Initialization

Welcome to the first assignment of "Improving Deep Neural Networks".

Training your neural network requires specifying an initial value of the weights. A well chosen initialization method will help learning.

If you completed the previous course of this specialization, you probably followed our instructions for weight initialization, and it has worked out so far. But how do you choose the initialization for a new neural network? In this notebook, you will see how different initializations lead to different results.

A well chosen initialization can:

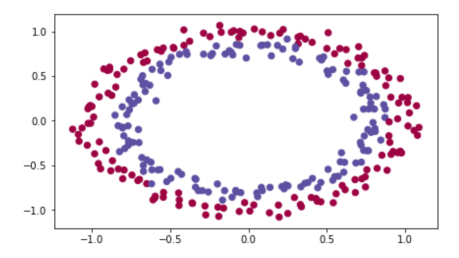
- · Speed up the convergence of gradient descent
- Increase the odds of gradient descent converging to a lower training (and generalization) error

To get started, run the following cell to load the packages and the planar dataset you will try to classify.

```
In [2]: import numpy as np
   import matplotlib.pyplot as plt
   import sklearn
   import sklearn.datasets
   from init_utils import sigmoid, relu, compute_loss, forward_propagation, backward_propagation
   from init_utils import update_parameters, predict, load_dataset, plot_decision_boundary, predict_dec

%matplotlib inline
   plt.rcParams['figure.figsize'] = (7.0, 4.0) # set default size of plots
   plt.rcParams['image.interpolation'] = 'nearest'
   plt.rcParams['image.cmap'] = 'gray'

# load image dataset: blue/red dots in circles
   train_X, train_Y, test_X, test_Y = load_dataset()
```



You would like a classifier to separate the blue dots from the red dots.

1 - Neural Network model

You will use a 3-layer neural network (already implemented for you). Here are the initialization methods you will experiment with:

• Zeros initialization -- setting initialization = "zeros" in the input argument.

- Random initialization -- setting initialization = "random" in the input argument. This initializes the weights to large random values.
- He initialization -- setting initialization = "he" in the input argument. This initializes the weights to random values scaled according to a paper by He et al., 2015.

Instructions: Please quickly read over the code below, and run it. In the next part you will implement the three initialization methods that this model() calls.

```
In [3]:
         def model(X, Y, learning rate = 0.01, num iterations = 15000, print cost = True, initialization = "h
              Implements a three-layer neural network: LINEAR->RELU->LINEAR->RELU->LINEAR->SIGMOID.
              Arguments:
              X -- input data, of shape (2, number of examples)
              Y -- true "label" vector (containing 0 for red dots; 1 for blue dots), of shape (1, number of ex
              learning_rate -- learning rate for gradient descent
              num_iterations -- number of iterations to run gradient descent
              print cost -- if True, print the cost every 1000 iterations
              initialization -- flag to choose which initialization to use ("zeros", "random" or "he")
              Returns:
              parameters -- parameters learnt by the model
              grads = \{\}
              costs = [] # to keep track of the loss
              m = X. shape[1] # number of examples
              layers dims = [X. shape[0], 10, 5, 1]
              # Initialize parameters dictionary.
              if initialization == "zeros":
                  parameters = initialize_parameters_zeros(layers_dims)
              elif initialization == "random":
                  parameters = initialize parameters random(layers dims)
              elif initialization == "he":
                  parameters = initialize parameters he(layers dims)
              # Loop (gradient descent)
              for i in range (0, num iterations):
                  # Forward propagation: LINEAR -> RELU -> LINEAR -> RELU -> LINEAR -> SIGMOID.
                  a3, cache = forward propagation(X, parameters)
                  # Loss
                  cost = compute loss(a3, Y)
                  # Backward propagation.
                  grads = backward propagation(X, Y, cache)
                  # Update parameters.
                  parameters = update parameters (parameters, grads, learning rate)
                  # Print the loss every 1000 iterations
                  if print cost and i \% 1000 == 0:
                      print("Cost after iteration {}: {}".format(i, cost))
                      costs. append (cost)
              # plot the loss
              plt. plot (costs)
              plt.ylabel('cost')
              plt.xlabel('iterations (per hundreds)')
              plt. title("Learning rate =" + str(learning_rate))
              plt. show()
              return parameters
```

