# **Gradient Checking**

Welcome to the final assignment for this week! In this assignment you will learn to implement and use gradient checking.

You are part of a team working to make mobile payments available globally, and are asked to build a deep learning model to detect fraud--whenever someone makes a payment, you want to see if the payment might be fraudulent, such as if the user's account has been taken over by a hacker.

But backpropagation is quite challenging to implement, and sometimes has bugs. Because this is a mission-critical application, your company's CEO wants to be really certain that your implementation of backpropagation is correct. Your CEO says, "Give me a proof that your backpropagation is actually working!" To give this reassurance, you are going to use "gradient checking".

Let's do it!

```
In [1]: # Packages
   import numpy as np
   from testCases import *
    from gc_utils import sigmoid, relu, dictionary_to_vector, vector_to_dictionary, gra
   dients_to_vector
```

# 1) How does gradient checking work?

Backpropagation computes the gradients  $\frac{\partial J}{\partial \theta}$ , where  $\theta$  denotes the parameters of the model. J is computed using forward propagation and your loss function.

Because forward propagation is relatively easy to implement, you're confident you got that right, and so you're almost 100% sure that you're computing the cost J correctly. Thus, you can use your code for computing J to verify the code for computing  $\frac{\partial J}{\partial \theta}$ .

Let's look back at the definition of a derivative (or gradient):

$$\frac{\partial J}{\partial \theta} = \lim_{\varepsilon \to 0} \frac{J(\theta + \varepsilon) - J(\theta - \varepsilon)}{2\varepsilon} \tag{1}$$

If you're not familiar with the " $\lim_{\varepsilon \to 0}$ " notation, it's just a way of saying "when  $\varepsilon$  is really really small."

We know the following:

- $\frac{\partial J}{\partial \theta}$  is what you want to make sure you're computing correctly.
- You can compute  $J(\theta + \varepsilon)$  and  $J(\theta \varepsilon)$  (in the case that  $\theta$  is a real number), since you're confident your implementation for J is correct.

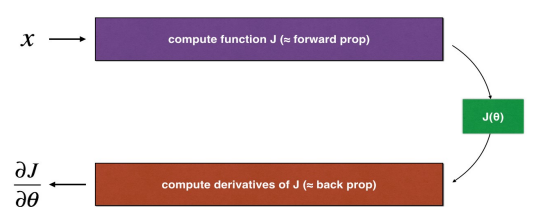
Lets use equation (1) and a small value for  $\varepsilon$  to convince your CEO that your code for computing  $\frac{\partial J}{\partial \theta}$  is correct!

# 2) 1-dimensional gradient checking

J = 8

Consider a 1D linear function  $J(\theta) = \theta x$ . The model contains only a single real-valued parameter  $\theta$ , and takes x as input.

You will implement code to compute J(.) and its derivative  $\frac{\partial J}{\partial \theta}$ . You will then use gradient checking to make sure your derivative computation for J is correct.



\*\*Figure 1\*\*: \*\*1D linear model\*\*

The diagram above shows the key computation steps: First start with x, then evaluate the function J(x) ("forward propagation"). Then compute the derivative  $\frac{\partial J}{\partial \theta}$  ("backward propagation").

**Exercise**: implement "forward propagation" and "backward propagation" for this simple function. I.e., compute both J(.) ("forward propagation") and its derivative with respect to  $\theta$  ("backward propagation"), in two separate functions.

```
In [2]: # GRADED FUNCTION: forward propagation
        def forward propagation(x, theta):
            Implement the linear forward propagation (compute J) presented in Figure 1 (J(t
        heta) = theta * x)
            Arguments:
            x -- a real-valued input
            theta -- our parameter, a real number as well
            Returns:
            J -- the value of function J, computed using the formula J(theta) = theta * x
            ### START CODE HERE ### (approx. 1 line)
            J = theta * x
            ### END CODE HERE ###
            return J
In [3]: x, theta = 2, 4
        J = forward_propagation(x, theta)
        print ("J = " + str(J))
```

### **Expected Output:**

```
 ** J ** 8
```

**Exercise**: Now, implement the backward propagation step (derivative computation) of Figure 1. That is, compute the derivative of  $J(\theta) = \theta x$  with respect to  $\theta$ . To save you from doing the calculus, you should get  $dtheta = \frac{\partial J}{\partial \theta} = x$ .

```
In [4]: # GRADED FUNCTION: backward propagation
        def backward_propagation(x, theta):
            Computes the derivative of J with respect to theta (see Figure 1).
            Arguments:
            x -- a real-valued input
            theta -- our parameter, a real number as well
            Returns:
            dtheta -- the gradient of the cost with respect to theta
            ### START CODE HERE ### (approx. 1 line)
            dtheta = x
            ### END CODE HERE ###
            return dtheta
In [5]: x, theta = 2, 4
        dtheta = backward propagation(x, theta)
        print ("dtheta = \overline{} + str(dtheta))
        dtheta = 2
```

### **Expected Output:**

\*\* dtheta \*\* 2

**Exercise**: To show that the <code>backward\_propagation()</code> function is correctly computing the gradient  $\frac{\partial J}{\partial \theta}$ , let's implement gradient checking.

#### Instructions:

- First compute "gradapprox" using the formula above (1) and a small value of  $\varepsilon$ . Here are the Steps to follow:
  - 1.  $\theta^+ = \theta + \varepsilon$
  - 2.  $\theta^- = \theta \varepsilon$
  - 3.  $J^+=J( heta^+)$
  - 4.  $J^-=J(\theta^-)$
  - 5.  $gradapprox = \frac{J^+ J^-}{2\varepsilon}$
- Then compute the gradient using backward propagation, and store the result in a variable "grad"
- Finally, compute the relative difference between "gradapprox" and the "grad" using the following formula:

$$difference = \frac{|| \operatorname{grad} - \operatorname{gradapprox} ||_{2}}{|| \operatorname{grad} ||_{2} + || \operatorname{gradapprox} ||_{2}}$$
 (2)

You will need 3 Steps to compute this formula:

- 1'. compute the numerator using np.linalg.norm(...)
- 2'. compute the denominator. You will need to call np.linalg.norm(...) twice.
- 3'. divide them.
- If this difference is small (say less than  $10^{-7}$ ), you can be quite confident that you have computed your gradient correctly. Otherwise, there may be a mistake in the gradient computation.

```
In [8]: # GRADED FUNCTION: gradient check
        def gradient_check(x, theta, epsilon = 1e-7):
            Implement the backward propagation presented in Figure 1.
            Arguments:
            x -- a real-valued input
            theta -- our parameter, a real number as well
            epsilon -- tiny shift to the input to compute approximated gradient with formul
        a(1)
            Returns:
            difference -- difference (2) between the approximated gradient and the backward
        propagation gradient
            11 11 11
            # Compute gradapprox using left side of formula (1). epsilon is small enough, y
        ou don't need to worry about the limit.
            ### START CODE HERE ### (approx. 5 lines)
            thetaplus = theta + epsilon
                                                             # Step 1
            thetaminus = theta - epsilon
                                                             # Step 2
            J plus = thetaplus * x
                                                            # Step 3
            J minus = thetaminus * x
                                                             # Step 4
            gradapprox = (J_plus - J_minus) / (2 * epsilon) # Step 5
            ### END CODE HERE ###
            # Check if gradapprox is close enough to the output of backward propagation()
            ### START CODE HERE ### (approx. 1 line)
            grad = backward propagation(x, theta)
            ### END CODE HERE ###
            ### START CODE HERE ### (approx. 1 line)
            numerator = np.linalg.norm(grad - gradapprox, ord=None, axis=None, keepdims=Tru
                   # Step 1'
            denominator = np.linalg.norm(grad, ord=None, axis=None, keepdims=True) + np.lin
        alg.norm(gradapprox, ord=None, axis=None, keepdims=True)
                                                                    # Step 2'
                                                                                # Step 3'
            difference = numerator / denominator
            ### END CODE HERE ###
            if difference < 1e-7:</pre>
                print ("The gradient is correct!")
            else:
                print ("The gradient is wrong!")
            return difference
In [9]: x, theta = 2, 4
        difference = gradient check(x, theta)
        print("difference = " + str(difference))
```

