Building your Recurrent Neural Network - Stepby Step

Welcome to Course 5's first assignment, where you'll be implementing key components of a Recurrent Neural Network, or RNN, in NumPy!

By the end of this assignment, you'll be able to:

- Define notation for building sequence models
- · Describe the architecture of a basic RNN
- Identify the main components of an LSTM
- Implement backpropagation through time for a basic RNN and an LSTM
- Give examples of several types of RNN

Recurrent Neural Networks (RNN) are very effective for Natural Language Processing and other sequence tasks because they have "memory." They can read inputs $x^{\langle t \rangle}$ (such as words) one at a time, and remember some contextual information through the hidden layer activations that get passed from one time step to the next. This allows a unidirectional (oneway) RNN to take information from the past to process later inputs. A bidirectional (two-way) RNN can take context from both the past and the future, much like Marty McFly.

Notation:

- Superscript [l] denotes an object associated with the l^{th} layer.
- Superscript (i) denotes an object associated with the i^{th} example.
- Superscript $\langle t \rangle$ denotes an object at the t^{th} time step.
- Subscript i denotes the i^{th} entry of a vector.

Example:

• $a_5^{(2)[3]<4>}$ denotes the activation of the 2nd training example (2), 3rd layer [3], 4th time step <4>, and 5th entry in the vector.

Pre-requisites

- You should already be familiar with numpy
- To refresh your knowledge of numpy, you can review course 1 of the specialization "Neural Networks and Deep Learning":
 - Specifically, review the week 2's practice assignment "Python Basics with Numpy (optional assignment)" (https://www.coursera.org/learn/neural-networks-deep-learning/programming/isoAV/python-basics-with-numpy)

Be careful when modifying the starter code!

When working on graded functions, please remember to only modify the code that is

between:

```
#### START CODE HERE
```

and:

```
#### END CODE HERE
```

• In particular, avoid modifying the first line of graded routines. These start with:

```
# GRADED FUNCTION: routine_name
```

The automatic grader (autograder) needs these to locate the function - so even a change in spacing will cause issues with the autograder, returning 'failed' if any of these are modified or missing. Now, let's get started!

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Packages

```
In [1]: import numpy as np
from rnn_utils import *
from public_tests import *
```

1 - Forward Propagation for the Basic Recurrent Neural Network

Later this week, you'll get a chance to generate music using an RNN! The basic RNN that you'll implement has the following structure:

In this example, $T_x = T_y$.

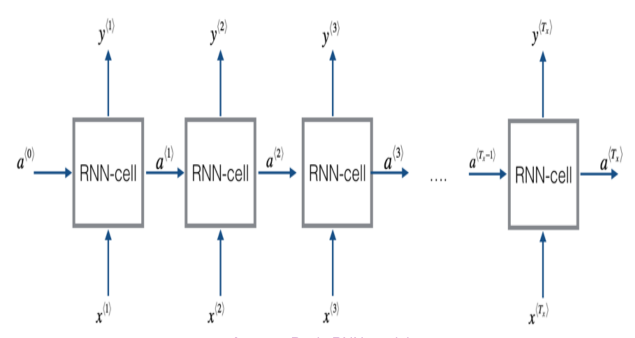


Figure 1: Basic RNN model

Dimensions of input x

Input with n_x number of units

- For a single time step of a single input example, $x^{(i)\langle t\rangle}$ is a one-dimensional input vector
- Using language as an example, a language with a 5000-word vocabulary could be one-hot encoded into a vector that has 5000 units. So $x^{(i)\langle t\rangle}$ would have the shape (5000,)
- The notation n_x is used here to denote the number of units in a single time step of a single training example

Time steps of size T_x

- A recurrent neural network has multiple time steps, which you'll index with t.
- In the lessons, you saw a single training example $x^{(i)}$ consisting of multiple time steps T_x . In this notebook, T_x will denote the number of timesteps in the longest sequence.

Batches of size m

- Let's say we have mini-batches, each with 20 training examples
- To benefit from vectorization, you'll stack 20 columns of $x^{(i)}$ examples
- For example, this tensor has the shape (5000,20,10)
- You'll use *m* to denote the number of training examples
- So, the shape of a mini-batch is (n_x, m, T_x)

3D Tensor of shape (n_x, m, T_x)

• The 3-dimensional tensor x of shape (n_x, m, T_x) represents the input x that is fed into the RNN

Taking a 2D slice for each time step: $x^{\langle t \rangle}$

- At each time step, you'll use a mini-batch of training examples (not just a single example)
- So, for each time step t, you'll use a 2D slice of shape (n_x, m)
- This 2D slice is referred to as $x^{\langle t \rangle}$. The variable name in the code is xt.

Definition of hidden state a

• The activation $a^{\langle t \rangle}$ that is passed to the RNN from one time step to another is called a "hidden state."

Dimensions of hidden state a

- Similar to the input tensor x, the hidden state for a single training example is a vector of length n_a
- If you include a mini-batch of m training examples, the shape of a mini-batch is (n_a, m)
- When you include the time step dimension, the shape of the hidden state is (n_a, m, T_x)
- You'll loop through the time steps with index t, and work with a 2D slice of the 3D tensor
- This 2D slice is referred to as $a^{(t)}$
- In the code, the variable names used are either a_prev or a_next, depending on the function being implemented
- The shape of this 2D slice is (n_a, m)

Dimensions of prediction \hat{y}

- Similar to the inputs and hidden states, \hat{y} is a 3D tensor of shape (n_v, m, T_v)
 - n_y : number of units in the vector representing the prediction
 - *m*: number of examples in a mini-batch
 - T_y : number of time steps in the prediction
- For a single time step t, a 2D slice $\hat{y}^{\langle t \rangle}$ has shape (n_v, m)
- In the code, the variable names are:
 - y_pred: \hat{y} • yt_pred: $\hat{y}^{\langle t \rangle}$

Here's how you can implement an RNN:

Steps:

- 1. Implement the calculations needed for one time step of the RNN.
- 2. Implement a loop over T_x time steps in order to process all the inputs, one at a time.

1.1 - RNN Cell

You can think of the recurrent neural network as the repeated use of a single cell. First, you'll implement the computations for a single time step. The following figure describes the operations for a single time step of an RNN cell:

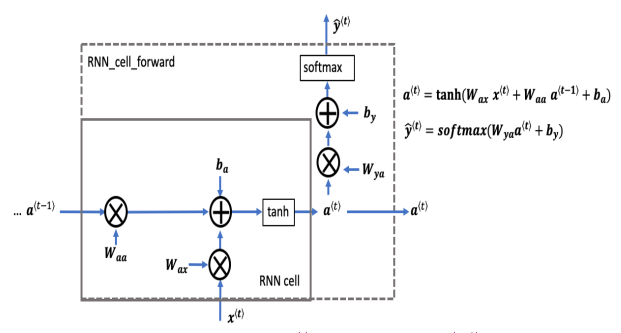


Figure 2: Basic RNN cell. Takes as input $x^{\langle t \rangle}$ (current input) and $a^{\langle t-1 \rangle}$ (previous hidden state containing information from the past), and outputs $a^{\langle t \rangle}$ which is given to the next RNN cell and also used to predict $\hat{y}^{\langle t \rangle}$

RNN cell versus RNN_cell_forward:

- Note that an RNN cell outputs the hidden state $a^{\langle t \rangle}$.
 - RNN cell is shown in the figure as the inner box with solid lines
- The function that you'll implement, <code>rnn_cell_forward</code> , also calculates the prediction $\hat{y}^{\langle t \rangle}$
 - RNN_cell_forward is shown in the figure as the outer box with dashed lines

Exercise 1 - rnn cell forward

Implement the RNN cell described in Figure 2.

Instructions:

1. Compute the hidden state with tanh activation:

$$a^{\langle t \rangle} = \tanh(W_{aa}a^{\langle t-1 \rangle} + W_{ax}x^{\langle t \rangle} + b_a)$$

- 2. Using your new hidden state $a^{\langle t \rangle}$, compute the prediction $\hat{y}^{\langle t \rangle} = softmax(W_{ya}a^{\langle t \rangle} + b_y)$. (The function softmax is provided)
- 3. Store $(a^{\langle t \rangle}, a^{\langle t-1 \rangle}, x^{\langle t \rangle}, parameters)$ in a cache
- 4. Return $a^{\langle t \rangle}$, $\hat{y}^{\langle t \rangle}$ and cache

Additional Hints

- A little more information on <u>numpy.tanh</u> (https://numpy.org/devdocs/reference/generated/numpy.tanh.html)
- In this assignment, there's an existing softmax function for you to use. It's located in the file 'rnn_utils.py' and has already been imported.
- For matrix multiplication, use <u>numpy.dot</u>
 (https://docs.scipy.org/doc/numpy/reference/generated/numpy.dot.html)

```
In [2]: # UNQ C1 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
        # GRADED FUNCTION: rnn_cell_forward
        def rnn_cell_forward(xt, a_prev, parameters):
            Implements a single forward step of the RNN-cell as described in F
            Arguments:
            xt -- your input data at timestep "t", numpy array of shape (n_x,
            a_prev -- Hidden state at timestep "t-1", numpy array of shape (n_
            parameters -- python dictionary containing:
                                Wax -- Weight matrix multiplying the input, nu
                                Waa -- Weight matrix multiplying the hidden st
                                Wya -- Weight matrix relating the hidden-state
                                ba -- Bias, numpy array of shape (n_a, 1)
                                by -- Bias relating the hidden-state to the ou
            Returns:
            a_next -- next hidden state, of shape (n_a, m)
            yt_pred -- prediction at timestep "t", numpy array of shape (n_y,
            cache -- tuple of values needed for the backward pass, contains (a
            # Retrieve parameters from "parameters"
            Wax = parameters["Wax"]
            Waa = parameters["Waa"]
            Wya = parameters["Wya"]
            ba = parameters["ba"]
            by = parameters["by"]
            ### START CODE HERE ### (≈2 lines)
            # compute next activation state using the formula given above
            a next = np.tanh(np.dot(Wax, xt) + np.dot(Waa, a prev) + ba)
            # compute output of the current cell using the formula given above
            vt pred = softmax(np.dot(Wya, a next) + by)
            ### END CODE HERE ###
            # store values you need for backward propagation in cache
            cache = (a_next, a_prev, xt, parameters)
```

return a_next, yt_pred, cache

```
In [3]: | np.random.seed(1)
        xt_tmp = np.random.randn(3, 10)
        a_prev_tmp = np.random.randn(5, 10)
        parameters_tmp = {}
        parameters_tmp['Waa'] = np.random.randn(5, 5)
        parameters_tmp['Wax'] = np.random.randn(5, 3)
        parameters tmp['Wya'] = np.random.randn(2, 5)
        parameters_tmp['ba'] = np.random.randn(5, 1)
        parameters tmp['by'] = np.random.randn(2, 1)
        a_next_tmp, yt_pred_tmp, cache_tmp = rnn_cell_forward(xt_tmp, a_prev_t
        print("a_next[4] = \n", a_next_tmp[4])
        print("a_next.shape = \n", a_next_tmp.shape)
        print("yt_pred[1] =\n", yt_pred_tmp[1])
        print("yt_pred.shape = \n", yt_pred_tmp.shape)
        # UNIT TESTS
        rnn_cell_forward_tests(rnn_cell_forward)
        a next[4] =
         [ 0.59584544  0.18141802  0.61311866  0.99808218  0.85016201  0.9998
        0978
         -0.18887155 0.99815551 0.6531151
                                              0.828720371
        a next.shape =
         (5, 10)
        yt_pred[1] =
         [0.9888161 0.01682021 0.21140899 0.36817467 0.98988387 0.88945212
         0.36920224 0.9966312 0.9982559 0.17746526]
        vt pred.shape =
         (2, 10)
        All tests passed
        Expected Output:
           a next[4] =
            [ 0.59584544  0.18141802  0.61311866  0.99808218  0.85016201
           0.99980978
            -0.18887155 0.99815551 0.6531151
                                                 0.828720371
           a_next.shape =
            (5, 10)
           yt_pred[1] =
             [ 0.9888161
                          0.01682021 0.21140899 0.36817467 0.98988387
           0.88945212
             0.36920224 0.9966312 0.9982559 0.177465261
           yt_pred.shape =
            (2, 10)
```

1.2 - RNN Forward Pass

- A recurrent neural network (RNN) is a repetition of the RNN cell that you've just built.
 - If your input sequence of data is 10 time steps long, then you will re-use the RNN cell 10 times
- Each cell takes two inputs at each time step:
 - $a^{(t-1)}$: The hidden state from the previous cell
 - $x^{(t)}$: The current time step's input data
- It has two outputs at each time step:
 - A hidden state $(a^{\langle t \rangle})$
 - A prediction $(y^{\langle t \rangle})$
- The weights and biases $(W_{aa}, b_a, W_{ax}, b_x)$ are re-used each time step
 - They are maintained between calls to rnn_cell_forward in the 'parameters' dictionary

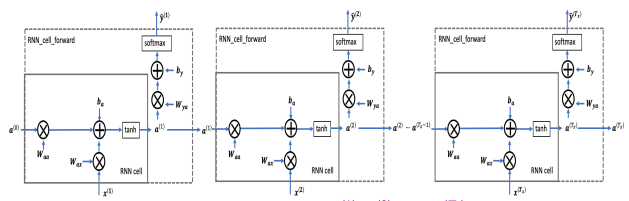


Figure 3: Basic RNN. The input sequence $x=(x^{\langle 1 \rangle},x^{\langle 2 \rangle},\ldots,x^{\langle T_x \rangle})$ is carried over T_x time steps. The network outputs $y=(y^{\langle 1 \rangle},y^{\langle 2 \rangle},\ldots,y^{\langle T_x \rangle})$.

Exercise 2 - rnn forward

Implement the forward propagation of the RNN described in Figure 3.

Instructions:

- Create a 3D array of zeros, a of shape (n_a, m, T_x) that will store all the hidden states computed by the RNN
- Create a 3D array of zeros, \hat{y} , of shape (n_y, m, T_x) that will store the predictions
 - Note that in this case, $T_y = T_x$ (the prediction and input have the same number of time steps)
- Initialize the 2D hidden state a_next by setting it equal to the initial hidden state, a_0
- At each time step *t*:
 - Get $x^{\langle t \rangle}$, which is a 2D slice of x for a single time step t
 - $x^{\langle t \rangle}$ has shape (n_x, m)
 - x has shape (n_x, m, T_x)
 - Update the 2D hidden state $a^{\langle t \rangle}$ (variable name a_next), the prediction $\hat{y}^{\langle t \rangle}$ and the cache by running rnn_cell_forward
 - $a^{\langle t \rangle}$ has shape (n_a, m)
 - Store the 2D hidden state in the 3D tensor a, at the t^{th} position
 - a has shape (n_a, m, T_x)
 - Store the 2D $\hat{y}^{\langle t \rangle}$ prediction (variable name yt_pred) in the 3D tensor \hat{y}_{pred} at the t^{th} position
 - $\hat{y}^{\langle t \rangle}$ has shape (n_v, m)
 - \hat{y} has shape (n_y, m, T_x)
 - Append the cache to the list of caches
- Return the 3D tensor a and \hat{y} , as well as the list of caches

Additional Hints

- Some helpful documentation on <u>np.zeros</u>
 (https://docs.scipy.org/doc/numpy/reference/generated/numpy.zeros.html)
- If you have a 3 dimensional numpy array and are indexing by its third dimension, you can use array slicing like this: var_name[:,:,i]

```
parameters -- python dictionary containing:
                    Waa -- Weight matrix multiplying the hidden st
                    Wax -- Weight matrix multiplying the input, nu
                    Wya -- Weight matrix relating the hidden-state
                    ba -- Bias numpy array of shape (n_a, 1)
                    by -- Bias relating the hidden-state to the ou
Returns:
a -- Hidden states for every time-step, numpy array of shape (n_a,
y_pred -- Predictions for every time-step, numpy array of shape (n
caches -- tuple of values needed for the backward pass, contains (
# Initialize "caches" which will contain the list of all caches
caches = []
# Retrieve dimensions from shapes of x and parameters["Wya"]
n_x, m, T_x = x. shape
n_y, n_a = parameters["Wya"].shape
### START CODE HERE ###
# initialize "a" and "y_pred" with zeros (≈2 lines)
a = np.zeros((n a,m,T x))
y_pred = np.zeros((n_y,m,T_x))
# Initialize a_next (≈1 line)
a_next = a0
# loop over all time-steps
for t in range(T x):
    # Update next hidden state, compute the prediction, get the ca
    a_next, yt_pred, cache = rnn_cell_forward(x[:,:,t], a_next, pa
    # Save the value of the new "next" hidden state in a (≈1 line)
    a[:,:,t] = a_next
    # Save the value of the prediction in y (≈1 line)
   y_pred[:,:,t] = yt_pred
    # Append "cache" to "caches" (≈1 line)
    caches.append(cache)
### END CODE HERE ###
# store values needed for backward propagation in cache
caches = (caches, x)
return a, y_pred, caches
```

```
In [5]: | np.random.seed(1)
        x_{tmp} = np.random.randn(3, 10, 4)
        a0\_tmp = np.random.randn(5, 10)
        parameters_tmp = {}
        parameters_tmp['Waa'] = np.random.randn(5, 5)
        parameters_tmp['Wax'] = np.random.randn(5, 3)
        parameters tmp['Wya'] = np.random.randn(2, 5)
        parameters tmp['ba'] = np.random.randn(5, 1)
        parameters tmp['by'] = np.random.randn(2, 1)
        a_tmp, y_pred_tmp, caches_tmp = rnn_forward(x_tmp, a0_tmp, parameters
        print("a[4][1] = \n", a_tmp[4][1])
        print("a.shape = \n", a_tmp.shape)
        print("y_pred[1][3] =\n", y_pred_tmp[1][3])
        print("y_pred.shape = \n", y_pred_tmp.shape)
        print("caches[1][1][3] =\n", caches_tmp[1][1][3])
        print("len(caches) = \n", len(caches_tmp))
        #UNIT TEST
        rnn_forward_test(rnn_forward)
        a[4][1] =
         [-0.99999375 \quad 0.77911235 \quad -0.99861469 \quad -0.99833267]
        a.shape =
         (5, 10, 4)
        y \text{ pred}[1][3] =
         [0.79560373 0.86224861 0.11118257 0.81515947]
        y_pred.shape =
         (2, 10, 4)
        caches[1][1][3] =
         [-1.1425182 -0.34934272 -0.20889423 0.58662319]
        len(caches) =
         2
        All tests passed
        Expected Output:
           a[4][1] =
            [-0.99999375 0.77911235 -0.99861469 -0.99833267]
           a.shape =
            (5, 10, 4)
           y_pred[1][3] =
            [ 0.79560373  0.86224861  0.11118257  0.81515947]
           y_pred.shape =
            (2, 10, 4)
           caches[1][1][3] =
            len(caches) =
```

2

Congratulations!

You've successfully built the forward propagation of a recurrent neural network from scratch. Nice work!

Situations when this RNN will perform better:

- This will work well enough for some applications, but it suffers from vanishing gradients.
- The RNN works best when each output $\hat{y}^{(t)}$ can be estimated using "local" context.
- "Local" context refers to information that is close to the prediction's time step *t*.
- More formally, local context refers to inputs $x^{\langle t' \rangle}$ and predictions $\hat{y}^{\langle t \rangle}$ where t' is close to t.

What you should remember:

- The recurrent neural network, or RNN, is essentially the repeated use of a single cell.
- A basic RNN reads inputs one at a time, and remembers information through the hidden layer activations (hidden states) that are passed from one time step to the next.
 - The time step dimension determines how many times to re-use the RNN cell
- Each cell takes two inputs at each time step:
 - The hidden state from the previous cell
 - The current time step's input data
- Each cell has two outputs at each time step:
 - A hidden state
 - A prediction

In the next section, you'll build a more complex model, the LSTM, which is better at addressing vanishing gradients. The LSTM is better able to remember a piece of information and save it for many time steps.

2 - Long Short-Term Memory (LSTM) Network

The following figure shows the operations of an LSTM cell:

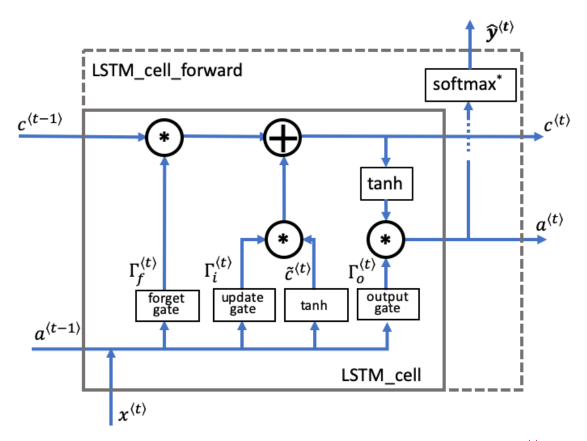


Figure 4: LSTM cell. This tracks and updates a "cell state," or memory variable $c^{\langle t \rangle}$ at every time step, which can be different from $a^{\langle t \rangle}$. Note, the softmax includes a dense layer and softmax.

Similar to the RNN example above, you'll begin by implementing the LSTM cell for a single time step. Then, you'll iteratively call it from inside a "for loop" to have it process an input with T_x time steps.

Overview of gates and states

Forget gate Γ_f

- Let's assume you are reading words in a piece of text, and plan to use an LSTM to keep track of grammatical structures, such as whether the subject is singular ("puppy") or plural ("puppies").
- If the subject changes its state (from a singular word to a plural word), the memory of the previous state becomes outdated, so you'll "forget" that outdated state.
- The "forget gate" is a tensor containing values between 0 and 1.
 - If a unit in the forget gate has a value close to 0, the LSTM will "forget" the stored

state in the corresponding unit of the previous cell state.