2 - Zero initialization

There are two types of parameters to initialize in a neural network:

```
• the weight matrices (W^{[1]},W^{[2]},W^{[3]},\dots,W^{[L-1]},W^{[L]})
• the bias vectors (b^{[1]},b^{[2]},b^{[3]},\dots,b^{[L-1]},b^{[L]})
```

Exercise: Implement the following function to initialize all parameters to zeros. You'll see later that this does not work well since it fails to "break symmetry", but lets try it anyway and see what happens. Use np.zeros((..,..)) with the correct shapes.

```
In [4]: | # GRADED FUNCTION: initialize_parameters_zeros
         def initialize parameters zeros(layers dims):
              Arguments:
              layer dims -- python array (list) containing the size of each layer.
              Returns:
              parameters -- python dictionary containing your parameters "W1", "b1", ..., "WL", "bL":
                              W1 -- weight matrix of shape (layers_dims[1], layers_dims[0])
                              b1 -- bias vector of shape (layers dims[1], 1)
                              WL -- weight matrix of shape (layers_dims[L], layers_dims[L-1])
                              bL -- bias vector of shape (layers dims[L], 1)
              parameters = {}
              L = len(layers dims)
                                              # number of layers in the network
              for 1 in range (1, L):
                  ### START CODE HERE ### (\approx 2 lines of code)
                  parameters['W' + str(1)] = np. zeros((layers dims[1], layers dims[1-1]))
                  parameters['b' + str(1)] = np. zeros((layers dims[1], 1))
                  ### END CODE HERE ###
             return parameters
```

Expected Output:

W2 = [[0. 0.]]b2 = [[0.]]

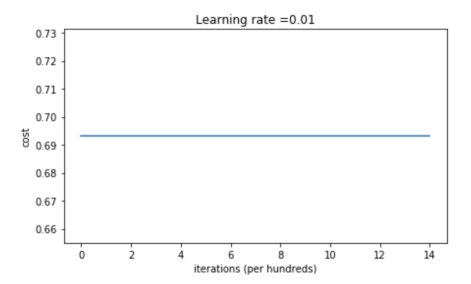
 $\begin{bmatrix} 0. \end{bmatrix}$

```
W1 [[ 0. 0. 0.] [ 0. 0. 0.]]
b1 [[ 0.] [ 0.]]
W2 [[ 0. 0.]]
b2 [[ 0.]]
```

Run the following code to train your model on 15,000 iterations using zeros initialization.

```
In [7]: parameters = model(train_X, train_Y, initialization = "zeros")
    print ("On the train set:")
    predictions_train = predict(train_X, train_Y, parameters)
    print ("On the test set:")
    predictions_test = predict(test_X, test_Y, parameters)
```

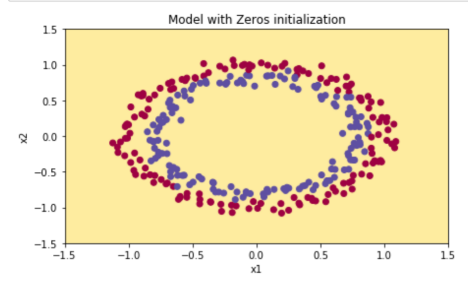
```
Cost after iteration 0: 0.6931471805599453
Cost after iteration 1000: 0.6931471805599453
Cost after iteration 2000: 0.6931471805599453
Cost after iteration 3000: 0.6931471805599453
Cost after iteration 4000: 0.6931471805599453
Cost after iteration 5000: 0.6931471805599453
Cost after iteration 6000: 0.6931471805599453
Cost after iteration 7000: 0.6931471805599453
Cost after iteration 8000: 0.6931471805599453
Cost after iteration 9000: 0.6931471805599453
Cost after iteration 10000: 0.6931471805599453
Cost after iteration 10000: 0.6931471805599453
Cost after iteration 12000: 0.6931471805599453
Cost after iteration 13000: 0.6931471805599453
Cost after iteration 13000: 0.6931471805599453
```



