid")
y_pos = y_values[y_values > 0]
y_rem = y_values[y_values <= 0]</pre>
plt.plot(y_rem, heaviside (y_rem), 'b')
plt.plot(y_pos, heaviside (y_pos), 'b')
plt.plot(y_values, logistic (y_values), 'r--')
#sns.regplot (y values, heaviside (y values), fit reg=False)
#sns.regplot (y_values, logistic (y_values), fit_reg=False)
```

Out[17]: [<matplotlib.lines.Line2D at 0x7f0ec257c2b0>]



```
In [18]: # Test cell: `logistic__check`
         assert logistic(np.log(3)) == 0.75
         assert logistic(-np.log(3)) == 0.25
         g_grid = logistic(my_theta[2] + my_theta[0]*x0_grid + my_theta[1]*x1_grid)
         plt.contourf (x0, x1, g grid)
         assert ((np.round(g_grid) - h_grid).astype(int) == 0).all()
         print ("\n(Passed.)")
```



Exercise 4 (optional; ungraded). Consider a set of 1-D points generated by a mixture of Gaussians. That is, suppose that there are two Gaussians. over the 1-dimensional variable,  $x \in (-\infty, +\infty)$ , that have the same variance ( $\sigma^2$ ) but different means ( $\mu_0$  and  $\mu_1$ ). Show that the conditional observing a point labeled "1" given x may be written as,

$$\Pr[l = 1 \mid x] \propto \frac{1}{1 + e^{-(\theta_0 x + \theta_1)}},$$

for a suitable definition of  $\theta_0$  and  $\theta_1$ .

Hints. Since the points come from Gaussian distributions,

$$\Pr[x \mid l] \equiv \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x-\mu_l)^2}{2\sigma^2}\right).$$

To rewrite  $Pr[l \mid x]$  in terms of  $Pr[x \mid l]$ , recall Bayes's rule (also: Bayes's theorem (https://en.wikipedia.org/wiki/Bayes%27\_theorem)):

$$Pr[l = 1 | x] = \frac{Pr[x | l = 1] Pr[l = 1]}{Pr[x]},$$

where the denominator can be expanded as

$$Pr[x] = Pr[x | l = 0] Pr[l = 0] + Pr[x | l = 1] Pr[l = 1].$$

You may assume the prior probabilities of observing a 0 or 1 are given by  $\Pr[l=0] \equiv p_0$  and  $\Pr[l=1] \equiv p_1$ .

The point of this derivation is to show you that the definition of the logistic function does not just arise out of thin air. It also hints that you might expect a final algorithm for logistic regression based on using G(y) as the discriminant will work well when the classes are best explained as a mixture of Gaussians.

**Generalizing to** d-dimensions. The preceding exercise can be generalized to d-dimensions. Let  $\theta$  and x be (d+1)-dimensional points. Then

$$\Pr\left[l = 1 \mid x\right] \propto \frac{1}{1 + \exp\left(-\theta^T x\right)}.$$

**Exercise 5** (*optional*; ungraded). Verify the following properties of the logistic function, G(y).

$$G(y) = \frac{e^y}{e^y + 1} \tag{P1}$$

$$G(-y) = 1 - G(y)$$
 (P2)

$$\frac{dG}{dy} = G(y)G(-y) \quad (P3)$$

$$\frac{d}{dy}[\ln G(y)] = G(-y)$$
 (P4)

$$\frac{d}{dy}\ln[1 - G(y)] = -G(y)$$
 (P5)

**Answers.** In all of the derivations below, we use the fact that G(y) > 0.

(P1). Multiply the numerator and denominator by  $e^{y}$ .

(P2). Start with the right-hand side, 1 - G(y), apply some algebra, and then apply (P1)

$$1 - G(y) = \frac{e^{y} + 1}{e^{y} + 1} - \frac{e^{y}}{e^{y} + 1} = \frac{1}{e^{y} + 1} \cdot \frac{e^{-y}}{e^{-y}} = \frac{e^{-y}}{e^{-y} + 1} = G(-y).$$

(P3). By direct calculation and application of (P1):

$$\frac{dG}{dy} = \frac{d}{dy} (1 + e^{-y})^{-1} = -(1 + e^{-y})^{-2} \cdot (-e^{-y}) = \underbrace{\frac{1}{1 + e^{-y}}}_{=G(y)} \cdot \underbrace{\frac{e^{-y}}{1 + e^{-y}}}_{=G(-y)} = G(y) \cdot G(-y).$$

(P4). By the chain rule and application of (P3):

$$\frac{d}{dy}\ln G(y) = \left(\frac{d}{dG}\ln G\right)\frac{dG}{dy} = \frac{1}{G(y)}\cdot G(y)G(-y) = G(-y).$$

(P5). By combining (P2), variable substitution and the chain rule, and (P4),

$$\frac{d}{dy}\ln[1 - G(y)] = \frac{d}{dy}\ln G(-y) = \underbrace{\left[\frac{d}{dz}\ln G(z)\right] \cdot \frac{dz}{dy}}_{\text{Let }z \equiv -y} = G(-z) \cdot (-1) = -G(y).$$

#### Part 2: Determining the discriminant via maximum likelihood estimation

Previously, you determined  $\theta$  for our synthetic dataset by hand. Can you compute a good  $\theta$  automatically? One of the standard techniques in s perform a *maximum likelihood estimation* (MLE) of a model's parameters,  $\theta$ . Indeed, you may have seen or used MLE to derive the normal equal regression in a more "statistically principled" way.

"Likelihood" as an objective function. MLE derives from the following idea. Consider the joint probability of observing all of the labels, given the parameters, *θ*:

$$\Pr[y \mid X, \theta].$$

Suppose these observations are independent and identically distributed (i.i.d.). Then the joint probability can be factored as the product of indiversal probabilities,

$$\begin{split} \Pr[y \mid X, \theta] &= \Pr[y_0, \dots, y_{m-1} \mid \hat{x}_0, \dots, \hat{x}_{m-1}, \theta] = \Pr[y_0 \mid \hat{x}_0, \theta] \cdots \Pr[y_{m-1} \mid \hat{x}_{m-1}, \theta] \\ &= \prod_{i=0}^{m-1} \Pr[y_i \mid \hat{x}_i, \theta]. \end{split}$$

The maximum likelihood principle says that you should choose  $\theta$  to maximize the chances (or "likelihood") of seeing these particular observation  $\Pr[y \mid X, \theta]$  is now an objective function to maximize.

For both mathematical and numerical reasons, we will use the logarithm of the likelihood, or log-likelihood, as the objective function instead. Le

the logarithm of the likelihood, or log-like 
$$\mathcal{L}(\theta; y, X) \equiv \log \left\{ \prod_{i=0}^{m-1} \Pr[y_i \mid \hat{x}_i, \theta] \right\}$$
$$= \sum_{i=0}^{m-1} \log \Pr[y_i \mid \hat{x}_i, \theta].$$

We are using the symbol log, which could be taken in any convenient base, such as the natural logarithm ( $log_2$  y).

The MLE fitting procedure then consists of two steps:

- For the problem at hand, decide on a model of  $\Pr[y_i \mid \hat{x}_i, \theta]$ .
- Run any optimization procedure to find the  $\theta$  that maximizes  $\mathcal{L}(\theta; y, X)$ .

# Part 3: MLE for logistic regression

Let's say you have decided that the logistic function,  $G(\hat{x}_i^T \theta) = G(\theta^T \hat{x}_i)$ , is a good model of the probability of producing a label  $y_i$  given the o Under the i.i.d. assumption, you can interpret the label  $y_i$  as the result of flipping a coin, or a Bernoulli trial (https://en.wikipedia.org/wiki/Bernou the probability of success  $(y_i = 1)$  is defined as  $g_i = g_i(\theta) \equiv G(\hat{x}_i^T \theta)$ . Thus,

$$\Pr[y_i \mid \hat{x}_i, \theta] \equiv g_i^{y_i} \cdot (1 - g_i)^{1 - y_i}.$$

The log-likelihood in turn becomes,

$$\mathcal{L}(\theta; y, X) = \sum_{i=0}^{m-1} y_i \ln g_i + (1 - y_i) \ln(1 - g_i)$$

$$= \sum_{i=0}^{m-1} y_i \ln \frac{g_i}{1 - g_i} + \ln(1 - g_i)$$

$$= \sum_{i=0}^{m-1} y_i \theta^T \hat{x}_i + \ln(1 - g_i).$$

You can write the log-likelihood more compactly in the language of linear algebra.

**Convention 1.** Let  $u \equiv (1, ..., 1)^T$  be a column vector of all ones, with its length inferred from context. Let  $A = (a_0 \quad a_1 \quad \cdots \quad a_{n-1})$  be a where  $\{a_i\}$  denote its *n* columns. Then, the sum of the columns is

$$\sum_{i=0}^{n-1} a_i = \begin{pmatrix} a_0 & a_1 & \cdots & a_{n-1} \end{pmatrix} \cdot \begin{pmatrix} 1 \\ 1 \\ \vdots \\ 1 \end{pmatrix} = Au.$$

**Convention 2.** Let  $A = (a_{ii})$  be any matrix and let f(y) be any function that we have defined by default to accept a scalar argument y and pro result. For instance,  $f(y) = \ln y$  or f(y) = G(y). Then, assume that B = f(A) applies  $f(\cdot)$  elementwise to A, returning a matrix B whose eleme  $b_{ii} = f(a_{ii}).$ 

With these notational conventions, convince yourself that these are two different ways to write the log-likelihood for logistic regression.

(V1) 
$$\mathcal{L}(\theta; y, X) = y^T \ln G(X\theta) + (u - y)^T \ln[u - G(X\theta)]$$
  
(V2)  $\mathcal{L}(\theta; y, X) = y^T X\theta + u^T \ln G(-X\theta)$ 

Exercise 6 (2 points). Implement the log-likelihood function in Python by defining a function with the following signature:

```
def log_likelihood (theta, y, X):
```

To compute the elementwise logarithm of a matrix or vector, use Numpy's <u>log</u> (https://docs.scipy.org/doc/numpy/reference/generated/numpy.log.html) function.

```
In [19]: def log_likelihood(theta, y, X):
         ### BEGIN SOLUTION
             u = np.ones((len (X), 1)) # column of all ones
             z = X.dot(theta)
             return y.T.dot(z) + u.T.dot(np.log(logistic(-z)))
         def log_likelihood_alt(theta, y, X):
             z = X.dot(theta)
             g = logistic(z)
             return y.T.dot(np.log(g)) + (1.0-y).T.dot(np.log(1.0-g))
         ### END SOLUTION
```

```
In [20]: # Test cell: `log likelihood check`
         if False:
             d soln = 10
             m soln = 1000
             theta_soln = np.random.random ((d_soln+1, 1)) * 2.0 - 1.0
             y_soln = np.random.randint (low=0, high=2, size=(m_soln, 1))
             X_{soln} = np.random.random ((m_{soln}, d_{soln+1})) * 2.0 - 1.0
             X_{soln[:, 0]} = 1.0
             L_soln = log_likelihood (theta_soln, y_soln, X_soln)
             np.savez compressed('log likelihood soln',
                                  d_soln, m_soln, theta_soln, y_soln, X_soln, L_soln)
         npzfile_soln = np.load('{}log_likelihood_soln.npz'.format(DATA_PATH))
         d_soln = npzfile_soln['arr 0']
         m soln = npzfile soln['arr 1']
         theta soln = npzfile soln['arr 2']
         y_soln = npzfile_soln['arr_3']
         X_soln = npzfile_soln['arr_4']
```

```
L_soln = npzfile_soln['arr_5']
L_you = log_likelihood(theta_soln, y_soln, X_soln)
your_err = np.max(np.abs(L_you/L_soln - 1.0))
cal{L}_{\tiny \mbox{solution}}}\right\|_\infty \approx %g' % your_err))
assert your err <= 1e-12
print ("\n(Passed.)")
\left\| \frac{\mathcal{L}_{\text{yours}} - \mathcal{L}_{\text{solution}}}{\mathcal{L}_{\text{solution}}} \right\|_{\infty} \approx 0
```

To optimize the log-likelihood with respect to the parameters,  $\theta$ , you want to "set the derivative to zero" and solve for  $\theta$ .

Part 4: Computing the MLE solution via gradient ascent: theory

For example, recall that in the case of linear regression via least squares minimization, carrying out this process produced an analytic solution for parameters, which was to solve the normal equations.

Unfortunately, for logistic regression---or for most log-likelihoods you are likely to ever write down---you cannot usually derive an analytic solut you will need to resort to numerical optimization procedures.

Gradient ascent, in 1-D. A simple numerical algorithm to maximize a function is gradient ascent (or steepest ascent). If instead you are minimize function, then the equivalent procedure is gradient (or steepest) descent. Here is the basic idea in 1-D.

Suppose we wish to find the maximum of a scalar function f(x) in one dimension. At the maximum,  $\frac{df(x)}{dx} = 0$ .

Suppose instead that  $\frac{df}{dx} \neq 0$  and consider the value of f at a nearby point, x + s, as given approximately by a truncated Taylor series:

$$f(x+s) = f(x) + s\frac{df(x)}{dx} + \mathcal{O}(s^2).$$

To make progress toward maximizing f(x), you'd like to choose s so that f(x+s) > f(x). One way is to choose  $s = \alpha \cdot \text{sign}\left(\frac{df}{dx}\right)$ , where 0"small:

$$f\left(x + \alpha \cdot \operatorname{sign}\left(\frac{df}{dx}\right)\right) \approx f(x) + \alpha \left|\frac{df}{dx}\right| + \mathcal{O}(\alpha^2).$$

If  $\alpha$  is small enough, then you can neglect the  $\mathcal{O}(\alpha^2)$  term and f(x+s) will be larger than f(x), thus making progress toward finding a maximum

This scheme is the basic idea: starting from some initial guess x, refine the guess by taking a small step s in the direction of the derivative, i.e.,

**Gradient ascent in higher dimensions.** Now suppose x is a vector rather than a scalar. Then the value of f at a nearby point f(x + s), where becomes

$$f(x + s) = f(x) + s^{T} \nabla_{x} f(x) + \mathcal{O}(\|s\|^{2}),$$

where  $\nabla_x f(x)$  is the gradient of f with respect to x. As in the 1-D case, you want a step s such that f(x+s) > f(x). To make as much progres let's choose s to be parallel to  $\nabla_x f(x)$ , that is, proportional to the gradient:

$$s \equiv \alpha \frac{\nabla_x f(x)}{\|\nabla_x f(x)\|}.$$

Again,  $\alpha$  is a fudge (or "gentle nudge?") factor. You need to choose it to be small enough that the high-order terms of the Taylor approximation negligible, yet large enough that you can make reasonable progress.

The gradient ascent procedure applied to MLE. Applying gradient ascent to the problem of maximizing the log-likelihood leads to the following

• Start with some initial guess,  $\theta(0)$ .

(Passed.)

- At each iteration  $t \ge 0$  of the procedure, let  $\theta(t)$  be the current guess.
- Compute the direction of steepest ascent by evaluating the gradient,  $\Delta_t \equiv \nabla_{\theta(t)} \{ \mathcal{L}(\theta(t); y, X) \}$ .
- Define the step to be  $s_t \equiv \alpha \frac{\Delta_t}{\|\Delta_t\|}$ , where  $\alpha$  is a suitably chosen fudge factor.
- Take a step in the direction of the gradient,  $\theta(t+1) \leftarrow \theta(t) + s_t$ .
- Stop when the parameters don't change much or after some maximum number of steps.

This procedure should remind you of one you saw in a prior notebook (the least mean square algorithm for online regression!). As was true at the tricky bit is how to choose  $\alpha$ .

There is at least one difference between this procedure and the online regression procedure you learned earlier. Here, we are optimizing the full dataset rather than processing data points one at a time. (That is, the step iteration variable t used above is not used in exactly same way as the step iteration in LMS.)

Another question is, how do we know this procedure will converge to the global maximum, rather than, say, a local maximum? For that need a deeper analysis of a specific  $\mathcal{L}(\theta; y, X)$ , to show, for instance, that it is convex in  $\theta$ .

## Implementing logistic regression using MLE by gradient ascent

Let's apply the gradient ascent procedure to the logistic regression problem, in order to determine a good  $\theta$ .

**Exercise 7** (optional; ungraded). Show the following.

$$\nabla_{\theta} \left\{ \mathcal{L}(\theta; y, X) \right\} = X^{T} \left[ y - G(X \cdot \theta) \right].$$

**Answer.** From (V2),

$$\mathcal{L}(\theta; y, X) = y^{T} X \theta + u^{T} \ln G(-X\theta).$$

Thus,

$$\nabla_{\theta} \{ \mathcal{L}(\theta; \mathbf{v}, X) \} = \nabla_{\theta} (\mathbf{v}^T X \theta) + \nabla_{\theta} (\mathbf{u}^T \ln G(-X \theta)).$$

Let's consider each term in turn.

For the first term, apply the gradient identities to obtain

$$\nabla_{\theta} (y^T X \theta) = \nabla_{\theta} (\theta^T X^T y) = X^T y.$$

For the second term, recall the scalar interpretation of  $u^T \ln G(-X\theta)$ .

$$u^{T} \ln G(-X\theta) = \sum_{j=0}^{m-1} \ln G\left(-\hat{x}_{j}^{T}\theta\right).$$

The *i*-th component of the gradient is

$$\frac{\partial}{\partial \theta_i} \sum_{j=0}^{m-1} \ln G \left( -\hat{x}_j^T \theta \right) = \sum_{j=0}^{m-1} \frac{\partial}{\partial \theta_i} \ln G \left( -\hat{x}_j^T \theta \right).$$

Let's evaluate the summand:

$$\frac{\partial}{\partial \theta_{i}} \ln G \left( -\hat{x}_{j}^{T} \theta \right) = \underbrace{\left[ \frac{d}{dz} \ln G(z) \right] \cdot \left[ \frac{\partial z}{\partial \theta_{i}} \right]}_{\text{Let } z \equiv -\hat{x}_{j}^{T} \theta} = G(-z) \cdot \frac{\partial}{\partial \theta_{i}} \left( -\hat{x}_{j}^{T} \theta \right) = -G \left( \hat{x}_{j}^{T} \theta \right) \cdot x_{ji}.$$

Thus, the *i*-th component of the gradient becomes

$$\left[\nabla_{\theta} \left( u^{T} \ln G(-X\theta) \right) \right]_{i} = -\sum_{j=0}^{m-1} G\left( \hat{x}_{j}^{T}\theta \right) \cdot x_{ji}.$$

In other words, the full gradient vector is

$$\nabla_{\theta} \left( u^T \ln G(-X\theta) \right) = -X^T G(X\theta).$$

Putting the two components together,

$$\nabla_{\theta} \{ \mathcal{L}(\theta; y, X) \} = X^{T} y - X^{T} G(X\theta) = X^{T} [y - G(X\theta)].$$

Exercise 8 (2 points). Implement a function to compute the gradient of the log-likelihood. Your function should have the signature,

```
def grad_log_likelihood (theta, y, X):
```

```
In [21]: def grad log likelihood(theta, y, X):
              """Returns the gradient of the log-likelihood."""
             ### BEGIN SOLUTION
             return X.T.dot(y - logistic(X.dot(theta)))
              ### END SOLUTION
```

```
In [22]: # Test cell: `grad_log_likelihood_code__check`
         if False:
             d grad soln = 6
             m \text{ grad soln} = 399
             theta_grad_soln = np.random.random((d_grad_soln+1, 1)) * 2.0 - 1.0
             y_grad_soln = np.random.randint(low=0, high=2, size=(m_grad_soln, 1))
             X_grad_soln = np.random.random((m_grad_soln, d_grad_soln+1)) * 2.0 - 1.0
             X_grad_soln[:, 0] = 1.0
             L_grad_soln = grad_log_likelihood(theta_grad_soln, y_grad_soln, X_grad_soln)
             np.savez_compressed('grad_log_likelihood_soln',
                                  d grad soln, m grad soln, theta grad soln, y grad soln, X grad soln, L o
         npzfile_grad_soln = np.load ('{}grad_log_likelihood_soln.npz'.format(DATA_PATH))
         d_grad_soln = npzfile_grad_soln['arr_0']
         m grad soln = npzfile grad soln['arr 1']
         theta_grad_soln = npzfile_grad_soln['arr_2']
         y_grad_soln = npzfile_grad_soln['arr_3']
```

X\_grad\_soln = npzfile\_grad\_soln['arr\_4']

```
L_grad_soln = npzfile_grad_soln['arr_5']
L_grad_you = grad_log_likelihood (theta_grad_soln, y_grad_soln, X_grad_soln)
your grad err = np.max (np.abs (L grad you/L grad soln - 1.0))
x{solution}}}{\nabla\, \mathcal{L}_{\tiny \mbox{solution}}}\right\|_\infty \approx %g' % your_gi
assert your grad err <= 1e-12
print ("\n(Passed.)")
\left\| \frac{\nabla \mathcal{L}_{\text{yours}} - \nabla \mathcal{L}_{\text{solution}}}{\nabla \mathcal{L}_{\text{solution}}} \right\|_{\infty} \approx 2.22045e - 16
(Passed.)
```

**Exercise 9** (4 points). Implement the gradient ascent procedure to determine  $\theta$ , and try it out on the sample data.

Recall the procedure (repeated from above):

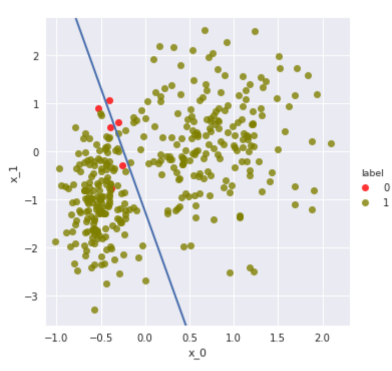
- Start with some initial guess,  $\theta(0)$ .
- At each iteration  $t \ge 0$  of the procedure, let  $\theta(t)$  be the current guess.
- Compute the direction of steepest ascent by evaluating the gradient,  $\Delta_t \equiv \nabla_{\theta(t)} \{ \mathcal{L}(\theta(t); y, X) \}$ .
- Define the step to be  $s_t \equiv \alpha \frac{\Delta_t}{\|\Delta_t\|}$ , where  $\alpha$  is a suitably chosen fudge factor.
- Take a step in the direction of the gradient,  $\theta(t+1) \leftarrow \theta(t) + s_t$ .
- Stop when the parameters don't change much or after some maximum number of steps.

In the code skeleton below, we've set up a loop to run a fixed number, MAX\_STEP, of gradient ascent steps. Also, when normalizing the step  $\Delta$ norm.

In your solution, we'd like you to store all guesses in the matrix thetas, so that you can later see how the  $\theta(t)$  values evolve. To extract particular column t, use the notation, theta[:, t:t+1]. This notation is necessary to preserve the "shape" of the column as a colur vector.

```
In [23]: ALPHA = 0.1
         MAX STEP = 250
         # Get the data coordinate matrix, X, and labels vector, y
         X = points
         y = labels.astype(dtype=float)
         # Store *all* guesses, for subsequent analysis
         thetas = np.zeros((3, MAX_STEP+1))
         for t in range(MAX_STEP):
              # Fill in the code to compute thetas[:, t+1:t+2]
              ### BEGIN SOLUTION
             theta t = thetas[:, t:t+1]
             delta_t = grad_log_likelihood(theta_t, y, X)
             delta_t = delta_t / np.linalg.norm(delta_t, ord=2)
             thetas[:, t+1:t+2] = theta_t + ALPHA*delta_t
              ### END SOLUTION
         theta_ga = thetas[:, MAX_STEP:]
         print("Your (hand) solution:", my theta.T.flatten())
         print("Computed solution:", theta_ga.T.flatten())
         print("\n=== Comparisons ===")
         display(Math (r'\dfrac{\theta 0}{\theta 2}:'))
         print("Your manual (hand-picked) solution is", my_theta[0]/my_theta[2], \
                , vs. MLE (via gradient ascent), which is", theta_ga[0]/theta_ga[2])
         display(Math (r'\dfrac{\theta_1}{\theta_2}:'))
         print("Your manual (hand-picked) solution is", my_theta[1]/my_theta[2], \
                ", vs. MLE (via gradient ascent), which is", theta_ga[1]/theta_ga[2])
         print("\n=== The MLE solution, visualized ===")
         ga_labels = gen_lin_discr_labels(points, theta_ga)
         df ga = df.copy()
         df_ga['label'] = mark_matches(ga_labels, labels).astype (dtype=int)
         plot_lin_discr(theta_ga, df_ga)
         Your (hand) solution: [-2. -0.5 -0.55]
         Computed solution: [-15.57666992 -3.03431905 -3.79328353]
         === Comparisons ===
         \frac{\theta_0}{\theta_2}:
         Your manual (hand-picked) solution is [3.63636364], vs. MLE (via gradient ascent), which is [4.
         Your manual (hand-picked) solution is [0.90909091], vs. MLE (via gradient ascent), which is [0.
```

=== The MLE solution, visualized ===



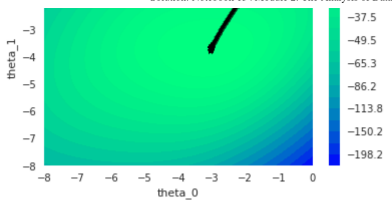
```
In [24]: | print ("\n=== Mismatch counts ===")
         my_labels = gen_lin_discr_labels (points, my_theta)
         my_mismatches = len (labels) - count_matches (labels, my_labels)
         print ("Your manual (hand-picked) solution has", num_mismatches, "mismatches.")
         ga_labels = gen_lin_discr_labels (points, theta_ga)
         ga_mismatches = len (labels) - count_matches (labels, ga_labels)
         print ("The MLE method produces", ga_mismatches, "mismatches.")
         assert ga mismatches <= 8</pre>
         print ("\n(Passed.)")
         === Mismatch counts ===
         Your manual (hand-picked) solution has 5 mismatches.
         The MLE method produces 6 mismatches.
         (Passed.)
```

The gradient ascent trajectory. Let's take a look at how gradient ascent progresses. (You might try changing the  $\alpha$  parameter and see how it results.)

```
In [25]: n_ll_grid = 100
         x1 = np.linspace(-8., 0., n_ll_grid)
         x2 = np.linspace(-8., 0., n_ll_grid)
         x1_grid, x2_grid = np.meshgrid(x1, x2)
         ll_grid = np.zeros((n_ll_grid, n_ll_grid))
         for i1 in range(n_ll_grid):
             for i2 in range(n_ll_grid):
                 theta_i1_i2 = np.array([[thetas[0, MAX_STEP]],
                                          [x1_grid[i1][i2]],
                                          [x2_grid[i1][i2]])
                  ll_grid[i1][i2] = log_likelihood(theta_i1_i2, y, X)
         # Determine a color scale
         def v(x):
             return -np.log(np.abs(x))
             return x
         def v_inv(v):
             return -np.exp(np.abs(v))
             return v
         v_min, v_max = v(ll_grid.min()), v(ll_grid.max())
         v_range = v_max - v_min
         v_breaks = v_inv(np.linspace(v_min, v_max, 20))
         p = plt.contourf(x1, x2, ll_grid, v_breaks, cmap=plt.cm.get_cmap("winter"))
         plt.xlabel('theta 0')
         plt.ylabel('theta_1')
         plt.title('log-likelihood')
         plt.colorbar()
         plt.plot(thetas[1, :], thetas[2, :], 'k*-')
```

Out[25]: [<matplotlib.lines.Line2D at 0x7f0ec2484dd8>]

```
log-likelihood
 0
                                                            -16.3
                                                            -21.5
-1
                                                            -28.4
-2
```



# Part 5 (optional): Numerical optimization via Newton's method

The fudge factor,  $\alpha$ , in gradient ascent should give you pause. Can you choose the step size or direction in a better or more principled way?

One idea is Newton's method (http://www.math.uiuc.edu/documenta/vol-ismp/13\_deuflhard-peter.pdf), summarized below.

This part of the notebook has additional exercises, but they are all worth 0 points. (So if you submit something that is incomplete or fail test cells, you won't lose any points.)

The basic idea, in 1-D. Suppose you start at a point x and, assuming you are not yet at the optimum, you have decided to take a step of size you at f(x + s).

How do you choose s? In gradient ascent, you do so by following the gradient, which points in an "upward" direction.

In Newton's method, you will pick s in a different way: choose s to maximize f(x + s).

That should strike you as circular; the whole problem from the beginning was to maximize f(x). The trick, in this case, is not to maximize f(x)rather, let's replace it with some approximation,  $q(s) \approx f(x+s)$ , and maximize q(s) instead.

A simple choice for q(s) is a quadratic function in s. This choice is motivated by two factors: (a) since it's quadratic, it should have some sort or point (and hopefully an actual maximum), and (b) it is a higher-order approximation than a linear one, and so hopefully more accurate than a line well.

$$f(x+s) \approx f(x) + s\frac{df}{dx} + \frac{1}{2}s^2\frac{d^2f}{dx^2} \equiv q(s).$$

To maximize q(s), take its derivative and then solve for the  $s_*$  such that  $q(s_*) = 0$ :

consists of the following steps:

$$\left. \frac{dq}{ds} \right|_{s=s} = \frac{df}{dx} + s_* \frac{d^2f}{dx^2} = 0$$

$$\implies s_* = -\frac{df}{dx} \left( \frac{d^2f}{dx^2} \right)^{-1}.$$

That is, the optimal step  $s_*$  is the negative of the first derivative of f divided by its second derivative.

**Generalizing to higher dimensions.** To see how this procedure works in higher dimensions, you will need not only the gradient of f(x), but als which is the moral equivalent of a second derivative.

Definition: the Hessian. Let f(v) be a function that takes a vector v of length n as input and returns a scalar. The Hessian of f(v) is an  $n \times n$  made nwhose entries are all  $n^2$  possible second-order partial derivatives with respect to the components of v. That is, let  $h_{ij}$  be the (i,j) element of Hdefine

$$h_{ij} \equiv \frac{\partial^2}{\partial v_i \partial v_j} f(v).$$

Armed with a Hessian, the Newton step is defined as follows, by direct analogy to the 1-D case. First, the Taylor series approximation of f(x +multidimensional variables is, as it happens,

$$f(x+s) \approx f(x) + s^T \nabla_x f + \frac{1}{2} s^T H_x(f) s \equiv q(s).$$

As in the 1-D case, we want to find an extreme point of q(s). Taking its "derivative" (gradient),  $\nabla_s q$ , and setting it to 0 yields,

$$\nabla_{s} q(s) = \nabla_{x} f(x) + H_{x}(f) s = 0$$
  
$$\implies H_{x}(f) \cdot s = -\nabla_{x} f(x).$$

In other words, to choose the next step s, Newton's method suggests that you must solve a system of linear equations, where the matrix is the and the right-hand side is the negative gradient of f.

**Summary: Newton's method.** Summarizing the main ideas from above, Newton's method to maximize the scalar objective function f(x) where

- Start with some initial guess x(0).
- At step t, compute the search direction s(t) by solving  $H_{x(t)}(f) \cdot s(t) = -\nabla_x f(x(t))$ .
- Compute a new (and hopefully improved) guess by the update,  $x(t+1) \leftarrow x(t) + s(t)$ .

#### Implementing logistic regression via a Newton-based MLE

To perform MLE for the logistic regression model using Newton's method, you need both the gradient of the log-likelihood as well as the Hessia already know how to compute the gradient from the preceding exercises; so what about the Hessian?

Notationally, that calculation will be a little bit easier to write down and program with the following definition.

Definition: Elementwise product. Let  $A \equiv (a_{ij})$  and  $B \equiv (b_{ij})$  be  $m \times n$  matrices. Denote the elementwise product of A and B by  $A \odot B$ . That  $C = A \odot B$ , then element  $c_{ij} = a_{ij} \cdot b_{ij}$ .

If A is  $m \times n$  but B is instead just  $m \times 1$ , then we will "auto-extend" B. Put differently, if B has the same number of rows as A but only 1 column take  $C = A \odot B$  to have elements  $c_{ii} = a_{ii} \cdot b_i$ .

In Python, you can use np.multiply() (http://docs.scipy.org/doc/numpy/reference/generated/numpy.multiply.html) for elementwise multiplic Numpy arrays.

Exercise 10 (optional; ungraded). Show that the Hessian of the log-likelihood for logistic regression is

```
H_{\theta}(\mathcal{L}(\theta; l, X)) = -(X \odot G(X\theta))^{T} (X \odot G(-X\theta)).
```

Exercise 11 (0 points). Implement a function to compute the Hessian of the log-likelihood. The signature of your function should be,

```
def hess_log_likelihood (theta, y, X):
    ...
```

assert vour hess err <= 1e-12</pre>

```
In [27]: ### BEGIN SOLUTION

def hess_log_likelihood(theta, y, X):
    """Returns the Hessian of the log-likelihood."""
    z = X.dot(theta)
    A = np.multiply(X, logistic(z))
    B = np.multiply(X, logistic(-z))
    return -A.T.dot(B)

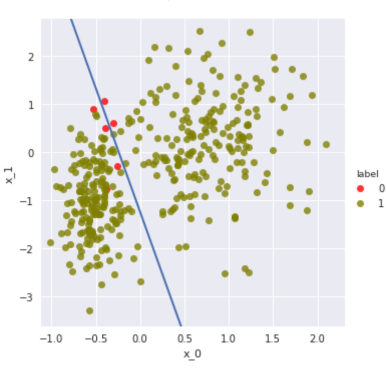
### END SOLUTION
```

```
In [28]: # Test cell: `hess log likelihood check`
        if False:
            d_{hess\_soln} = 20
            m hess soln = 501
            theta hess soln = np.random.random ((d hess soln+1, 1)) * 2.0 - 1.0
            y_hess_soln = np.random.randint (low=0, high=2, size=(m_hess_soln, 1))
            X_hess_soln = np.random.random ((m_hess_soln, d_hess_soln+1)) * 2.0 - 1.0
            X_hess_soln[:, 0] = 1.0
            L hess soln = hess log likelihood (theta hess soln, y hess soln, X hess soln)
            np.savez compressed ('hess log likelihood soln',
                                d_hess_soln, m_hess_soln, theta_hess_soln, y_hess_soln, X_hess_soln, L_
        npzfile hess soln = np.load ('{}hess log likelihood soln.npz'.format(DATA PATH))
        d_hess_soln = npzfile_hess_soln['arr_0']
        m hess soln = npzfile hess soln['arr 1']
        theta_hess_soln = npzfile_hess_soln['arr_2']
        y hess soln = npzfile hess soln['arr 3']
        X hess soln = npzfile hess soln['arr 4']
        L hess soln = npzfile hess soln['arr 5']
        L hess you = hess log likelihood(theta hess soln, y hess soln, X hess soln)
        your_hess_err = np.max(np.abs(L_hess_you/L_hess_soln - 1.0))
        }}\right\| \infty \approx %g' % your hess err))
```

```
print ("\n(Passed.)")  \left\| \frac{H_{\text{yours}} - H_{\text{solution}}}{H_{\text{solution}}} \right\|_{\infty} \approx 9.74776e - 14  (Passed.)
```

**Exercise 12** (0 points). Finish the implementation of a Newton-based MLE procedure for the logistic regression problem.

```
In [29]: MAX_STEP = 10
         # Get the data coordinate matrix, X, and labels vector, 1
         X = points
         y = labels.astype(dtype=float)
         # Store *all* guesses, for subsequent analysis
         thetas_newt = np.zeros((3, MAX_STEP+1))
         for t in range(MAX_STEP):
             ### BEGIN SOLUTION
             theta_t = thetas_newt[:, t:t+1]
             g_t = grad_log_likelihood(theta_t, y, X)
             H t = hess_log_likelihood(theta_t, y, X)
             s_t = np.linalg.solve(H_t, -g_t)
             thetas_newt[:, t+1:t+2] = theta_t + s_t
             ### END SOLUTION
         theta_newt = thetas_newt[:, MAX_STEP:]
         print ("Your (hand) solution:", my_theta.T.flatten())
         print ("Computed solution:", theta_newt.T.flatten())
         print ("\n=== Comparisons ===")
         display (Math (r'\dfrac{\theta_0}{\theta_2}:'))
         print ("Your manual (hand-picked) solution is", my_theta[0]/my_theta[2], \
               ", vs. MLE (via Newton's method), which is", theta newt[0]/theta newt[2])
         display (Math (r'\dfrac{\theta_1}{\theta_2}:'))
         print ("Your manual (hand-picked) solution is", my_theta[1]/my_theta[2], \
                , vs. MLE (via Newton's method), which is", theta_newt[1]/theta_newt[2])
         print ("\n=== The MLE solution, visualized ===")
         newt_labels = gen_lin_discr_labels(points, theta_newt)
         df_newt = df.copy()
         df newt['label'] = mark matches(newt labels, labels).astype (dtype=int)
         plot_lin_discr(theta_newt, df_newt)
         Your (hand) solution: [-2. -0.5 -0.55]
         Computed solution: [-15.63082207 -3.04255951 -3.76500606]
         === Comparisons ===
         Your manual (hand-picked) solution is [3.63636364], vs. MLE (via Newton's method), which is [4.
         Your manual (hand-picked) solution is [0.90909091], vs. MLE (via Newton's method), which is [0.
         === The MLE solution, visualized ===
```



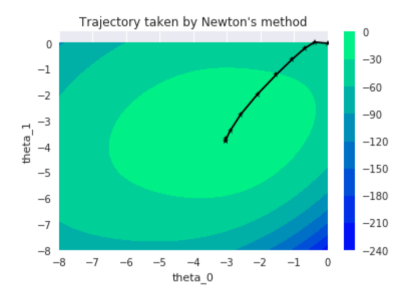
```
In [30]: # Test cell: `logreg_mle_newt__check`
    print ("\n=== Mismatch counts ===")
```

```
my_labels = gen_lin_discr_labels (points, my_theta)
my mismatches = len (labels) - count matches (labels, my labels)
print ("Your manual (hand-picked) solution has", num mismatches, "mismatches.")
newt_labels = gen_lin_discr_labels (points, theta_newt)
newt mismatches = len (labels) - count matches (labels, newt labels)
print ("The MLE+Newton method produces", newt_mismatches, "mismatches.")
assert newt_mismatches <= ga_mismatches</pre>
print ("\n(Passed.)")
=== Mismatch counts ===
Your manual (hand-picked) solution has 5 mismatches.
The MLE+Newton method produces 6 mismatches.
(Passed.)
```

The following cell creates a contour plot of the log-likelihood, as done previously in this notebook. Add code to display the trajectory taken by I method.

```
In [31]: p = plt.contourf(x1, x2, ll_grid, cmap=plt.cm.get_cmap("winter"))
         plt.xlabel('theta_0')
         plt.ylabel('theta_1')
         plt.title('Trajectory taken by Newton\'s method')
         plt.colorbar()
         plt.plot(thetas_newt[1, :], thetas_newt[2, :], 'k*-')
```

Out[31]: [<matplotlib.lines.Line2D at 0x7f0ec2609b38>]



How many steps does this optimization procedure take compared to gradient ascent? What is the tradeoff?

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