• If a unit in the forget gate has a value close to 1, the LSTM will mostly remember the corresponding value in the stored state.

Equation

$$\mathbf{\Gamma}_f^{\langle t \rangle} = \sigma(\mathbf{W}_f[\mathbf{a}^{\langle t-1 \rangle}, \mathbf{x}^{\langle t \rangle}] + \mathbf{b}_f)$$
 (1)

Explanation of the equation:

- ullet W_f contains weights that govern the forget gate's behavior.
- The previous time step's hidden state $[a^{\langle t-1\rangle}]$ and current time step's input $x^{\langle t\rangle}$ are concatenated together and multiplied by $\mathbf{W_f}$.
- A sigmoid function is used to make each of the gate tensor's values $\Gamma_f^{\langle t \rangle}$ range from 0 to 1.
- The forget gate $\Gamma_f^{\langle t \rangle}$ has the same dimensions as the previous cell state $c^{\langle t-1 \rangle}$.
- This means that the two can be multiplied together, element-wise.
- Multiplying the tensors $\Gamma_f^{\langle t \rangle} * \mathbf{c}^{\langle t-1 \rangle}$ is like applying a mask over the previous cell state.
- If a single value in $\Gamma_f^{\langle t \rangle}$ is 0 or close to 0, then the product is close to 0.
 - This keeps the information stored in the corresponding unit in $\mathbf{c}^{\langle t-1 \rangle}$ from being remembered for the next time step.
- Similarly, if one value is close to 1, the product is close to the original value in the previous cell state.
 - The LSTM will keep the information from the corresponding unit of $\mathbf{c}^{\langle t-1 \rangle}$, to be used in the next time step.

Variable names in the code

The variable names in the code are similar to the equations, with slight differences.

ullet Wf : forget gate weight \mathbf{W}_f

• bf : forget gate bias \mathbf{b}_f

• ft : forget gate $\Gamma_f^{\langle t \rangle}$

Candidate value $\tilde{\mathbf{c}}^{\langle t \rangle}$

- The candidate value is a tensor containing information from the current time step that **may** be stored in the current cell state $\mathbf{c}^{\langle t \rangle}$.
- The parts of the candidate value that get passed on depend on the update gate.
- The candidate value is a tensor containing values that range from -1 to 1.
- The tilde "~" is used to differentiate the candidate $\tilde{\mathbf{c}}^{\langle t \rangle}$ from the cell state $\mathbf{c}^{\langle t \rangle}$.

Equation

$$\tilde{\mathbf{c}}^{\langle t \rangle} = \tanh(\mathbf{W}_c[\mathbf{a}^{\langle t-1 \rangle}, \mathbf{x}^{\langle t \rangle}] + \mathbf{b}_c)$$
(3)

Explanation of the equation

• The tanh function produces values between -1 and 1.

Variable names in the code

• cct : candidate value $\tilde{\mathbf{c}}^{\langle t \rangle}$

Update gate Γ_i

- You use the update gate to decide what aspects of the candidate $\tilde{\mathbf{c}}^{(t)}$ to add to the cell state $c^{\langle t \rangle}$.
- The update gate decides what parts of a "candidate" tensor $\tilde{\mathbf{c}}^{\langle t \rangle}$ are passed onto the cell state $\mathbf{c}^{\langle t \rangle}$.
- The update gate is a tensor containing values between 0 and 1.
 - When a unit in the update gate is close to 1, it allows the value of the candidate $\tilde{\mathbf{c}}^{\langle t \rangle}$ to be passed onto the hidden state $\mathbf{c}^{\langle t \rangle}$
 - When a unit in the update gate is close to 0, it prevents the corresponding value in the candidate from being passed onto the hidden state.
- Notice that the subscript "i" is used and not "u", to follow the convention used in the literature.

Equation

$$\mathbf{\Gamma}_{i}^{\langle t \rangle} = \sigma(\mathbf{W}_{i}[a^{\langle t-1 \rangle}, \mathbf{x}^{\langle t \rangle}] + \mathbf{b}_{i})$$
 (2)

Explanation of the equation

- Similar to the forget gate, here $\mathbf{\Gamma}_i^{\langle t \rangle}$, the sigmoid produces values between 0 and 1.
- The update gate is multiplied element-wise with the candidate, and this product ($\Gamma_i^{\langle t \rangle} * \tilde{c}^{\langle t \rangle}$) is used in determining the cell state $\mathbf{c}^{\langle t \rangle}$.

Variable names in code (Please note that they're different than the equations)

In the code, you'll use the variable names found in the academic literature. These variables don't use "u" to denote "update".

- Wi is the update gate weight W_i (not "Wu")
- bi is the update gate bias \mathbf{b}_i (not "bu")
- it is the forget gate $\Gamma_i^{\langle t \rangle}$ (not "ut")

Cell state $\mathbf{c}^{\langle t \rangle}$

- The cell state is the "memory" that gets passed onto future time steps.
- The new cell state $\mathbf{c}^{\langle t \rangle}$ is a combination of the previous cell state and the candidate value.

Equation

$$\mathbf{c}^{\langle t \rangle} = \mathbf{\Gamma}_f^{\langle t \rangle} * \mathbf{c}^{\langle t-1 \rangle} + \mathbf{\Gamma}_i^{\langle t \rangle} * \tilde{\mathbf{c}}^{\langle t \rangle}$$
(4)

Explanation of equation

- ullet The previous cell state ${f c}^{\langle t-1
 angle}$ is adjusted (weighted) by the forget gate ${f \Gamma}_f^{\langle t
 angle}$
- and the candidate value $\tilde{\mathbf{c}}^{\langle t \rangle}$, adjusted (weighted) by the update gate $\mathbf{\Gamma}_i^{\langle t \rangle}$

Variable names and shapes in the code

- c : cell state, including all time steps, c shape (n_a, m, T_x)
- c_next : new (next) cell state, $\mathbf{c}^{\langle t \rangle}$ shape (n_a, m)
- c_prev: previous cell state, $\mathbf{c}^{(t-1)}$, shape (n_a, m)

Output gate Γ_o

- The output gate decides what gets sent as the prediction (output) of the time step.
- The output gate is like the other gates, in that it contains values that range from 0 to 1.

Equation

$$\mathbf{\Gamma}_o^{\langle t \rangle} = \sigma(\mathbf{W}_o[\mathbf{a}^{\langle t-1 \rangle}, \mathbf{x}^{\langle t \rangle}] + \mathbf{b}_o) \tag{5}$$

Explanation of the equation

- The output gate is determined by the previous hidden state ${\bf a}^{\langle t-1 \rangle}$ and the current input ${\bf x}^{\langle t \rangle}$
- The sigmoid makes the gate range from 0 to 1.

Variable names in the code

ullet Wo : output gate weight, W_o

• bo : output gate bias, b_o

• ot : output gate, $\Gamma_a^{\langle t \rangle}$

Hidden state $\mathbf{a}^{\langle t \rangle}$

- The hidden state gets passed to the LSTM cell's next time step.
- It is used to determine the three gates $(\Gamma_f, \Gamma_u, \Gamma_o)$ of the next time step.
- The hidden state is also used for the prediction $v^{\langle t \rangle}$.

Equation

$$\mathbf{a}^{\langle t \rangle} = \mathbf{\Gamma}_o^{\langle t \rangle} * \tanh(\mathbf{c}^{\langle t \rangle}) \tag{6}$$

Explanation of equation

- The hidden state ${\bf a}^{\langle t \rangle}$ is determined by the cell state ${\bf c}^{\langle t \rangle}$ in combination with the output gate Γ_o .
- The cell state state is passed through the tanh function to rescale values between -1 and 1.
- The output gate acts like a "mask" that either preserves the values of $\tanh(\mathbf{c}^{\langle t \rangle})$ or keeps those values from being included in the hidden state $\mathbf{a}^{\langle t \rangle}$

Variable names and shapes in the code

- a : hidden state, including time steps. **a** has shape (n_a, m, T_x)
- a_prev: hidden state from previous time step. $\mathbf{a}^{(t-1)}$ has shape (n_a, m)
- a_next : hidden state for next time step. $\mathbf{a}^{\langle t \rangle}$ has shape (n_a, m)

Prediction $\mathbf{y}_{pred}^{\langle t \rangle}$

• The prediction in this use case is a classification, so you'll use a softmax.

The equation is:

$$\mathbf{y}_{pred}^{\langle t \rangle} = \operatorname{softmax}(\mathbf{W}_{y} \mathbf{a}^{\langle t \rangle} + \mathbf{b}_{y})$$

Variable names and shapes in the code

- y_pred : prediction, including all time steps. \mathbf{y}_{pred} has shape (n_y, m, T_x) . Note that $(T_y = T_x)$ for this example.
- yt_pred : prediction for the current time step t. $\mathbf{y}_{pred}^{\langle t \rangle}$ has shape (n_y, m)

2.1 - LSTM Cell

Exercise 3 - Istm cell forward

Implement the LSTM cell described in Figure 4.

Instructions:

1. Concatenate the hidden state $a^{(t-1)}$ and input $x^{(t)}$ into a single matrix:

$$concat = \begin{bmatrix} a^{\langle t-1 \rangle} \\ \chi^{\langle t \rangle} \end{bmatrix}$$

- 2. Compute all formulas (1 through 6) for the gates, hidden state, and cell state.
- 3. Compute the prediction $y^{\langle t \rangle}$.

Additional Hints

- You can use <u>numpy.concatenate</u> (https://docs.scipy.org/doc/numpy/reference/generated/numpy.concatenate.html).
 Check which value to use for the axis parameter.
- The functions sigmoid() and softmax are imported from rnn_utils.py.
- Some docs for <u>numpy.tanh</u>
 (https://docs.scipy.org/doc/numpy/reference/generated/numpy.tanh.html)
- Use numpy.dot.html) for matrix multiplication.
- Notice that the variable names Wi, bi refer to the weights and biases of the **update** gate. There are no variables named "Wu" or "bu" in this function.

```
bc -- Bias of the first "tanh", numpy array of
                    Wo — Weight matrix of the output gate, numpy
                    bo -- Bias of the output gate, numpy array of
                    Wy -- Weight matrix relating the hidden-state
                    by -- Bias relating the hidden-state to the ou
Returns:
a_next -- next hidden state, of shape (n_a, m)
c_next -- next memory state, of shape (n_a, m)
yt_pred -- prediction at timestep "t", numpy array of shape (n_y,
cache -- tuple of values needed for the backward pass, contains (a
Note: ft/it/ot stand for the forget/update/output gates, cct stand
      c stands for the cell state (memory)
.....
# Retrieve parameters from "parameters"
Wf = parameters["Wf"] # forget gate weight
bf = parameters["bf"]
Wi = parameters["Wi"] # update gate weight (notice the variable na
bi = parameters["bi"] # (notice the variable name)
Wc = parameters["Wc"] # candidate value weight
bc = parameters["bc"]
Wo = parameters["Wo"] # output gate weight
bo = parameters["bo"]
Wy = parameters["Wy"] # prediction weight
by = parameters["by"]
# Retrieve dimensions from shapes of xt and Wy
n_x, m = xt.shape
n_y, n_a = Wy.shape
### START CODE HERE ###
# Concatenate a prev and xt (≈1 line)
concat = np.concatenate((a_prev,xt),axis=0)
# Compute values for ft, it, cct, c_next, ot, a_next using the for
ft = sigmoid(np.dot(Wf,concat)+bf)
it = sigmoid(np.dot(Wi,concat)+bi)
cct = np.tanh(np.dot(Wc,concat)+bc)
c_next = it*cct + ft*c_prev
ot = sigmoid(np.dot(Wo,concat)+bo)
a_next = ot*np.tanh(c_next)
# Compute prediction of the LSTM cell (≈1 line)
yt_pred = softmax(np.dot(Wy,a_next)+by)
### END CODE HERE ###
# store values needed for backward propagation in cache
cache = (a_next, c_next, a_prev, c_prev, ft, it, cct, ot, xt, para
return a_next, c_next, yt_pred, cache
```

```
In [7]: | np.random.seed(1)
        xt_tmp = np.random.randn(3, 10)
        a_prev_tmp = np.random.randn(5, 10)
        c_prev_tmp = np.random.randn(5, 10)
        parameters_tmp = {}
        parameters_tmp['Wf'] = np.random.randn(5, 5 + 3)
        parameters tmp['bf'] = np.random.randn(5, 1)
        parameters tmp['Wi'] = np.random.randn(5, 5 + 3)
        parameters tmp['bi'] = np.random.randn(5, 1)
        parameters_tmp['Wo'] = np.random.randn(5, 5 + 3)
        parameters tmp['bo'] = np.random.randn(5, 1)
        parameters_tmp['Wc'] = np.random.randn(5, 5 + 3)
        parameters_tmp['bc'] = np.random.randn(5, 1)
        parameters tmp['Wy'] = np.random.randn(2, 5)
        parameters tmp['by'] = np.random.randn(2, 1)
        a_next_tmp, c_next_tmp, yt_tmp, cache_tmp = lstm_cell_forward(xt_tmp,
        print("a_next[4] = \n", a_next_tmp[4])
        print("a_next.shape = ", a_next_tmp.shape)
print("c_next[2] = \n", c_next_tmp[2])
        print("c_next.shape = ", c_next_tmp.shape)
        print("yt[1] =", yt_tmp[1])
        print("yt.shape = ", yt_tmp.shape)
        print("cache[1][3] =\n", cache_tmp[1][3])
        print("len(cache) = ", len(cache_tmp))
        # UNIT TEST
        lstm_cell_forward_test(lstm_cell_forward)
        a next[4] =
         [-0.66408471 0.0036921
                                   0.02088357 0.22834167 -0.85575339 0.0013
        8482
          0.76566531 0.34631421 - 0.00215674 0.43827275
        a_next.shape = (5, 10)
        c next[2] =
         2942
          0.76449811 - 0.0981561 - 0.74348425 - 0.26810932
        c next.shape = (5, 10)
        yt[1] = [0.79913913 \ 0.15986619 \ 0.22412122 \ 0.15606108 \ 0.97057211 \ 0.311
        46381
         0.00943007 0.12666353 0.39380172 0.07828381]
        yt.shape = (2, 10)
        cache[1][3] =
         [-0.16263996 \ 1.03729328 \ 0.72938082 \ -0.54101719 \ 0.02752074 \ -0.3082
        1874
          0.07651101 -1.03752894 1.41219977 -0.37647422]
        len(cache) = 10
        All tests passed
```

Expected Output:

```
a_next[4] =
 [-0.66408471 0.0036921
                       0.00138482
 0.76566531 0.34631421 - 0.00215674 0.43827275
a_next.shape = (5, 10)
c next[2] =
 0.11832942
 0.76449811 - 0.0981561 - 0.74348425 - 0.26810932
c_next.shape = (5, 10)
yt[1] = [0.79913913 0.15986619 0.22412122 0.15606108 0.97]
057211 0.31146381
 0.00943007 0.12666353 0.39380172 0.07828381]
yt.shape = (2, 10)
cache[1][3] =
 [-0.16263996 \quad 1.03729328 \quad 0.72938082 \quad -0.54101719 \quad 0.02752074
-0.30821874
 0.07651101 -1.03752894 1.41219977 -0.376474221
len(cache) = 10
```

2.2 - Forward Pass for LSTM

Now that you have implemented one step of an LSTM, you can iterate this over it using a for loop to process a sequence of T_x inputs.

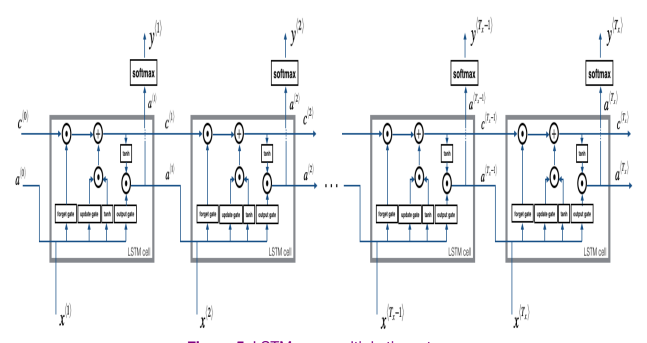


Figure 5: LSTM over multiple time steps.

Exercise 4 - Istm forward

Implement lstm_forward() to run an LSTM over T_x time steps.

Instructions

- Get the dimensions n_x , n_a , n_y , m, T_x from the shape of the variables: x and parameters
- Initialize the 3D tensors a, c and y
 - a: hidden state, shape (n_a, m, T_x)
 - c: cell state, shape (n_a, m, T_x)
 - y: prediction, shape (n_v, m, T_x) (Note that $T_v = T_x$ in this example)
 - Note Setting one variable equal to the other is a "copy by reference". In other words, don't do `c = a', otherwise both these variables point to the same underlying variable.
- Initialize the 2D tensor $a^{\langle t \rangle}$
 - $a^{\langle t \rangle}$ stores the hidden state for time step t. The variable name is a_next.
 - $a^{\langle 0 \rangle}$, the initial hidden state at time step 0, is passed in when calling the function. The variable name is a0 .
 - $a^{\langle t \rangle}$ and $a^{\langle 0 \rangle}$ represent a single time step, so they both have the shape (n_a, m)
 - Initialize $a^{\langle t \rangle}$ by setting it to the initial hidden state $(a^{\langle 0 \rangle})$ that is passed into the function.
- Initialize $c^{\langle t \rangle}$ with zeros.
 - The variable name is c_next
 - $c^{\langle t \rangle}$ represents a single time step, so its shape is (n_a, m)
 - **Note**: create c_next as its own variable with its own location in memory. Do not initialize it as a slice of the 3D tensor c. In other words, **don't** do c_next = c[:,:,0].
- For each time step, do the following:
 - From the 3D tensor x, get a 2D slice $x^{(t)}$ at time step t
 - Call the lstm_cell_forward function that you defined previously, to get the hidden state, cell state, prediction, and cache
 - Store the hidden state, cell state and prediction (the 2D tensors) inside the 3D tensors
 - Append the cache to the list of caches

```
bf — Bias of the forget gate, numpy array of
                    Wi -- Weight matrix of the update gate, numpy
                    bi -- Bias of the update gate, numpy array of
                    Wc -- Weight matrix of the first "tanh", numpy
                    bc -- Bias of the first "tanh", numpy array of
                    Wo -- Weight matrix of the output gate, numpy
                    bo -- Bias of the output gate, numpy array of
                    Wy -- Weight matrix relating the hidden-state
                    by -- Bias relating the hidden-state to the ou
Returns:
a -- Hidden states for every time-step, numpy array of shape (n_a,
y -- Predictions for every time-step, numpy array of shape (n_y, m
c -- The value of the cell state, numpy array of shape (n_a, m, T_
caches -- tuple of values needed for the backward pass, contains (
# Initialize "caches", which will track the list of all the caches
caches = []
### START CODE HERE ###
Wy = parameters['Wy'] # saving parameters['Wy'] in a local variabl
# Retrieve dimensions from shapes of x and parameters['Wy'] (≈2 li
n_x, m, T_x = x.shape
n_y, n_a = Wy.shape
# initialize "a", "c" and "y" with zeros (≈3 lines)
a = np.zeros((n_a,m,T_x))
c = np.zeros((n_a,m,T_x))
y = np.zeros((n y,m,T x))
# Initialize a_next and c_next (≈2 lines)
a_next = a0[:,:]
c_next = np.zeros((n_a,m))
# loop over all time-steps
for t in range(T x):
   # Get the 2D slice 'xt' from the 3D input 'x' at time step 't'
   xt = x[:,:,t]
    # Update next hidden state, next memory state, compute the pre
    a_next, c_next, yt, cache = lstm_cell_forward(xt, a_next, c_next)
    # Save the value of the new "next" hidden state in a (≈1 line)
    a[:,:,t] = a next
    # Save the value of the next cell state (≈1 line)
    c[:,:,t] = c next
    # Save the value of the prediction in y (≈1 line)
   y[:,:,t] = yt
    # Append the cache into caches (≈1 line)
    caches.append(cache)
### END CODE HERE ###
# store values needed for backward propagation in cache
```

Wt -- Weight matrix of the forget gate, numpy

```
caches = (caches, x)
return a, y, c, caches
```

```
In [9]: | np.random.seed(1)
        x \text{ tmp} = \text{np.random.randn}(3, 10, 7)
        a0\_tmp = np.random.randn(5, 10)
        parameters_tmp = {}
        parameters_tmp['Wf'] = np.random.randn(5, 5 + 3)
        parameters_tmp['bf'] = np.random.randn(5, 1)
        parameters_tmp['Wi'] = np.random.randn(5, 5 + 3)
        parameters tmp['bi'] = np.random.randn(5, 1)
        parameters tmp['Wo'] = np.random.randn(5, 5 + 3)
        parameters tmp['bo'] = np.random.randn(5, 1)
        parameters_{tmp}['Wc'] = np.random.randn(5, 5 + 3)
        parameters_tmp['bc'] = np.random.randn(5, 1)
        parameters_tmp['Wy'] = np.random.randn(2, 5)
        parameters_tmp['by'] = np.random.randn(2, 1)
        a tmp, y tmp, c tmp, caches tmp = lstm forward(x tmp, a0 tmp, paramete)
        print("a[4][3][6] = ", a_tmp[4][3][6])
        print("a.shape = ", a_tmp.shape)
print("y[1][4][3] =", y_tmp[1][4][3])
        print("y.shape = ", y_tmp.shape)
        print("caches[1][1][1] = \n", caches_tmp[1][1][1])
        print("c[1][2][1]", c_tmp[1][2][1])
        print("len(caches) = ", len(caches_tmp))
        # UNIT TEST
        lstm_forward_test(lstm_forward)
        a[4][3][6] = 0.17211776753291672
        a.shape = (5, 10, 7)
        v[1][4][3] = 0.9508734618501101
        y.shape = (2, 10, 7)
        caches[1][1][1] =
          0.82797464 0.23009474 0.76201118 -0.22232814 -0.20075807 0.1865
        6139
           0.41005165]
        c[1][2][1] -0.8555449167181981
        len(caches) = 2
```

All tests passed

Expected Output:

```
a[4][3][6] = 0.172117767533

a.shape = (5, 10, 7)

y[1][4][3] = 0.95087346185

y.shape = (2, 10, 7)

caches[1][1][1] =

[ 0.82797464     0.23009474     0.76201118     -0.22232814     -0.20075807

0.18656139

     0.41005165]

c[1][2][1]     -0.855544916718

len(caches) = 2
```

Congratulations!

You have now implemented the forward passes for both the basic RNN and the LSTM. When using a deep learning framework, implementing the forward pass is sufficient to build systems that achieve great performance. The framework will take care of the rest.

What you should remember:

- An LSTM is similar to an RNN in that they both use hidden states to pass along information, but an LSTM also uses a cell state, which is like a long-term memory, to help deal with the issue of vanishing gradients
- An LSTM cell consists of a cell state, or long-term memory, a hidden state, or short-term memory, along with 3 gates that constantly update the relevancy of its inputs:
 - A **forget** gate, which decides which input units should be remembered and passed along. It's a tensor with values between 0 and 1.
 - If a unit has a value close to 0, the LSTM will "forget" the stored state in the previous cell state.
 - If it has a value close to 1, the LSTM will mostly remember the corresponding value.
 - An update gate, again a tensor containing values between 0 and 1. It decides on what information to throw away, and what new information to add.
 - When a unit in the update gate is close to 1, the value of its candidate is passed on to the hidden state.
 - When a unit in the update gate is close to 0, it's prevented from being passed onto the hidden state.
 - And an output gate, which decides what gets sent as the output of the time step.

Let's recap all you've accomplished so far. You have:

- Used notation for building sequence models
- Become familiar with the architecture of a basic RNN and an LSTM, and can describe their components

The rest of this notebook is optional, and will not be graded, but as always, you are encouraged to push your own understanding! Good luck and have fun.

3 - Backpropagation in Recurrent Neural Networks (OPTIONAL / UNGRADED)

In modern deep learning frameworks, you only have to implement the forward pass, and the framework takes care of the backward pass, so most deep learning engineers do not need to bother with the details of the backward pass. If, however, you are an expert in calculus (or are just curious) and want to see the details of backprop in RNNs, you can work through this optional portion of the notebook.

When in an earlier <u>course (https://www.coursera.org/learn/neural-networks-deep-learning/lecture/0VSHe/derivatives-with-a-computation-graph)</u> you implemented a simple (fully connected) neural network, you used backpropagation to compute the derivatives with respect to the cost to update the parameters. Similarly, in recurrent neural networks you can calculate the derivatives with respect to the cost in order to update the parameters. The backprop equations are quite complicated, and so were not derived in lecture. However, they're briefly presented for your viewing pleasure below.

Note that this notebook does not implement the backward path from the Loss 'J' backwards to 'a'. This would have included the dense layer and softmax, which are a part of the forward path. This is assumed to be calculated elsewhere and the result passed to rnn_backward in 'da'. It is further assumed that loss has been adjusted for batch size (m) and division by the number of examples is not required here.

This section is optional and ungraded, because it's more difficult and has fewer details regarding its implementation. Note that this section only implements key elements of the full path!

Onward, brave one:

3.1 - Basic RNN Backward Pass

Begin by computing the backward pass for the basic RNN cell. Then, in the following sections, iterate through the cells.

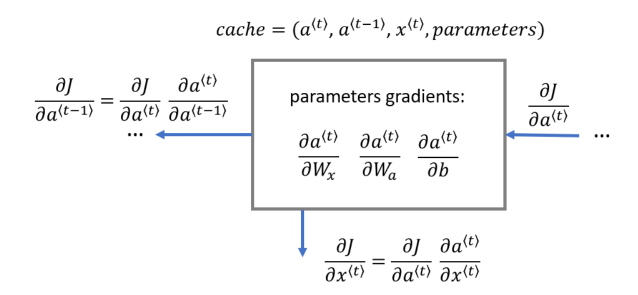


Figure 6: The RNN cell's backward pass. Just like in a fully-connected neural network, the derivative of the cost function J backpropagates through the time steps of the RNN by following the chain rule from calculus. Internal to the cell, the chain rule is also used to calculate $(\frac{\partial J}{\partial W_{ax}}, \frac{\partial J}{\partial W_{aa}}, \frac{\partial J}{\partial b})$ to update the parameters (W_{ax}, W_{aa}, b_a) . The operation can utilize the cached results from the forward path.

Recall from lecture that the shorthand for the partial derivative of cost relative to a variable is dVariable . For example, $\frac{\partial J}{\partial W_{ax}}$ is dW_{ax} . This will be used throughout the remaining sections.

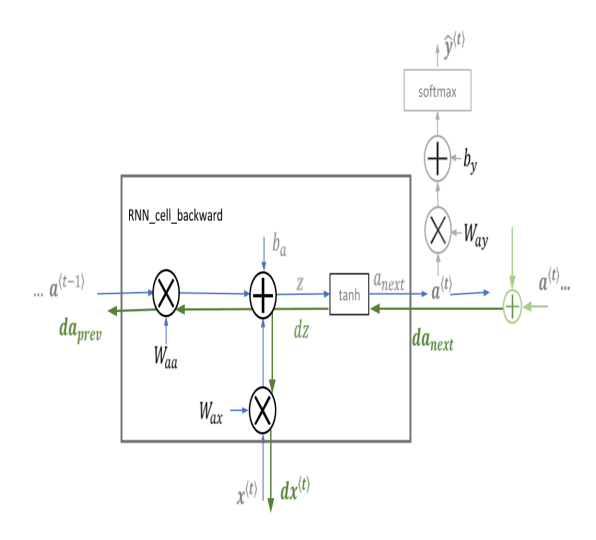


Figure 7: This implementation of `rnn_cell_backward` does **not** include the output dense layer and softmax which are included in `rnn_cell_forward`.

 da_{next} is $\frac{\partial J}{\partial a^{\langle t \rangle}}$ and includes loss from previous stages and current stage output logic. The addition shown in green will be part of your implementation of <code>rnn_backward</code>.

Equations

To compute rnn_cell_backward, you can use the following equations. It's a good exercise to derive them by hand. Here, * denotes element-wise multiplication while the absence of a symbol indicates matrix multiplication.

$$a^{\langle t \rangle} = \tanh(W_{ax} x^{\langle t \rangle} + W_{aa} a^{\langle t-1 \rangle} + b_a)$$
 (-)

$$\frac{\partial \tanh(x)}{\partial x} = 1 - \tanh^2(x) \tag{-}$$

$$dtanh = da_{next} * (1 - \tanh^2(W_{ax}x^{\langle t \rangle} + W_{aa}a^{\langle t-1 \rangle} + b_a))$$
 (0)

$$dW_{ax} = dtanh \cdot x^{\langle t \rangle T} \tag{1}$$

$$dW_{aa} = dtanh \cdot a^{(t-1)T} \tag{2}$$

$$db_a = \sum_{batch} dtanh \tag{3}$$

$$dx^{\langle t \rangle} = W_{ax}^T \cdot dtanh \tag{4}$$

$$da_{prev} = W_{aa}^{T} \cdot dtanh \tag{5}$$

Exercise 5 - rnn cell backward

Implementing rnn cell backward.

The results can be computed directly by implementing the equations above. However, you have an option to simplify them by computing 'dz' and using the chain rule. This can be further simplified by noting that $\tanh(W_{ax}x^{\langle t\rangle}+W_{aa}a^{\langle t-1\rangle}+b_a)$ was computed and saved as a_next in the forward pass.

To calculate dba, the 'batch' above is a sum across all 'm' examples (axis= 1). Note that you should use the keepdims = True option.

It may be worthwhile to review Course 1 <u>Derivatives with a Computation Graph</u> (https://www.coursera.org/learn/neural-networks-deep-learning/lecture/0VSHe/derivatives-with-a-computation-graph) through <u>Backpropagation Intuition</u> (https://www.coursera.org/learn/neural-networks-deep-learning/lecture/6dDj7/backpropagation-intuition-optional), which decompose the calculation into steps using the chain rule.

Matrix vector derivatives are described here (http://cs231n.stanford.edu/vecDerivs.pdf),

though the equations above incorporate the required transformations.

Note: rnn_cell_backward does **not** include the calculation of loss from $y\langle t\rangle$. This is incorporated into the incoming da_next . This is a slight mismatch with rnn_cell_forward , which includes a dense layer and softmax.

Note on the code:

 $dx^{\langle t \rangle}$ is represented by dxt, dW_{ax} is represented by dWax, da_{prev} is represented by da_prev, dW_{aa} is represented by dWaa, db_a is represented by dba,

dz is not derived above but can optionally be derived by students to simplify the repeated calculations.

```
In [50]: # UNGRADED FUNCTION: rnn cell backward
         def rnn_cell_backward(da_next, cache):
             Implements the backward pass for the RNN-cell (single time-step).
             Arguments:
             da next -- Gradient of loss with respect to next hidden state
             cache -- python dictionary containing useful values (output of rnd
             Returns:
             gradients -- python dictionary containing:
                                 dx -- Gradients of input data, of shape (n_x,
                                 da prev -- Gradients of previous hidden state,
                                 dWax -- Gradients of input-to-hidden weights,
                                 dWaa -- Gradients of hidden-to-hidden weights,
                                 dba -- Gradients of bias vector, of shape (n a
             .....
             # Retrieve values from cache
             (a_next, a_prev, xt, parameters) = cache
             # Retrieve values from parameters
             Wax = parameters["Wax"]
             Waa = parameters["Waa"]
             Wya = parameters["Wya"]
             ba = parameters["ba"]
             by = parameters["by"]
             ### START CODE HERE ###
             # compute the gradient of dtanh term using a_next and da_next (≈1
             dtanh = da next*(1-a next**2)
             # compute the gradient of the loss with respect to Wax (≈2 lines)
             dxt = np.dot(Wax.T,dtanh)
             dWax = np.dot(dtanh,xt.T)
             # compute the gradient with respect to Waa (≈2 lines)
             da prev = np.dot(Waa.T,dtanh)
             dWaa = np.dot(dtanh,a_prev.T)
             # compute the gradient with respect to b (≈1 line)
             dba = np.sum(dtanh,axis=1,keepdims=True)
             ### END CODE HERE ###
             # Store the gradients in a python dictionary
             gradients = {"dxt": dxt, "da_prev": da_prev, "dWax": dWax, "dWaa":
             return gradients
```

```
In [51]: | np.random.seed(1)
          xt_tmp = np.random.randn(3,10)
          a_prev_tmp = np.random.randn(5,10)
          parameters_tmp = {}
          parameters_tmp['Wax'] = np.random.randn(5,3)
          parameters_tmp['Waa'] = np.random.randn(5,5)
          parameters tmp['Wya'] = np.random.randn(2,5)
          parameters tmp['ba'] = np.random.randn(5,1)
          parameters tmp['by'] = np.random.randn(2,1)
          a_next_tmp, yt_tmp, cache_tmp = rnn_cell_forward(xt_tmp, a_prev_tmp, p
          da next tmp = np.random.randn(5,10)
          gradients tmp = rnn cell backward(da next tmp, cache tmp)
          print("gradients[\"dxt\"][1][2] =", gradients_tmp["dxt"][1][2])
print("gradients[\"dxt\"].shape =", gradients_tmp["dxt"].shape)
          print("gradients[\"da_prev\"][2][3] =", gradients_tmp["da_prev"][2][3]
          print("gradients[\"da_prev\"].shape =", gradients_tmp["da_prev"].shape
          print("gradients[\"dWax\"][3][1] =", gradients_tmp["dWax"][3][1])
print("gradients[\"dWax\"].shape =", gradients_tmp["dWax"].shape)
print("gradients[\"dWaa\"][1][2] =", gradients_tmp["dWaa"][1][2])
          print("gradients[\"dWaa\"].shape =", gradients_tmp["dWaa"].shape)
          print("gradients[\"dba\"][4] =", gradients_tmp["dba"][4])
          print("gradients[\"dba\"].shape =", gradients_tmp["dba"].shape)
          gradients ["dxt"] [1] [2] = -1.3872130506020925
          gradients["dxt"].shape = (3, 10)
          gradients ["da_prev"] [2] [3] = -0.15239949377395495
          gradients["da prev"].shape = (5, 10)
          gradients["dWax"][3][1] = 0.4107728249354584
          gradients["dWax"].shape = (5, 3)
          gradients["dWaa"][1][2] = 1.1503450668497135
          gradients["dWaa"].shape = (5, 5)
```

gradients["dba"][4] = [0.20023491]
gradients["dba"].shape = (5, 1)

```
gradients["dxt"][1][2] =
                                 -1.3872130506
    gradients["dxt"].shape =
                                          (3, 10)
 gradients["da_prev"][2][3] = -0.152399493774
gradients["da_prev"].shape =
                                          (5, 10)
   gradients["dWax"][3][1] =
                               0.410772824935
  gradients["dWax"].shape =
                                           (5, 3)
   gradients["dWaa"][1][2] =
                                 1.15034506685
  gradients["dWaa"].shape =
                                           (5, 5)
        gradients["dba"][4] =
                                  [ 0.20023491]
    gradients["dba"].shape =
                                           (5, 1)
```

Exercise 6 - rnn backward

Computing the gradients of the cost with respect to $a^{\langle t \rangle}$ at every time step t is useful because it is what helps the gradient backpropagate to the previous RNN cell. To do so, you need to iterate through all the time steps starting at the end, and at each step, you increment the overall db_a , dW_{aa} , dW_{ax} and you store dx.

Instructions:

Implement the rnn_backward function. Initialize the return variables with zeros first, and then loop through all the time steps while calling the rnn_cell_backward at each time time step, updating the other variables accordingly.

- Note that this notebook does not implement the backward path from the Loss 'J' backwards to 'a'.
 - This would have included the dense layer and softmax which are a part of the forward path.
 - This is assumed to be calculated elsewhere and the result passed to rnn backward in 'da'.
 - You must combine this with the loss from the previous stages when calling rnn_cell_backward (see figure 7 above).
- It is further assumed that loss has been adjusted for batch size (m).
 - Therefore, division by the number of examples is not required here.

```
In [52]: # UNGRADED FUNCTION: rnn_backward

def rnn_backward(da, caches):
```

```
Implement the backward pass for a RNN over an entire sequence of i
Arguments:
da -- Upstream gradients of all hidden states, of shape (n_a, m, T
caches —— tuple containing information from the forward pass (rnn
Returns:
gradients -- python dictionary containing:
                    dx -- Gradient w.r.t. the input data, numpy-ar
                    da0 -- Gradient w.r.t the initial hidden state
                    dWax -- Gradient w.r.t the input's weight matr
                    dWaa -- Gradient w.r.t the hidden state's weig
                    dba -- Gradient w.r.t the bias, of shape (n_a,
1111111
### START CODE HERE ###
# Retrieve values from the first cache (t=1) of caches (≈2 lines)
(caches, x) = caches
(a1, a0, x1, parameters) = caches[0]
# Retrieve dimensions from da's and x1's shapes (≈2 lines)
n_a, m, T_x = da.shape
n_x, m = x1.shape
# initialize the gradients with the right sizes (≈6 lines)
dx = np.zeros((n_x,m,T_x))
dWax = np.zeros((n_a,n_x))
dWaa = np.zeros((n a,n a))
dba = np.zeros((n_a,1))
da0 = np.zeros((n_a,m))
da_prevt = da[:,:,-1]
# Loop through all the time steps
for t in reversed(range(T x)):
    # Compute gradients at time step t. Choose wisely the "da next
    gradients = rnn_cell_backward(da[:,:,t], caches[t])
    # Retrieve derivatives from gradients (≈ 1 line)
    dxt, da_prevt, dWaxt, dWaat, dbat = gradients["dxt"], gradient
    # Increment global derivatives w.r.t parameters by adding thei
    dx[:, :, t] = dxt
    dWax += dWaxt
    dWaa += dWaat
    dba += dbat
# Set da0 to the gradient of a which has been backpropagated throu
da0 = da prevt
### END CODE HERE ###
# Store the gradients in a python dictionary
gradients = {"dx": dx, "da0": da0, "dWax": dWax, "dWaa": dWaa, "dba
return gradients
```

```
In [53]: |np.random.seed(1)
         x_{tmp} = np.random.randn(3,10,4)
         a0 tmp = np.random.randn(5,10)
         parameters_tmp = {}
         parameters_tmp['Wax'] = np.random.randn(5,3)
         parameters_tmp['Waa'] = np.random.randn(5,5)
         parameters tmp['Wya'] = np.random.randn(2,5)
         parameters tmp['ba'] = np.random.randn(5,1)
         parameters tmp['by'] = np.random.randn(2,1)
         a_tmp, y_tmp, caches_tmp = rnn_forward(x_tmp, a0_tmp, parameters_tmp)
         da_{tmp} = np.random.randn(5, 10, 4)
         gradients_tmp = rnn_backward(da_tmp, caches_tmp)
         print("gradients[\"dx\"][1][2] =", gradients_tmp["dx"][1][2])
         print("gradients[\"dx\"].shape =", gradients_tmp["dx"].shape)
         print("gradients[\"da0\"][2][3] =", gradients_tmp["da0"][2][3])
         print("gradients[\"da0\"].shape =", gradients_tmp["da0"].shape)
         print("gradients[\"dWax\"][3][1] =", gradients_tmp["dWax"][3][1])
         print("gradients[\"dWax\"].shape =", gradients_tmp["dWax"].shape)
print("gradients[\"dWaa\"][1][2] =", gradients_tmp["dWaa"][1][2])
         print("gradients[\"dWaa\"].shape =", gradients_tmp["dWaa"].shape)
         print("gradients[\"dba\"][4] =", gradients_tmp["dba"][4])
         print("gradients[\"dba\"].shape =", gradients_tmp["dba"].shape)
         gradients ["dx"][1][2] = [-0.15028183 -0.34554547 0.02071758 0.01483]
         3171
         gradients["dx"].shape = (3, 10, 4)
         gradients ["da0"] [2] [3] = -0.17268893183890754
         gradients["da0"].shape = (5, 10)
         qradients["dWax"][3][1] = 4.081485734449453
         gradients["dWax"].shape = (5, 3)
         gradients["dWaa"][1][2] = 1.056012342849445
         gradients["dWaa"].shape = (5, 5)
         gradients["dba"][4] = [-0.12427391]
         gradients["dba"].shape = (5, 1)
```

gradients["dx"][1][2] =	[-2.07101689 -0.59255627 0.02466855 0.01483317]		
gradients["dx"].shape =	(3, 10, 4)		
gradients["da0"][2][3] =	-0.314942375127		
gradients["da0"].shape =	(5, 10)		
gradients["dWax"][3][1] =	11.2641044965		
gradients["dWax"].shape =	(5, 3)		
gradients["dWaa"][1][2] =	2.30333312658		
gradients["dWaa"].shape =	(5, 5)		
gradients["dba"][4] =	[-0.74747722]		
gradients["dba"].shape =	(5, 1)		

3.2 - LSTM Backward Pass

1. One Step Backward

The LSTM backward pass is slightly more complicated than the forward pass.

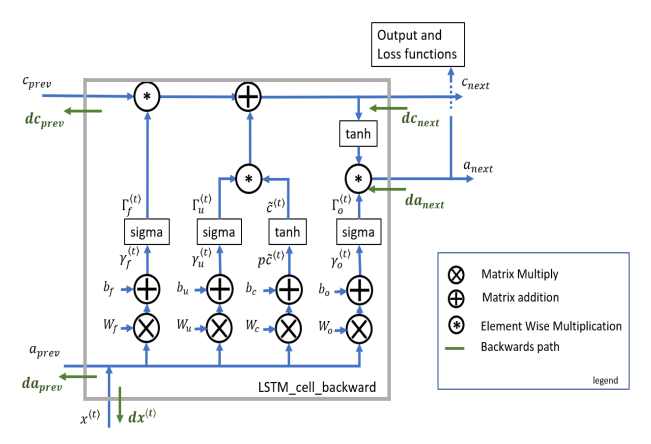


Figure 8: LSTM Cell Backward. Note the output functions, while part of the `lstm_cell_forward`, are not included in `lstm_cell_backward`

The equations for the LSTM backward pass are provided below. (If you enjoy calculus exercises feel free to try deriving these from scratch yourself.)

2. Gate Derivatives

Note the location of the gate derivatives (γ ..) between the dense layer and the activation function (see graphic above). This is convenient for computing parameter derivatives in the next step.

$$d\gamma_o^{\langle t \rangle} = da_{next} * \tanh(c_{next}) * \Gamma_o^{\langle t \rangle} * \left(1 - \Gamma_o^{\langle t \rangle}\right)$$

$$dp\widetilde{c}^{\langle t \rangle} = \left(dc_{next} * \Gamma_u^{\langle t \rangle} + \Gamma_o^{\langle t \rangle} * (1 - \tanh^2(c_{next})) * \Gamma_u^{\langle t \rangle} * da_{next}\right) * \left(1 - \left(\widetilde{c}^{\langle t \rangle}\right)^2\right)$$

$$d\gamma_u^{\langle t \rangle} = \left(dc_{next} * \widetilde{c}^{\langle t \rangle} + \Gamma_o^{\langle t \rangle} * (1 - \tanh^2(c_{next})) * \widetilde{c}^{\langle t \rangle} * da_{next}\right) * \Gamma_u^{\langle t \rangle} * \left(1 - \Gamma_u^{\langle t \rangle}\right)$$

$$d\gamma_f^{\langle t \rangle} = \left(dc_{next} * c_{prev} + \Gamma_o^{\langle t \rangle} * (1 - \tanh^2(c_{next})) * c_{prev} * da_{next}\right) * \Gamma_f^{\langle t \rangle} * \left(1 - \Pi_u^{\langle t \rangle}\right)$$

3. Parameter Derivatives

$$dW_f = d\gamma_f^{\langle t \rangle} \begin{bmatrix} a_{prev} \\ x_t \end{bmatrix}_T^T \tag{11}$$

$$dW_{u} = d\gamma_{u}^{\langle t \rangle} \begin{bmatrix} a_{prev} \\ x_{t} \end{bmatrix}^{T}$$

$$(12)$$

$$dW_{c} = dp\widetilde{c}^{\langle t \rangle} \begin{bmatrix} x_{t} \\ a_{prev} \\ x_{t} \end{bmatrix}^{T}$$

$$dW_{o} = d\gamma_{o}^{\langle t \rangle} \begin{bmatrix} a_{prev} \\ x_{t} \end{bmatrix}^{T}$$
(13)

$$dW_o = d\gamma_o^{\langle t \rangle} \begin{bmatrix} a_{prev} \\ x_t \end{bmatrix}^T \tag{14}$$

To calculate db_f, db_u, db_c, db_o you just need to sum across all 'm' examples (axis= 1) on $d\gamma_f^{\langle t \rangle}, d\gamma_u^{\langle t \rangle}, dp\widetilde{c}^{\langle t \rangle}, d\gamma_o^{\langle t \rangle}$ respectively. Note that you should have the keepdims = True option.

$$db_f = \sum d\gamma_f^{\langle t \rangle} \tag{15}$$

$$db_u = \sum_{u}^{batch} d\gamma_u^{\langle t \rangle} \tag{16}$$

$$db_c = \sum_{batch}^{batch} d\gamma_c^{\langle t \rangle} \tag{17}$$

$$db_o = \sum_{batch}^{batch} d\gamma_o^{\langle t \rangle} \tag{18}$$

Finally, you will compute the derivative with respect to the previous hidden state, previous memory state, and input.

$$da_{prev} = W_f^T d\gamma_f^{\langle t \rangle} + W_u^T d\gamma_u^{\langle t \rangle} + W_c^T dp \widetilde{c}^{\langle t \rangle} + W_o^T d\gamma_o^{\langle t \rangle}$$
(19)

Here, to account for concatenation, the weights for equations 19 are the first n_a, (i.e. $W_f = W_f[:,:n_a]$ etc...)

$$dc_{prev} = dc_{next} * \Gamma_f^{\langle t \rangle} + \Gamma_o^{\langle t \rangle} * (1 - \tanh^2(c_{next})) * \Gamma_f^{\langle t \rangle} * da_{next}$$
 (20)

$$dx^{\langle t \rangle} = W_f^T d\gamma_f^{\langle t \rangle} + W_u^T d\gamma_u^{\langle t \rangle} + W_c^T dp \tilde{c}^{\langle t \rangle} + W_o^T d\gamma_o^{\langle t \rangle}$$
(21)

where the weights for equation 21 are from n_a to the end, (i.e. $W_f = W_f[:, n_a:]$ etc...)

Exercise 7 - Istm cell backward

Implement lstm_cell_backward by implementing equations 7-21 below.

Note:

In the code:

```
d\gamma_u^{\langle t \rangle} is represented by dit,
                    d\gamma_{\rm f}^{\langle t \rangle} is represented by dft
In [42]: # UNGRADED FUNCTION: lstm_cell_backward
                    def lstm_cell_backward(da_next, dc_next, cache):
                            Implement the backward pass for the LSTM-cell (single time-step).
                            Arguments:
                            da_next -- Gradients of next hidden state, of shape (n_a, m)
                            dc_next -- Gradients of next cell state, of shape (n_a, m)
                            cache -- cache storing information from the forward pass
                            Returns:
                            gradients -- python dictionary containing:
                                                                       dxt -- Gradient of input data at time-step t,
                                                                       da_prev -- Gradient w.r.t. the previous hidder
                                                                       dc_prev -- Gradient w.r.t. the previous memory
                                                                       dWf -- Gradient w.r.t. the weight matrix of th
                                                                       dWi -- Gradient w.r.t. the weight matrix of th
                                                                       dWc -- Gradient w.r.t. the weight matrix of th
                                                                       dWo -- Gradient w.r.t. the weight matrix of th
                                                                       dbf -- Gradient w.r.t. biases of the forget ga
                                                                       dbi -- Gradient w.r.t. biases of the update ga
                                                                       dbc -- Gradient w.r.t. biases of the memory ga
                                                                       dbo -- Gradient w.r.t. biases of the output ga
                            # Retrieve information from "cache"
                            (a_next, c_next, a_prev, c_prev, ft, it, cct, ot, xt, parameters)
                            ### START CODE HERE ###
                            # Retrieve dimensions from xt's and a next's shape (≈2 lines)
                            n_x, m = xt.shape
                            n_a, m = a_next.shape
                            # Compute gates related derivatives. Their values can be found by
                            dot = da_next*np.tanh(c_next)*ot*(1-ot)
                            dcct = (dc_next*it + ot*(1-(np_tanh(c_next))**2)*it*da_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-cd_next)*(1-
                            dit = (dc next*cct + ot*(1-(np.tanh(c next))**2)*cct*da next)*it*(
                            dft = (dc_next*c_prev + ot*(1-(np.tanh(c_next))**2)*c_prev*da_next
                            # Compute parameters related derivatives. Use equations (11)-(18)
                            dWf = np.dot(dft,np.concatenate((a_prev,xt)).T)
                            dWi = np.dot(dit,np.concatenate((a_prev,xt)).T)
                            dWc = np.dot(dcct,np.concatenate((a prev,xt)).T)
                            dWo = np.dot(dot,np.concatenate((a_prev,xt)).T)
                            dbf = np.sum(dft,axis=1,keepdims=True)
                            dbi = np.sum(dit,axis=1,keepdims=True)
```

 $d\gamma_o^{\langle t \rangle}$ is represented by dot, $dp\widetilde{c}^{\langle t \rangle}$ is represented by dcct,

```
In [43]: |np.random.seed(1)
         xt_tmp = np.random.randn(3,10)
         a prev_tmp = np.random.randn(5,10)
         c_prev_tmp = np.random.randn(5,10)
         parameters tmp = {}
         parameters_tmp['Wf'] = np.random.randn(5, 5+3)
         parameters_tmp['bf'] = np.random.randn(5,1)
         parameters tmp['Wi'] = np.random.randn(5, 5+3)
         parameters tmp['bi'] = np.random.randn(5,1)
         parameters_tmp['Wo'] = np.random.randn(5, 5+3)
         parameters tmp['bo'] = np.random.randn(5,1)
         parameters_tmp['Wc'] = np.random.randn(5, 5+3)
         parameters tmp['bc'] = np.random.randn(5,1)
         parameters_tmp['Wy'] = np.random.randn(2,5)
         parameters tmp['by'] = np.random.randn(2,1)
         a_next_tmp, c_next_tmp, yt_tmp, cache_tmp = lstm_cell_forward(xt tmp,
         da_next_tmp = np.random.randn(5,10)
         dc_next_tmp = np.random.randn(5,10)
         gradients_tmp = lstm_cell_backward(da_next_tmp, dc_next_tmp, cache_tmp
         print("gradients[\"dxt\"][1][2] =", gradients_tmp["dxt"][1][2])
         print("gradients[\"dxt\"].shape =", gradients_tmp["dxt"].shape)
         print("gradients[\"da_prev\"][2][3] =", gradients_tmp["da_prev"][2][3]
         print("gradients[\"da_prev\"].shape =", gradients_tmp["da_prev"].shape
         print("gradients[\"dc_prev\"][2][3] =", gradients_tmp["dc_prev"][2][3]
print("gradients[\"dc_prev\"].shape =", gradients_tmp["dc_prev"].shape
         print("gradients[\"dWf\"][3][1] =", gradients_tmp["dWf"][3][1])
         print("gradients[\"dWf\"].shape =", gradients_tmp["dWf"].shape)
         print("gradients[\"dWi\"][1][2] =", gradients_tmp["dWi"][1][2])
         print("gradients[\"dWi\"].shape =", gradients_tmp["dWi"].shape)
         print("gradients[\"dWc\"][3][1] =", gradients_tmp["dWc"][3][1])
         print("gradients[\"dWc\"].shape =", gradients_tmp["dWc"].shape)
         print("gradients[\"dWo\"][1][2] =", gradients_tmp["dWo"][1][2])
         print("gradients[\"dWo\"].shape =", gradients_tmp["dWo"].shape)
```

```
print( gradients[\ ubi\ ][4] = , gradients_tmp[ ubi ][4])
print("gradients[\"dbf\"].shape =", gradients_tmp["dbf"].shape)
print("gradients[\"dbi\"][4] =", gradients_tmp["dbi"][4])
print("gradients[\"dbc\"].shape =", gradients_tmp["dbc"][4])
print("gradients[\"dbc\"].shape =", gradients_tmp["dbc"].shape)
print("gradients[\"dbo\"][4] =", gradients_tmp["dbo"].shape)
print("gradients[\"dbo\"][4] =", gradients_tmp["dbo"].shape)
```

```
gradients ["dxt"] [1] [2] = 3.2305591151091875
gradients["dxt"].shape = (3, 10)
gradients ["da prev"] [2] [3] = -0.06396214197109236
gradients["da_prev"].shape = (8, 10)
gradients["dc prev"][2][3] = 0.7975220387970015
gradients["dc_prev"].shape = (5, 10)
gradients["dWf"][3][1] = -0.1479548381644968
gradients["dWf"].shape = (5, 8)
gradients["dWi"][1][2] = 1.0574980552259903
gradients["dWi"].shape = (5, 8)
gradients["dWc"][3][1] = 2.3045621636876668
gradients["dWc"].shape = (5, 8)
gradients["dWo"][1][2] = 0.3313115952892109
gradients["dWo"].shape = (5, 8)
gradients["dbf"][4] = [0.18864637]
gradients["dbf"].shape = (5, 1)
gradients ["dbi"] [4] = [-0.40142491]
gradients["dbi"].shape = (5, 1)
qradients["dbc"][4] = [0.25587763]
gradients["dbc"].shape = (5, 1)
qradients["dbo"][4] = [0.13893342]
gradients["dbo"].shape = (5, 1)
```

gradients["dxt"][1][2] =	3.23055911511	
gradients["dxt"].shape =	(3, 10)	
gradients["da_prev"][2][3] =	-0.0639621419711	
gradients["da_prev"].shape =	(5, 10)	
gradients["dc_prev"][2][3] =	0.797522038797	
gradients["dc_prev"].shape =	(5, 10)	
gradients["dWf"][3][1] =	-0.147954838164	
gradients["dWf"].shape =	(5, 8)	
gradients["dWi"][1][2] =	1.05749805523	
gradients["dWi"].shape =	(5, 8)	
gradients["dWc"][3][1] =	2.30456216369	
gradients["dWc"].shape =	(5, 8)	
gradients["dWo"][1][2] =	0.331311595289	
gradients["dWo"].shape =	(5, 8)	
gradients["dbf"][4] =	[0.18864637]	
gradients["dbf"].shape =	(5, 1)	
gradients["dbi"][4] =	[-0.40142491]	
gradients["dbi"].shape =	(5, 1)	
gradients["dbc"][4] =	[0.25587763]	
gradients["dbc"].shape =	(5, 1)	
gradients["dbo"][4] =	[0.13893342]	
gradients["dbo"].shape =	(5, 1)	

3.3 Backward Pass through the LSTM RNN

This part is very similar to the rnn_backward function you implemented above. You will first create variables of the same dimension as your return variables. You will then iterate over all the time steps starting from the end and call the one step function you implemented for LSTM at each iteration. You will then update the parameters by summing them individually. Finally return a dictionary with the new gradients.

Exercise 8 - Istm backward

Implement the lstm_backward function.

Instructions: Create a for loop starting from T_x and going backward. For each step, call $lstm_cell_backward$ and update your old gradients by adding the new gradients to them. Note that dxt is not updated, but is stored.

```
In [58]: # UNGRADED FUNCTION: lstm_backward
         def lstm_backward(da, caches):
             Implement the backward pass for the RNN with LSTM-cell (over a who
             Arguments:
             da -- Gradients w.r.t the hidden states, numpy-array of shape (n_a
             caches —— cache storing information from the forward pass (lstm fd
             Returns:
             gradients -- python dictionary containing:
                                 dx -- Gradient of inputs, of shape (n_x, m, T
                                 da0 -- Gradient w.r.t. the previous hidden sta
                                 dWf -- Gradient w.r.t. the weight matrix of th
                                 dWi -- Gradient w.r.t. the weight matrix of th
                                 dWc -- Gradient w.r.t. the weight matrix of th
                                 dWo -- Gradient w.r.t. the weight matrix of th
                                 dbf -- Gradient w.r.t. biases of the forget ga
                                 dbi -- Gradient w.r.t. biases of the update ga
                                 dbc -- Gradient w.r.t. biases of the memory ga
                                 dbo -- Gradient w.r.t. biases of the output ga
             .....
             # Retrieve values from the first cache (t=1) of caches.
             (caches, x) = caches
             (a1, c1, a0, c0, f1, i1, cc1, o1, x1, parameters) = caches[0]
             ### START CODE HERE ###
             # Retrieve dimensions from da's and x1's shapes (≈2 lines)
             n_a, m, T_x = da.shape
             n_x, m = x1.shape
```

```
# initialize the gradients with the right sizes (≈12 lines)
dx = np.zeros((n_x,m,T_x))
da0 = np.zeros((n_a,m))
da_prevt = da[:,:,-1]
dc_prevt = da_prevt*caches[-1][7]*(1-caches[-1][1]**2) ## this
dWf = np.zeros((n_a,n_a+n_x))
dWi = np.zeros((n a,n a+n x))
dWc = np.zeros((n_a,n_a+n_x))
dWo = np.zeros((n a,n a+n x))
dbf = np.zeros((n_a,1))
dbi = np.zeros((n a.1))
dbc = np.zeros((n_a,1))
dbo = np.zeros((n_a,1))
# loop back over the whole sequence
for t in reversed(range(T x)):
      print("T: ",t)
    # Compute all gradients using lstm_cell_backward
    gradients = lstm_cell_backward(da[:,:,t], dc_prevt, caches[t])
    # Store or add the gradient to the parameters' previous step's
    # gradients = {"dxt": dxt, "da prev": da prev, "dc prev": dc p
             "dWc": dWc,"dbc": dbc, "dWo": dWo,"dbo": dbo}
    da_prevt = gradients['da_prev']
    dc_prevt = gradients['dc_prev']
    dx[:,:,t] = gradients['dxt']
    dWf += gradients['dWf']
    dWi += gradients['dWi']
    dWc += gradients['dWc']
    dWo += gradients['dWo']
    dbf += gradients['dbf']
    dbi += gradients['dbi']
    dbc += gradients['dbc']
    dbo += gradients['dbo']
# Set the first activation's gradient to the backpropagated gradie
da0 = da_prevt[:,:]
### END CODE HERE ###
# Store the gradients in a python dictionary
gradients = {"dx": dx, "da0": da0, "dWf": dWf, "dbf": dbf, "dWi": d
            "dWc": dWc, "dbc": dbc, "dWo": dWo, "dbo": dbo}
return gradients
```

```
In [59]: np.random.seed(1)
x_tmp = np.random.randn(3,10,7)
a0_tmp = np.random.randn(5,10)

parameters_tmp = {}
parameters_tmp['Wf'] = np.random.randn(5, 5+3)
parameters_tmp['bf'] = np.random.randn(5,1)
parameters_tmp['Wi'] = np.random.randn(5, 5+3)
narameters_tmp['hi'] = np.random.randn(5,1)
```

```
parameters_tmp['Wc'] = np.random.randn(5, 5+3)
parameters_tmp['bc'] = np.random.randn(5,1)
parameters_tmp['Wy'] = np.zeros((2,5))
                                             # unused, but needed for
parameters_tmp['by'] = np.zeros((2,1))
                                             # unused, but needed for
a_tmp, y_tmp, c_tmp, caches_tmp = lstm_forward(x_tmp, a0_tmp, paramete
da tmp = np.random.randn(5, 10, 4)
gradients tmp = lstm backward(da tmp, caches tmp)
print("gradients[\"dx\"][1][2] =", gradients_tmp["dx"][1][2])
print("gradients[\"dx\"].shape =", gradients_tmp["dx"].shape)
print("gradients[\"da0\"][2][3] =", gradients_tmp["da0"][2][3])
print("gradients[\"da0\"].shape =", gradients_tmp["da0"].shape)
print("gradients[\"dWf\"][3][1] =", gradients_tmp["dWf"][3][1])
print("gradients[\"dWf\"].shape =", gradients_tmp["dWf"].shape)
print("gradients[\"dWi\"][1][2] =", gradients_tmp["dWi"][1][2])
print("gradients[\"dWi\"].shape =", gradients_tmp["dWi"].shape)
print("gradients[\"dWc\"][3][1] =", gradients_tmp["dWc"][3][1])
print("gradients[\"dWc\"].shape =", gradients_tmp["dWc"].shape)
print("gradients[\"dWo\"][1][2] =", gradients_tmp["dWo"][1][2])
print("gradients[\"dWo\"].shape =", gradients_tmp["dWo"].shape)
print("gradients[\"dbf\"][4] =", gradients_tmp["dbf"][4])
print("gradients[\"dbf\"].shape =", gradients_tmp["dbf"].shape)
print("gradients[\"dbi\"][4] =", gradients_tmp["dbi"][4])
print("gradients[\"dbi\"].shape =", gradients tmp["dbi"].shape)
print("gradients[\"dbc\"][4] =", gradients_tmp["dbc"][4])
print("gradients[\"dbc\"].shape =", gradients_tmp["dbc"].shape)
print("gradients[\"dbo\"][4] =", gradients_tmp["dbo"][4])
print("gradients[\"dbo\"].shape =", gradients_tmp["dbo"].shape)
T:
    3
T:
   2
T:
    1
T:
gradients ["dx"][1][2] = [0.0027341 0.2244089 -0.21607787]
                                                               0.01155
883]
gradients["dx"].shape = (3, 10, 4)
gradients ["da0"] [2] [3] = -0.02340087780988835
qradients["da0"].shape = (8, 10)
gradients ["dWf"] [3] [1] = -0.08858050474102298
gradients["dWf"].shape = (5, 8)
gradients["dWi"][1][2] = 0.09975777657186694
gradients["dWi"].shape = (5, 8)
gradients ["dWc"] [3] [1] = -0.11348715302595871
gradients["dWc"].shape = (5, 8)
gradients["dWo"][1][2] = 0.04843891314443014
gradients["dWo"].shape = (5, 8)
gradients["dbf"][4] = [-0.06970858]
gradients["dbf"].shape = (5, 1)
gradients["dbi"][4] = [-0.29085818]
```

parameters_tmp['Wo'] = np.random.randn(5, 5+3)
parameters_tmp['bo'] = np.random.randn(5,1)

```
gradients["dbi"].shape = (5, 1)
gradients["dbc"][4] = [-0.28250795]
gradients["dbc"].shape = (5, 1)
gradients["dbo"][4] = [-0.29798344]
gradients["dbo"].shape = (5, 1)
```

```
gradients["dx"][1][2] = [0.00218254 0.28205375 -0.48292508 -0.43281115]
 gradients["dx"].shape =
                                                                     (3, 10, 4)
  gradients["da0"][2][3] =
                                                             0.312770310257
gradients["da0"].shape =
                                                                       (5, 10)
 gradients["dWf"][3][1] =
                                                           -0.0809802310938
gradients["dWf"].shape =
                                                                        (5, 8)
  gradients["dWi"][1][2] =
                                                              0.40512433093
gradients["dWi"].shape =
                                                                        (5, 8)
 gradients["dWc"][3][1] =
                                                           -0.0793746735512
gradients["dWc"].shape =
                                                                        (5, 8)
                                                             0.038948775763
 gradients["dWo"][1][2] =
gradients["dWo"].shape =
                                                                        (5, 8)
     gradients["dbf"][4] =
                                                                [-0.15745657]
gradients["dbf"].shape =
                                                                        (5, 1)
     gradients["dbi"][4] =
                                                                [-0.50848333]
 gradients["dbi"].shape =
                                                                        (5, 1)
    gradients["dbc"][4] =
                                                                [-0.42510818]
gradients["dbc"].shape =
                                                                        (5, 1)
    gradients["dbo"][4] =
                                                               [-0.17958196]
gradients["dbo"].shape =
                                                                        (5, 1)
```

Congratulations on completing this assignment!

You now understand how recurrent neural networks work! In the next exercise, you'll use an RNN to build a character-level language model. See you there!