On the train set: Accuracy: 0.5 On the test set: Accuracy: 0.5

The performance is really bad, and the cost does not really decrease, and the algorithm performs no better than random guessing. Why? Lets look at the details of the predictions and the decision boundary:

```
In [9]: plt.title("Model with Zeros initialization")
    axes = plt.gca()
    axes.set_xlim([-1.5, 1.5])
    axes.set_ylim([-1.5, 1.5])
    plot_decision_boundary(lambda x: predict_dec(parameters, x.T), train_X, train_Y)
```



The model is predicting 0 for every example.

In general, initializing all the weights to zero results in the network failing to break symmetry. This means that every neuron in each layer will learn the same thing, and you might as well be training a neural network with $n^{[l]} = 1$ for every layer, and the network is no more powerful than a linear classifier such as logistic regression.

What you should remember:

- The weights $W^{[l]}$ should be initialized randomly to break symmetry.
- It is however okay to initialize the biases $b^{[l]}$ to zeros. Symmetry is still broken so long as $W^{[l]}$ is initialized randomly.

3 - Random initialization

To break symmetry, lets intialize the weights randomly. Following random initialization, each neuron can then proceed to learn a different function of its inputs. In this exercise, you will see what happens if the weights are intialized randomly, but to very large values.

Exercise: Implement the following function to initialize your weights to large random values (scaled by *10) and your biases to zeros. Use np. random. randn(..,..) * 10 for weights and np. zeros((.., ..)) for biases. We are using a fixed np. random. seed(..) to make sure your "random" weights match ours, so don't worry if running several times your code gives you always the same initial values for the parameters.

```
In [10]: | # GRADED FUNCTION: initialize_parameters_random
           def initialize_parameters_random(layers_dims):
               Arguments:
               layer dims -- python array (list) containing the size of each layer.
               Returns:
               parameters -- python dictionary containing your parameters "W1", "b1", ..., "WL", "bL":
                                W1 -- weight matrix of shape (layers dims[1], layers dims[0])
                                b1 -- bias vector of shape (layers dims[1], 1)
                                WL -- weight matrix of shape (layers dims[L], layers dims[L-1])
                                bL -- bias vector of shape (layers dims[L], 1)
                                                 # This seed makes sure your "random" numbers will be the as ours
               np. random. seed (3)
               parameters = {}
               L = len(layers dims)
                                                 # integer representing the number of layers
               for 1 in range (1, L):
                    ### START CODE HERE ### (pprox 2 lines of code)
                   parameters['W' + str(1)] = np. random. randn(layers_dims[1], layers_dims[1-1])*10
                   parameters['b' + str(1)] = np. zeros((layers_dims[1], 1))
                    ### END CODE HERE ###
               return parameters
           parameters = initialize_parameters_random([3, 2, 1])
In [11]:
           print("W1 = " + str(parameters["W1"]))
           print("b1 = " + str(parameters["b1"]))
           print("W2 = " + str(parameters["W2"]))
           print("b2 = " + str(parameters["b2"]))
           W1 = [ [ 17.88628473 ]
                                 4.36509851
            [-18.63492703 -2.77388203 -3.54758979]]
           b1 = \lceil \lceil 0. \rceil
           [ 0.]]
           W2 = [[-0.82741481 -6.27000677]]
           b2 = \lceil \lceil 0. \rceil \rceil
           Expected Output:
                           W1 [[ 17.88628473 4.36509851 0.96497468] [-18.63492703 -2.77388203 -3.54758979]]
                            b1
                                                                                            [[ 0.] [ 0.]]
```

Run the following code to train your model on 15,000 iterations using random initialization.

[[-0.82741481 -6.27000677]]

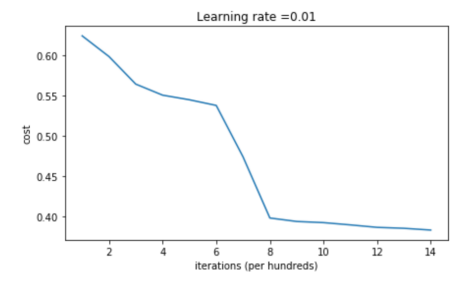
[[.0.]]

W2

b2

```
In [13]: parameters = model(train_X, train_Y, initialization = "random")
    print ("On the train set:")
    predictions_train = predict(train_X, train_Y, parameters)
    print ("On the test set:")
    predictions_test = predict(test_X, test_Y, parameters)
```

```
Cost after iteration 0: inf
Cost after iteration 1000: 0.6237287551108738
Cost after iteration 2000: 0.5981106708339466
Cost after iteration 3000: 0.5638353726276827
Cost after iteration 4000: 0.550152614449184
Cost after iteration 5000: 0.5444235275228304
Cost after iteration 6000: 0.5374184054630083
Cost after iteration 7000: 0.47357131493578297
Cost after iteration 8000: 0.39775634899580387
Cost after iteration 9000: 0.3934632865981078
Cost after iteration 10000: 0.39202525076484457
Cost after iteration 11000: 0.38921493051297673
Cost after iteration 12000: 0.38614221789840486
Cost after iteration 13000: 0.38497849983013926
Cost after iteration 14000: 0.38278397192120406
```

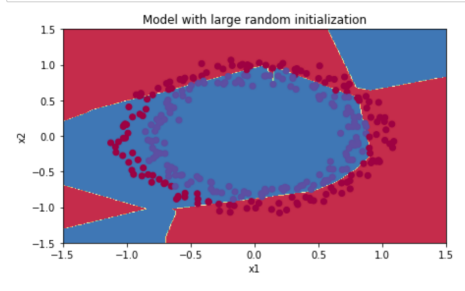


On the train set: Accuracy: 0.83 On the test set: Accuracy: 0.86

If you see "inf" as the cost after the iteration 0, this is because of numerical roundoff; a more numerically sophisticated implementation would fix this. But this isn't worth worrying about for our purposes.

Anyway, it looks like you have broken symmetry, and this gives better results. than before. The model is no longer outputting all 0s.

```
In [15]: plt.title("Model with large random initialization")
   axes = plt.gca()
   axes.set_xlim([-1.5, 1.5])
   axes.set_ylim([-1.5, 1.5])
   plot_decision_boundary(lambda x: predict_dec(parameters, x.T), train_X, train_Y)
```



Observations:

In [14]:

print (predictions train)

- The cost starts very high. This is because with large random-valued weights, the last activation (sigmoid) outputs results that are very close to 0 or 1 for some examples, and when it gets that example wrong it incurs a very high loss for that example. Indeed, when $\log(a^{[3]}) = \log(0)$, the loss goes to infinity.
- Poor initialization can lead to vanishing/exploding gradients, which also slows down the optimization algorithm.
- If you train this network longer you will see better results, but initializing with overly large random numbers slows down the optimization.

In summary:

- Initializing weights to very large random values does not work well.
- Hopefully intializing with small random values does better. The important question is: how small should be these random values be? Lets find out in the next part!

4 - He initialization

Finally, try "He Initialization"; this is named for the first author of He et al., 2015. (If you have heard of "Xavier initialization", this is similar except Xavier initialization uses a scaling factor for the weights $W^{[l]}$ of $sqrt(1./layers_dims[1-1])$ where He initialization would use $sqrt(2./layers_dims[1-1])$.)

Exercise: Implement the following function to initialize your parameters with He initialization.

Hint: This function is similar to the previous $initialize_parameters_random(...)$. The only difference is that instead of multiplying np. random. randn(...,..) by 10, you will multiply it by $\sqrt{\frac{2}{dimension of the previous layer}}$, which is what He initialization recommends for layers with a ReLU activation.

```
In [16]:
           # GRADED FUNCTION: initialize parameters he
           def initialize_parameters_he(layers_dims):
               Arguments:
               layer dims -- python array (list) containing the size of each layer.
               Returns:
               parameters -- python dictionary containing your parameters "W1", "b1", ..., "WL", "bL":
                                W1 -- weight matrix of shape (layers_dims[1], layers_dims[0])
                                b1 -- bias vector of shape (layers dims[1], 1)
                                WL -- weight matrix of shape (layers dims[L], layers dims[L-1])
                                bL — bias vector of shape (layers dims[L], 1)
               np. random. seed (3)
               parameters = {}
               L = len(layers_dims) - 1 # integer representing the number of layers
               for 1 in range (1, L + 1):
                    ### START CODE HERE ### (\approx 2 lines of code)
                   parameters['W' + str(1)] = np. random. randn(layers dims[1], layers dims[1-1])*np. sqrt(2./layer
                   parameters['b' + str(1)] = np. zeros((layers dims[1], 1))
                    ### END CODE HERE ###
               return parameters
In [17]:
           parameters = initialize parameters he([2, 4, 1])
           print("W1 = " + str(parameters["W1"]))
           print("b1 = " + str(parameters["b1"]))
           print("W2 = " + str(parameters["W2"]))
           print("b2 = " + str(parameters["b2"]))
           W1 = [[1.78862847 \ 0.43650985]]
            [ 0.09649747 -1.8634927 ]
            [-0.2773882 -0.35475898]
            [-0.08274148 -0.62700068]
           b1 = \lceil \lceil 0. \rceil
            [ 0.]
            \begin{bmatrix} 0. \end{bmatrix}
            [ 0. ]]
           W2 = [[-0.03098412 -0.33744411 -0.92904268 0.62552248]]
           b2 = \lceil \lceil 0. \rceil \rceil
```

Expected Output:

```
W1 [[ 1.78862847 0.43650985] [ 0.09649747 -1.8634927 ] [-0.2773882 -0.35475898] [-0.08274148 -0.62700068]]
b1 [[ 0.] [ 0.] [ 0.] [ 0.] [ 0.]
W2 [[-0.03098412 -0.33744411 -0.92904268 0.62552248]]
```

b2 [[0.]]

Run the following code to train your model on 15,000 iterations using He initialization.

```
In [ ]: parameters = model(train_X, train_Y, initialization = "he")
    print ("On the train set:")
    predictions_train = predict(train_X, train_Y, parameters)
    print ("On the test set:")
    predictions_test = predict(test_X, test_Y, parameters)
```

```
In [ ]: plt.title("Model with He initialization")
    axes = plt.gca()
    axes.set_xlim([-1.5, 1.5])
    axes.set_ylim([-1.5, 1.5])
    plot_decision_boundary(lambda x: predict_dec(parameters, x.T), train_X, train_Y)
```

Observations:

• The model with He initialization separates the blue and the red dots very well in a small number of iterations.

5 - Conclusions

You have seen three different types of initializations. For the same number of iterations and same hyperparameters the comparison is:

| Problem/Comment | Train accuracy | Model |
|-------------------------|----------------|---|
| fails to break symmetry | 50% | 3-layer NN with zeros initialization |
| too large weights | 83% | 3-layer NN with large random initialization |
| recommended method | 99% | 3-layer NN with He initialization |

What you should remember from this notebook:

- · Different initializations lead to different results
- Random initialization is used to break symmetry and make sure different hidden units can learn different things
- Don't intialize to values that are too large
- · He initialization works well for networks with ReLU activations.

```
In [ ]:
```