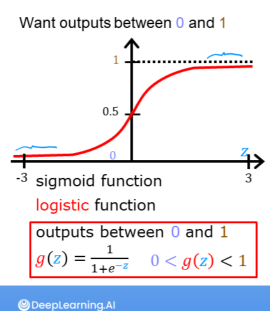
## **Optional Lab: Logistic Regression**

In this ungraded lab, you will

- explore the sigmoid function (also known as the logistic function)
- explore logistic regression; which uses the sigmoid function

# In [1]: import numpy as np %matplotlib widget import matplotlib.pyplot as plt from plt\_one\_addpt\_onclick import plt\_one\_addpt\_onclick from lab\_utils\_common import draw\_vthresh plt.style.use('./deeplearning.mplstyle')

#### **Sigmoid or Logistic Function**



As discussed in the lecture videos, for a classification task, we can start by using our linear regression model,

$$f_{\mathbf{w},b}(\mathbf{x}^{(i)}) = \mathbf{w} \cdot \mathbf{x}^{(i)} + b$$
, to predict  $y$  given  $x$ .

- However, we would like the predictions of our classification model to be between 0 and 1 since our output variable y is either 0 or 1.
- This can be accomplished by using a "sigmoid function" which maps all input values to values between 0 and 1.

Let's implement the sigmoid function and see this for ourselves.

## Formula for Sigmoid function

The formula for a sigmoid function is as follows -

$$g(z) = \frac{1}{1 + e^{-z}} \tag{1}$$

In the case of logistic regression, z (the input to the sigmoid function), is the output of a linear regression model.

- In the case of a single example, z is scalar.
- in the case of multiple examples, z may be a vector consisting of m values, one for each example.
- The implementation of the sigmoid function should cover both of these potential input formats. Let's implement this in Python.

NumPy has a function called exp() (https://numpy.org/doc/stable/reference/generated/numpy.exp.html), which offers a convenient way to calculate the exponential ( $e^z$ ) of all elements in the input array (z).

It also works with a single number as an input, as shown below.

```
In [2]: # Input is an array.
    input_array = np.array([1,2,3])
    exp_array = np.exp(input_array)

print("Input to exp:", input_array)
print("Output of exp:", exp_array)

# Input is a single number
input_val = 1
exp_val = np.exp(input_val)

print("Input to exp:", input_val)
print("Output of exp:", exp_val)

Input to exp: [1 2 3]
Output of exp: [2.72 7.39 20.09]
Input to exp: 1
Output of exp: 2.718281828459045
```

The sigmoid function is implemented in python as shown in the cell below.

Let's see what the output of this function is for various value of z

```
In [4]: # Generate an array of evenly spaced values between -10 and 10
        z_{tmp} = np.arange(-10,11)
        # Use the function implemented above to get the sigmoid values
        y = sigmoid(z tmp)
        # Code for pretty printing the two arrays next to each other
        np.set_printoptions(precision=3)
        print("Input (z), Output (sigmoid(z))")
        print(np.c_[z_tmp, y])
        Input (z), Output (sigmoid(z))
         [[-1.000e+01 4.540e-05]
         [-9.000e+00 1.234e-04]
          [-8.000e+00 \quad 3.354e-04]
          [-7.000e+00 9.111e-04]
         [-6.000e+00 \quad 2.473e-03]
          [-5.000e+00 6.693e-03]
          [-4.000e+00 1.799e-02]
```

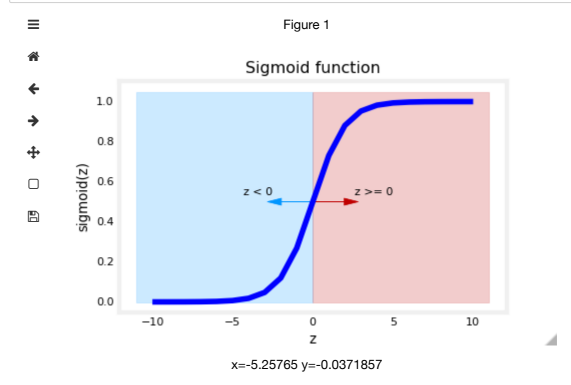
The values in the left column are z, and the values in the right column are sigmoid(z). As you can see, the input values to the sigmoid range from -10 to 10, and the output values range from 0 to 1.

Now, let's try to plot this function using the matplotlib library.