The gradient is correct! difference = 2.91933588329e-10

#### **Expected Output**: The gradient is correct!

\*\* difference \*\* 2.9193358103083e-10

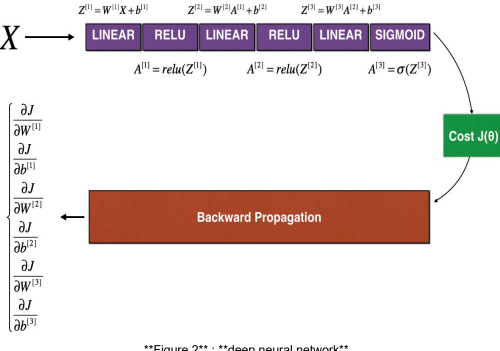
9/12/2017, 2:43 PM 5 of 12

Congrats, the difference is smaller than the  $10^{-7}$  threshold. So you can have high confidence that you've correctly computed the gradient in <code>backward propagation()</code>.

Now, in the more general case, your cost function J has more than a single 1D input. When you are training a neural network,  $\theta$  actually consists of multiple matrices  $W^{[l]}$  and biases  $b^{[l]}$ ! It is important to know how to do a gradient check with higher-dimensional inputs. Let's do it!

# 3) N-dimensional gradient checking

The following figure describes the forward and backward propagation of your fraud detection model.



\*\*Figure 2\*\*: \*\*deep neural network\*\*

\*LINEAR -> RELU -> LINEAR -> SIGMOID\*

Let's look at your implementations for forward propagation and backward propagation.

```
In [10]: def forward_propagation_n(X, Y, parameters):
             Implements the forward propagation (and computes the cost) presented in Figure
         3.
             Arguments:
             X -- training set for m examples
             Y -- labels for m examples
             parameters -- python dictionary containing your parameters "W1", "b1", "W2", "b
         2", "W3", "b3":
                             W1 -- weight matrix of shape (5, 4)
                             b1 -- bias vector of shape (5, 1)
                             W2 -- weight matrix of shape (3, 5)
                             b2 -- bias vector of shape (3, 1)
                             W3 -- weight matrix of shape (1, 3)
                             b3 -- bias vector of shape (1, 1)
             Returns:
             cost -- the cost function (logistic cost for one example)
             # retrieve parameters
             m = X.shape[1]
             W1 = parameters["W1"]
             b1 = parameters["b1"]
             W2 = parameters["W2"]
             b2 = parameters["b2"]
             W3 = parameters["W3"]
             b3 = parameters["b3"]
             \# LINEAR -> RELU -> LINEAR -> RELU -> LINEAR -> SIGMOID
             Z1 = np.dot(W1, X) + b1
             A1 = relu(Z1)
             Z2 = np.dot(W2, A1) + b2
             A2 = relu(Z2)
             Z3 = np.dot(W3, A2) + b3
             A3 = sigmoid(Z3)
             # Cost
             logprobs = np.multiply(-np.log(A3), Y) + np.multiply(-np.log(1 - A3), 1 - Y)
             cost = 1./m * np.sum(logprobs)
             cache = (Z1, A1, W1, b1, Z2, A2, W2, b2, Z3, A3, W3, b3)
             return cost, cache
```

Now, run backward propagation.

```
In [32]: def backward_propagation_n(X, Y, cache):
             Implement the backward propagation presented in figure 2.
             Arguments:
             X -- input datapoint, of shape (input size, 1)
             Y -- true "label"
             cache -- cache output from forward propagation n()
             gradients -- A dictionary with the gradients of the cost with respect to each p
         arameter, activation and pre-activation variables.
             m = X.shape[1]
             (Z1, A1, W1, b1, Z2, A2, W2, b2, Z3, A3, W3, b3) = cache
             dZ3 = A3 - Y
             dW3 = 1./m * np.dot(dZ3, A2.T)
             db3 = 1./m * np.sum(dZ3, axis=1, keepdims = True)
             dA2 = np.dot(W3.T, dZ3)
             dZ2 = np.multiply(dA2, np.int64(A2 > 0))
             dW2 = 1./m * np.dot(dZ2, A1.T) * 2
             In Order to correct it, change
             dW2 = 1./m * np.dot(dZ2, A1.T)
             db2 = 1./m * np.sum(dZ2, axis=1, keepdims = True)
             dA1 = np.dot(W2.T, dZ2)
             dZ1 = np.multiply(dA1, np.int64(A1 > 0))
             dW1 = 1./m * np.dot(dZ1, X.T)
             db1 = 4./m * np.sum(dZ1, axis=1, keepdims = True)
             In Order to correct it, change
             db1 = 1./m * np.sum(dZ1, axis=1, keepdims = True)
             11 11 11
             gradients = {"dZ3": dZ3, "dW3": dW3, "db3": db3,
                           "dA2": dA2, "dZ2": dZ2, "dW2": dW2, "db2": db2,
                           "dA1": dA1, "dZ1": dZ1, "dW1": dW1, "db1": db1}
             return gradients
```

You obtained some results on the fraud detection test set but you are not 100% sure of your model. Nobody's perfect! Let's implement gradient checking to verify if your gradients are correct.

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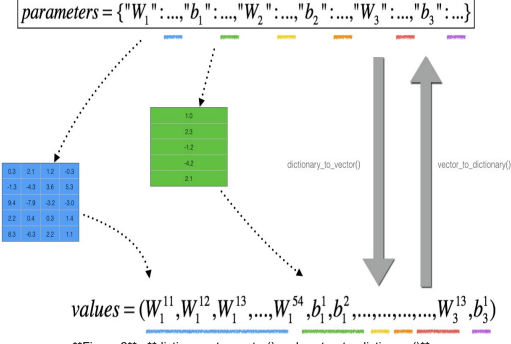
#### How does gradient checking work?.

As in 1) and 2), you want to compare "gradapprox" to the gradient computed by backpropagation. The formula is still:

$$\frac{\partial J}{\partial \theta} = \lim_{\varepsilon \to 0} \frac{J(\theta + \varepsilon) - J(\theta - \varepsilon)}{2\varepsilon} \tag{1}$$

However,  $\theta$  is not a scalar anymore. It is a dictionary called "parameters". We implemented a function "dictionary\_to\_vector()" for you. It converts the "parameters" dictionary into a vector called "values", obtained by reshaping all parameters (W1, b1, W2, b2, W3, b3) into vectors and concatenating them.

The inverse function is "vector to dictionary" which outputs back the "parameters" dictionary.



\*\*Figure 2\*\*: \*\*dictionary\_to\_vector() and vector\_to\_dictionary()\*\*
You will need these functions in gradient\_check\_n()

We have also converted the "gradients" dictionary into a vector "grad" using gradients\_to\_vector(). You don't need to worry about that.

**Exercise**: Implement gradient\_check\_n().

Instructions: Here is pseudo-code that will help you implement the gradient check.

For each i in num parameters:

- To compute J plus[i]:
  - 1. Set  $heta^+$  to np.copy(parameters\_values)
  - 2. Set  $\theta_i^+$  to  $\theta_i^+ + \varepsilon$
  - 3. Calculate  $J_i^{t+}$  using to forward\_propagation\_n(x, y, vector\_to\_dictionary( $heta^+$ )).
- ullet To compute <code>J\_minus[i]</code>: do the same thing with  $heta^-$
- ullet Compute  $gradapprox[i] = rac{J_i^+ J_i^-}{2arepsilon}$

Thus, you get a vector gradapprox, where gradapprox[i] is an approximation of the gradient with respect to parameter\_values[i]. You can now compare this gradapprox vector to the gradients vector from backpropagation. Just like for the 1D case (Steps 1', 2', 3'), compute:

$$difference = \frac{\|grad - gradapprox\|_2}{\|grad\|_2 + \|gradapprox\|_2}$$
(3)

10 of 12

```
In [33]: # GRADED FUNCTION: gradient check n
         def gradient_check_n(parameters, gradients, X, Y, epsilon = 1e-7):
             Checks if backward propagation n computes correctly the gradient of the cost ou
         tput by forward propagation n
             Arguments:
             parameters -- python dictionary containing your parameters "W1", "b1", "W2", "b
         2", "W3", "b3":
             grad -- output of backward propagation n, contains gradients of the cost with r
         espect to the parameters.
             x -- input datapoint, of shape (input size, 1)
             y -- true "label"
             epsilon -- tiny shift to the input to compute approximated gradient with formul
         a(1)
             Returns:
             difference -- difference (2) between the approximated gradient and the backward
         propagation gradient
             # Set-up variables
             parameters_values, _ = dictionary_to_vector(parameters)
             grad = gradients_to_vector(gradients)
             num_parameters = parameters_values.shape[0]
             J plus = np.zeros((num parameters, 1))
             J_minus = np.zeros((num_parameters, 1))
             gradapprox = np.zeros((num parameters, 1))
             # Compute gradapprox
             for i in range(num parameters):
                 # Compute J plus[i]. Inputs: "parameters values, epsilon". Output = "J plus
         [i]".
                 \# " " is used because the function you have to outputs two parameters but w
         e only care about the first one
                 ### START CODE HERE ### (approx. 3 lines)
                                                                       # Step 1
                 thetaplus = np.copy(parameters values)
                 thetaplus[i][0] = thetaplus[i][0] + epsilon
                                                                        # Step 2
                 J_plus[i], _ = forward_propagation_n(X, Y, vector_to_dictionary(thetaplus))
         # Step 3
                 ### END CODE HERE ###
                 # Compute J minus[i]. Inputs: "parameters values, epsilon". Output = "J min
         us[i]".
                 ### START CODE HERE ### (approx. 3 lines)
                                                                       # Step 1
                 thetaminus = np.copy(parameters values)
                 thetaminus[i][0] = thetaminus[i][0] - epsilon
                                                                       # Step 2
                 J_minus[i], _ = forward propagation_n(X, Y, vector_to_dictionary(thetaminus
         )) # Step 3
                 ### END CODE HERE ###
                 # Compute gradapprox[i]
                 ### START CODE HERE ### (approx. 1 line)
                 gradapprox[i] = (J_plus[i] - J_minus[i]) / (2 * epsilon)
                 ### END CODE HERE ###
             # Compare gradapprox to backward propagation gradients by computing difference.
             ### START CODE HERE ### (approx. 1 line)
             numerator = np.linalg.norm(grad - gradapprox, ord=None, axis=None, keepdims=Fal
             denominator = np.linalg.norm(grad, ord=None, axis=None, keepdims=False) + np.li
         nalg.norm(gradapprox. ord=None. axis=None. keepdims=False) # Step 2'
```

```
In [34]: X, Y, parameters = gradient_check_n_test_case()

cost, cache = forward_propagation_n(X, Y, parameters)
gradients = backward_propagation_n(X, Y, cache)
difference = gradient_check_n(parameters, gradients, X, Y)

There is a mistake in the backward propagation! difference = 0.285093156781
```

#### **Expected output:**

It seems that there were errors in the <code>backward\_propagation\_n</code> code we gave you! Good that you've implemented the gradient check. Go back to <code>backward\_propagation</code> and try to find/correct the errors (Hint: check dW2 and db1). Rerun the gradient check when you think you've fixed it. Remember you'll need to re-execute the cell defining <code>backward\_propagation\_n()</code> if you modify the code.

Can you get gradient check to declare your derivative computation correct? Even though this part of the assignment isn't graded, we strongly urge you to try to find the bug and re-run gradient check until you're convinced backprop is now correctly implemented.

#### Note

- Gradient Checking is slow! Approximating the gradient with  $\frac{\partial J}{\partial \theta} \approx \frac{J(\theta+\varepsilon)-J(\theta-\varepsilon)}{2\varepsilon}$  is computationally costly. For this reason, we don't run gradient checking at every iteration during training. Just a few times to check if the gradient is correct.
- Gradient Checking, at least as we've presented it, doesn't work with dropout. You would usually run the gradient check algorithm without dropout to make sure your backprop is correct, then add dropout.

Congrats, you can be confident that your deep learning model for fraud detection is working correctly! You can even use this to convince your CEO. :)

#### What you should remember from this notebook:

- Gradient checking verifies closeness between the gradients from backpropagation and the numerical approximation of the gradient (computed using forward propagation).
- Gradient checking is slow, so we don't run it in every iteration of training. You would usually run it only to make sure your code is correct, then turn it off and use backprop for the actual learning process.

```
In [ ]:
```