[-3.000e+00 4.743e-02]
[-2.000e+00 1.192e-01]
[-1.000e+00 2.689e-01]
[ 0.000e+00 5.000e-01]
[ 1.000e+00 7.311e-01]
[ 2.000e+00 8.808e-01]
[ 3.000e+00 9.526e-01]
[ 4.000e+00 9.820e-01]
[ 5.000e+00 9.933e-01]
[ 6.000e+00 9.975e-01]
[ 7.000e+00 9.991e-01]
[ 8.000e+00 9.997e-01]
[ 9.000e+00 9.999e-01]
[ 1.000e+01 1.000e+00]

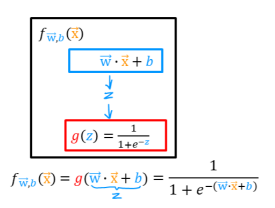
```
In [5]: # Plot z vs sigmoid(z)
fig,ax = plt.subplots(1,1,figsize=(5,3))
ax.plot(z_tmp, y, c="b")

ax.set_title("Sigmoid function")
ax.set_ylabel('sigmoid(z)')
ax.set_xlabel('z')
draw_vthresh(ax,0)
```



As you can see, the sigmoid function approaches 0 as z goes to large negative values and approaches 1 as z goes to large positive values.

## **Logistic Regression**



"logistic regression"

A logistic regression model applies the sigmoid to the familiar linear regression model as shown below:

$$f_{\mathbf{w},b}(\mathbf{x}^{(i)}) = g(\mathbf{w} \cdot \mathbf{x}^{(i)} + b) \qquad (2)$$

where

$$g(z) = \frac{1}{1 + e^{-z}} \tag{3}$$

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### Interpretation of logistic regression output

```
f_{\overrightarrow{w},b}(\overrightarrow{x}) = \frac{1}{1 + e^{-(\overrightarrow{w} \cdot \overrightarrow{x} + b)}}
f_{\overrightarrow{w},b}(\overrightarrow{x}) = P(y = 1 | \overrightarrow{x}; \overrightarrow{w}, b)
f_{\overrightarrow{w},b}(\overrightarrow{x}) = P(y = 1 | \overrightarrow{x}; \overrightarrow{w}, b)
f_{\overrightarrow{w},b}(\overrightarrow{x}) = P(y = 1 | \overrightarrow{x}; \overrightarrow{w}, b)
f_{\overrightarrow{w},b}(\overrightarrow{x}) = 0.7
```

Let's apply logistic regression to the categorical data example of tumor classification. First, load the examples and initial values for the parameters.

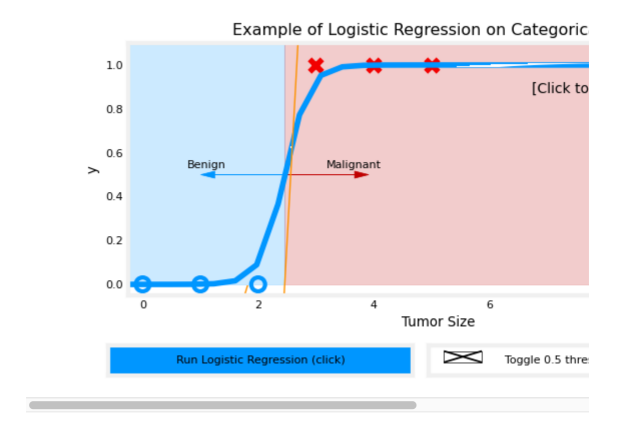
```
In [12]: x_train = np.array([0., 1, 2, 3, 4, 5])
y_train = np.array([0, 0, 0, 1, 1, 1])

w_in = np.zeros((1))
b_in = 0
```

Try the following steps:

- Click on 'Run Logistic Regression' to find the best logistic regression model for the given training data
  - Note the resulting model fits the data quite well.
  - Note, the orange line is z' or  $\mathbf{w} \cdot \mathbf{x}^{(i)} + b$  above. It does not match the line in a linear regression model. Further improve these results by applying a *threshold*.
- Tick the box on the 'Toggle 0.5 threshold' to show the predictions if a threshold is applied.
  - These predictions look good. The predictions match the data
  - Now, add further data points in the large tumor size range (near 10), and re-run logistic regression.
  - unlike the linear regression model, this model continues to make correct predictions

In [13]: plt.close('all')
addpt = plt\_one\_addpt\_onclick( x\_train,y\_train, w\_in, b\_in, logistic



## **Congratulations!**

You have explored the use of the sigmoid function in logistic regression.

In [ ]: