HOMEWORK 5: NEURAL NETWORKS

10-601 Introduction to Machine Learning (Spring 2018)
Carnegie Mellon University

https://piazza.com/cmu/spring2018/10601

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Summary In this assignment, you will build a handwriting recognition system using a neural network. As a warmup, Section 1 will lead you through an on-paper example of how to implement a neural network. Then, in Section 2, you will implement an end-to-end system that learns to perform handwritten letter classification.

START HERE: Instructions

- Collaboration Policy: Collaboration on solving the homework is allowed, after you have thought about the problems on your own. It is also OK to get clarification (but not solutions) from books or online resources, again after you have thought about the problems on your own. There are two requirements: first, cite your collaborators fully and completely (e.g., "Jane explained to me what is asked in Question 3.4"). Second, write your solution *independently*: close the book and all of your notes, and send collaborators out of the room, so that the solution comes from you only. See the collaboration policy on the website for more information: http://www.cs.cmu.edu/~mgormley/courses/10601-s18/about.html
- Late Submission Policy: See the late submission policy here: http://www.cs.cmu.edu/~mgormley/courses/10601-s18/about.html
- Submitting your work: You will use Gradescope to submit answers to all questions, and Autolab to submit your code. Please follow instructions at the end of this PDF to correctly submit all your code to Autolab.
 - Gradescope: For written problems such as derivations, proofs, or plots we will be using Gradescope (https://gradescope.com/). Please use ther provided template. Submissions can be handwritten onto the template, but should be labeled and clearly legible. If your writing is not legible, you will not be awarded marks. Alternatively, submissions can be written in LaTeX. Regrade requests can be made, however this gives the TA the opportunity to regrade your entire paper, meaning if additional mistakes are found then points will be deducted. Each derivation/proof should be completed on a separate page.
 - **Autolab:** You will submit your code for programming questions on the homework to Autolab (https://autolab.andrew.cmu.edu/). After uploading your code, our grading

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scripts will autograde your assignment by running your program on a virtual machine (VM). The software installed on the VM is identical to that on linux.andrew.cmu.edu, so you should check that your code runs correctly there. If developing locally, check that the version number of the programming language environment (e.g. Python 2.7, Octave 3.8.2, Open-JDK 1.8.0, g++ 4.8.5) and versions of permitted libraries (e.g. numpy 1.7.1) match those on linux.andrew.cmu.edu. (Octave users: Please make sure you do not use any Matlab-specific libraries in your code that might make it fail against our tests.) You have a total of 10 Autolab submissions. Use them wisely. In order to not waste Autolab submissions, we recommend debugging your implementation on your local machine (or the linux servers) and making sure your code is running correctly first before any Autolab submission.

• Materials: Download from Autolab the tar file ("Download handout"). The tar file will contain all the data that you will need in order to complete this assignment.

1 Written Questions [20 points]

Answer the following questions in the HW5 solutions template provided. Then upload your solutions to Gradescope. You may use LATEX or print the template and hand-write your answers then scan it in. Failure to use the template may result in a penalty.

1.1 Example Feed Forward and Backpropagation [10 points]

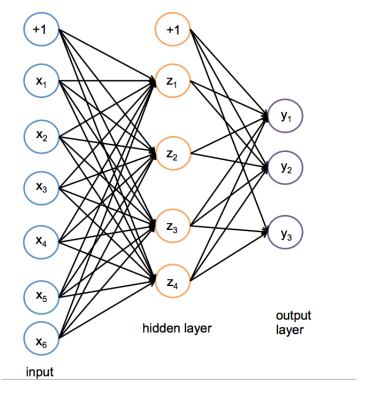


Figure 1.1: A One Hidden Layer Neural Network

Consider the neural network with one hidden layer shown in Figure 1.1. The inputs have 6 features $(x_1,...,x_6)$, the hidden layer has 4 nodes $(z_1,...,z_4)$, and the output is a probability distribution (y_1,y_2,y_3) over 3 classes. We also add a bias to the input, x_0 , as well as to the hidden layer, z_0 and set them to 1. α is the matrix of weights from the inputs to the hidden layer and β is the matrix of weights from the hidden layer to the output layer. $\alpha_{j,i}$ represents the weight going to the node z_j in the hidden layer from the node x_i in the input layer (e.g. $\alpha_{1,2}$ is the weight from x_2 to z_1), and β is defined similarly. We will use a sigmoid activation function for the hidden layer and a softmax for the output layer.

Equivalently, we define each of the following. The input:

$$\mathbf{x} = (x_1, x_2, x_3, x_4, x_5, x_6) \tag{1.1}$$

Linear combination at first (hidden) layer:

$$a_j = \alpha_0 + \sum_{i=1}^6 \alpha_{j,i} * x_i, \ \forall j \in \{1, \dots, 6\}$$
 (1.2)

Activation at first (hidden) layer:

$$z_j = \sigma(a_j) = \frac{1}{1 + \exp(-a_j)}, \ \forall j \in \{1, \dots, 6\}$$
 (1.3)

Linear combination at second (output) layer:

$$b_k = \beta_0 + \sum_{j=1}^4 \beta_{k,j} * z_j, \ \forall k \in \{1, \dots, 3\}$$
 (1.4)

Activation at second (output) layer:

$$\hat{y}_k = \frac{\exp(b_k)}{\sum_{l=1}^{3} \exp(b_l)}, \ \forall k \in \{1, \dots, 3\}$$
(1.5)

Note that the linear combination equations can be written equivalently as the product of the transpose of the weight matrix with the input vector. We can even fold in the bias term α_0 by thinking of $x_0 = 1$, and fold in β_0 by thinking of $z_0 = 1$.

We will use cross entropy loss, $\ell(\hat{\mathbf{y}}, \mathbf{y})$. If \mathbf{y} represents our target output, which will be a one-hot vector representing the correct class, and $\hat{\mathbf{y}}$ represents the output of the network, the loss is calculated by:

$$\ell(\hat{\mathbf{y}}, \mathbf{y}) = -\sum_{i=1}^{3} y_i \log(\hat{y}_i)$$
(1.6)

When doing prediction, we will predict the argmax of the output layer. For example, if $\hat{y}_1 = 0.3$, $\hat{y}_2 = 0.2$, $\hat{y}_3 = 0.5$ we would predict class 3. If the true class from the training data was 2 we would have a one-hot vector \mathbf{y} with values $y_1 = 0$, $y_2 = 1$, $y_3 = 0$.

1. [4 points] We initialize the weights as:

$$\alpha = \begin{bmatrix} 1 & 2 & -3 & 0 & 1 & -3 \\ 3 & 1 & 2 & 1 & 0 & 2 \\ 2 & 2 & 2 & 2 & 2 & 1 \\ 1 & 0 & 2 & 1 & -2 & 2 \end{bmatrix}$$

$$\beta = \begin{bmatrix} 1 & 2 & -2 & 1 \\ 1 & -1 & 1 & 2 \\ 3 & 1 & -1 & 1 \end{bmatrix}$$

And weights on the bias terms ($\alpha_{i,0}$ and $\beta_{i,0}$) are initialized to 1.

You are given a training example $x^{(1)}=(1,1,0,0,1,1)$ with label class 2, so $y^{(1)}=(0,1,0)$. Using the initial weights, run the feed forward of the network over this example (without rounding) and then answer the following questions. In your responses, round to four decimal places (if the answer is an integer you need not include trailing zeros). (Note: the superscript (1) simply indicates that a value corresponds to using training example $x^{(1)}$)

- (a) What is $a_1^{(1)}$?
- (b) What is $z_1^{(1)}$?
- (c) What is $a_3^{(1)}$?
- (d) What is $z_3^{(1)}$?
- (e) What is $b_2^{(1)}$?
- (f) What is $\hat{y}_{2}^{(1)}$?
- (g) Which class would we predict on this example?
- (h) What is the total loss on this example?
- 2. **[5 points]** Now use the results of the previous question to run backpropagation over the network and update the weights. Use learning rate $\eta = 1$.

Do your backpropagation calculations without rounding then answer the following questions, then in your responses, round to four decimal places

- (a) What is the updated value of $\beta_{2,1}$?
- (b) What is the updated weight of the hidden layer bias term applied to y_1 (eg $\beta_{1,0}$)?
- (c) What is the updated value of $\alpha_{3,4}$?
- (d) What is the updated weight of the input layer bias term applied to z_2 (eg $\alpha_{2,0}$)?
- (e) Once we've updated all of our weights if we ran feed forward over the same example again, which class would we predict?
- 3. [1 points] Suppose you are now given a collection of training examples $\mathcal{D} = \{(\mathbf{x}^{(i)}, \mathbf{y}^{(i)})\}$. Explain in English why the cross-entropy loss averaged over these examples is exactly the same quantity as the negative average conditional log-likelihood of the data.

1.2 Empirical Questions [10 points]

The following questions should be completed after you work through the programming portion of this assignment (Section 2).

For these questions, use the large dataset.

Use the following values for the hyperparameters unless otherwise specified:

Paramater	Value
Number of Hidden Units	50
Weight Initialization	0.1
Learning Rate	0.01

Table 1.1: Default values of hyperparameters for experiments in Section 1.2.

For the following questions, submit your solutions to Gradescope. Do **not** include any visualization-related code when submitting to Autolab! Note: we expect it to take about **5 minutes** to train each of these networks.

- 4. [4 points] Train a single hidden layer neural network using the hyperparameters mentioned in Table 1.1, except for the number of hidden units which should vary among 5, 20, 50, 100, and 200. Run the optimization for 100 epochs each time.
 - Plot the average training cross-entropy (sum of the cross-entropy terms over the training dataset divided by the total number of training examples) on the y-axis vs number of hidden units on the x-axis. On the same figure, plot the average validation cross-entropy.
- 5. [1 points] Examine and comment on the plots of training and validation cross-entropy. What is the effect of changing the number of hidden units?
- 6. **[4 points]** Train a single hidden layer neural network using the hyperparameters mentioned in Table 1.1, except for the learning rate which should vary among 0.1, 0.01, and 0.001. Run the optimization for 100 epochs each time.
 - Plot the average training cross-entropy on the y-axis vs the number of epochs on the x-axis for the mentioned learning rates. On the same figure, plot the average validation cross-entropy loss. You may make a separate figure for each learning rate.
- 7. [1 points] Examine and comment on the plots of training and validation cross-entropy. How does adjusting the learning rate affect the convergence of cross-entropy of each dataset?

2 Programming [80 points]

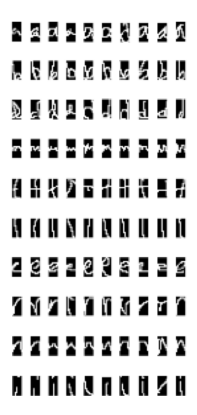


Figure 2.1: 10 Random Images of Each of 10 Letters in OCR

Your goal in this assignment is to label images of handwritten letters by implementing a Neural Network from scratch. You will implement all of the functions needed to initialize, train, evaluate, and make predictions with the network. The programs you write will be automatically graded using the Autolab system. You may write your programs in **Octave**, **Python**, **Java**, **or C++**. However, you should use the same language for all parts below.

2.1 The Task and Datasets

Materials Download the tar file from Autolab ("Download handout"). The tar file will contain all the data that you will need in order to complete this assignment.

Datasets We will be using a subset of an Optical Character Recognition (OCR) dataset. This data includes images of all 26 handwritten letters; our subset will include only the letters "a," "e," "g," "i," "l," "n," "o," "r," "t," and "u." The handout contains three datasets drawn from this data: a small dataset with 60 samples per class (50 for training and 10 for validation), a medium dataset with 600 samples per class (500 for training and 100 for validation), and a large dataset with 1000 samples per class (900 for training and 100 for validation). Figure 2.1 shows a random sample of 10 images of few letters from the dataset.

File Format Each dataset (small, medium, and large) consists of two csv files—train and validation. Each row contains 129 columns separated by commas. The first column contains the label and columns 2 to 129

represent the pixel values of a 16×8 image in a row major format. Label 0 corresponds to "a," 1 to "e," 2 to "g," 3 to "i," 4 to "l," 5 to "n," 6 to "o," 7 to "r," 8 to "t," and 9 to "u." Because the original images are black-and-white (not grayscale), the pixel values are either 0 or 1. However, you should write your code to accept arbitrary pixel values in the range [0,1]. The images in Figure 2.1 were produced by converting these pixel values into .png files for visualization. Observe that no feature engineering has been done here; instead the neural network you build will *learn* features appropriate for the task of character recognition.

2.2 Model Definition

In this assignment, you will implement a single-hidden-layer neural network with a sigmoid activation function for the hidden layer, and a softmax on the output layer. Let the input ectors \mathbf{x} be of length M, the hidden layer \mathbf{z} consist of D hidden units, and the output layer $\hat{\mathbf{y}}$ be a probability distribution over K classes. That is, each element y_k of the output vector represents the probability of \mathbf{x} belonging to the class k.

$$\hat{y}_k = \frac{\exp(b_k)}{\sum_{l=1}^K \exp(b_l)}$$

$$b_k = \beta_{k,0} + \sum_{j=1}^D \beta_{kj} z_j$$

$$z_j = \frac{1}{1 + \exp(-a_j)}$$

$$a_j = \alpha_{j,0} + \sum_{m=1}^M \alpha_{jm} x_m$$

We can compactly express this model by assuming that $x_0=1$ is a bias feature on the input and that $z_0=1$ is also fixed. In this way, we have two parameter matrices $\alpha \in \mathbb{R}^{D\times M+1}$ and $\beta \in \mathbb{R}^{K\times D+1}$. The extra 0th column of each matrix (i.e. $\alpha_{\cdot,0}$ and $\beta_{\cdot,0}$) hold the bias parameters.

$$\hat{y}_k = \frac{\exp(b_k)}{\sum_{l=1}^K \exp(b_l)}$$

$$b_k = \sum_{j=0}^D \beta_{kj} z_j$$

$$z_j = \frac{1}{1 + \exp(-a_j)}$$

$$a_j = \sum_{m=0}^M \alpha_{jm} x_m$$

The objective function we will use for training the neural network is the average cross entropy over the training dataset $\mathcal{D} = \{(\mathbf{x}^{(i)}, \mathbf{y}^{(i)})\}$:

$$J(\alpha, \beta) = -\frac{1}{N} \sum_{i=1}^{N} \sum_{k=1}^{K} y_k^{(i)} \log(\hat{y}_k)$$
 (2.1)

In Equation 2.1, J is a function of the model parameters α and β because \hat{y}_k is implicitly a function of $\mathbf{x}^{(i)}$, α , and β since it is the output of the neural network applied to $\mathbf{x}^{(i)}$. Of course, \hat{y}_k and $y_k^{(i)}$ are the kth components of $\hat{\mathbf{y}}$ and $\mathbf{y}^{(i)}$ respectively.

To train, you should optimize this objective function using stochastic gradient descent (SGD), where the gradient of the parameters for each training example is computed via backpropagation.

2.2.1 Initialization

In order to use a deep network, we must first initialize the weights and biases in the network. This is typically done with a random initialization, or initializing the weights from some other training procedure. For this assignment, we will be using two possible initialization:

RANDOM The weights are initialized randomly from a uniform distribution from -0.1 to 0.1. The bias parameters are initialized to zero.

ZERO All weights are initialized to 0.

You must support both of these initialization schemes.

2.3 Implementation

Write a program $neuralnet.{py|java|cpp|m}$ that implements an optical character recognizer using a one hidden layer neural network with sigmoid activations. Your program should learn the parameters of the model on the training data, report the cross-entropy at the end of each epoch on both train and validation data, and at the end of training write out its predictions and error rates on both datasets.

Your implementation must satisfy the following requirements:

- Use a **sigmoid** activation function on the hidden layer and **softmax** on the output layer to ensure it forms a proper probability distribution.
- Number of **hidden units** for the hidden layer should be determined by a command line flag.
- Support two different **initialization strategies**, as described in Section 2.2.1, selecting between them via a command line flag.
- Use stochastic gradient descent (SGD) to optimize the parameters for one hidden layer neural network. The number of **epochs** will be specified as a command line flag.
- Set the **learning rate** via a command line flag.
- Perform stochastic gradient descent updates on the training data in the order that the data is given
 in the input file. Although you would typically shuffle training examples when using stochastic gradient descent, in order to autograde the assignment, we ask that you DO NOT shuffle trials in this
 assignment.
- You may assume that the input data will always have the same *number* of features (i.e. number of columns) and the same output label space (i.e. $\{0,1,\ldots,9\}$. Other than these assumptions, do not hard-code any aspects of the data sets into your code. We will autograde your programs on multiple (hidden) data sets that include different examples.
- Do *not* use any machine learning libraries. You may use supported linear algebra packages. See Section 2.3.1 for more details.

Implementing a neural network can be tricky: the parameters are not just a simple vector, but a collection of many parameters; computational efficiency of the model itself becomes essential; the initialization strategy dramatically impacts overall learning quality; other aspects we will *not* change (e.g. activation function, optimization method) also have a large effect. These *tips* should help you along the way:

- Try to "vectorize" your code as much as possible—this is particularly important for Python and Octave. For example, in Python, you want to avoid for-loops and instead rely on numpy calls to perform operations such as matrix multiplication, transpose, subtraction, etc. over an entire numpy array at once. Why? Because these operations are actually implemented in fast C code, which won't get bogged down the way a high-level scripting language like Python will.
- For low level languages such as Java/C++, the use of primitive arrays and for-loops would not pose any computational efficiency problems—however, it is still helpful to make use of a linear algebra library to cut down on the number of lines of code you will write.
- Implement a finite difference test to check whether your implementation of backpropagation is correctly computing gradients. If you choose to do this, comment out this functionality once your backward pass starts giving correct results and before submitting to Autolab—since it will otherwise slow down your code.

2.3.1 Command Line Arguments

The autograder runs and evaluates the output from the files generated, using the following command:

```
For Python: $ python neuralnet.py [args...]

For Java: $ javac -cp "./lib/ejml-v0.33-libs/*:./" neuralnet.java  
$ java -cp "./lib/ejml-v0.33-libs/*:./" neuralnet [args...]

For C++: $ g++ -g -std=c++11 -I./lib neuralnet.cpp; ./a.out [args...]

For Octave: $ octave -qH neuralnet.m [args...]
```

Where above [args...] is a placeholder for nine command-line arguments: <train_input> <validation_input> <train_out> <validation_out> <metrics_out> <num_epoch> <hidden_units> <init_flag> <learning_rate>. These arguments are described in detail below:

- 1. <train_input>: path to the training input .csv file (see Section 2.1)
- 2. <validation_input>: path to the validation input .csv file (see Section 2.1)
- 3. <train_out>: path to output .labels file to which the prediction on the *training* data should be written (see Section 2.3.2)
- 4. <validation_out>: path to output .labels file to which the prediction on the *validation* data should be written (see Section 2.3.2)
- 5. <metrics_out>: path of the output .txt file to which metrics such as train and validation error should be written (see Section 2.3.3)
- 6. <num_epoch>: integer specifying the number of times backpropogation loops through all of the training data (e.g., if <num_epoch> equals 5, then each training example will be used in backpropogation 5 times).

- 7. <hidden_units>: positive integer specifying the number of hidden units.
- 8. <init_flag>: integer taking value 1 or 2 that specifies whether to use RANDOM or ZERO initialization (see Section 2.2.1 and Section 2.3)—that is, if init_flag==1 initialize your weights randomly from a uniform distribution over the range [-0.1,0.1] (i.e. RANDOM), if init_flag==2 initialize all weights to zero (i.e. ZERO). For both settings, always initialize bias terms to zero.
- 9. <learning_rate>: float value specifying the learning rate for SGD.

As an example, if you implemented your program in Python, the following command line would run your program with 5 hidden units on the small data provided in the handout for 4 epochs using Zero initialization and a learning rate of 0.1.

```
$ python neuralnet.py smalltrain.csv smallvalidation.csv \
model1train_out.labels model1val_out.labels model1metrics_out.txt \
5 4 2 0.1
```

Linear Algebra Libraries When implementing a neural network, it is often more convenient to have a linear algebra library at your disposal. In this assignment, Java users may use EJML^a and C++ users Eigen^b. Details below. (As usual, Python users have numpy; Octave users have built-in matrix support.)

Java EJML is a pure Java linear algebra package with three interfaces. We strongly recommend using the SimpleMatrix interface. Autolab will use EJML version 3.3. The command line arguments above demonstrate how we will call you code. The classpath inclusion -cp "./lib/ejml-v0.33-libs/*:./" will ensure that all the EJML jars are on the classpath as well as your code.

C++ Eigen is a header-only library, so there is no linking to worry about—just #include whatever components you need. Autolab will use Eigen version 3.3.4. The command line arguments above demonstrate how we will call you code. The argument -I./lib will include the lib/Eigen subdirectory, which contains all the headers.

We have included the correct versions of EJML/Eigen in the handout.tar for your convenience. Do **not** include EJML or Eigen in your Autolab submission tar; the autograder will ensure that they are in place.

```
"https://ejml.org
bhttp://eigen.tuxfamily.org/
```

2.3.2 Output: Labels Files

Your program should write two output .labels files containing the predictions of your model on training data (<train_out>) and validation data (<validation_out>). Each should contain the predicted labels for each example printed on a new line. Use \n to create a new line.

Your labels should exactly match those of a reference implementation – this will be checked by the autograder by running your program and evaluating your output file against the reference solution.

Note: You should output your predicted labels using the same *integer* identifiers as the original training data. You should also insert an empty line (again using \n) at the end of each sequence (as is done in the input data files). The first few lines of an example output file are given below.

```
6
4
2
2
```

2.3.3 Output Metrics

Generate a file where you report the following metrics:

cross entropy After each Stochastic Gradient Descent (SGD) epoch, report mean cross entropy on the training data crossentropy (train) and validation data crossentropy (validation) (See Equation 2.1). These two cross-entropy values should be reported at the end of each epoch and prefixed by the epoch number. For example, after the second pass through the training examples, these should be prefixed by epoch=2. The total number of train losses you print out should equal num_epoch—likewise for the total number of validation losses.

error After the final epoch (i.e. when training has completed fully), report the final training error error (train) and validation error error (validation).

A sample output is given below. It contains the train and validation losses for the first 2 epochs and the final error rate when using the command given above.

```
epoch=1 crossentropy(train): 2.18506276114
epoch=1 crossentropy(validation): 2.18827302588
epoch=2 crossentropy(train): 1.90103257727
epoch=2 crossentropy(validation): 1.91363803461
error(train): 0.728
error(validation): 0.77
```

Take care that your output has the exact same format as shown above. There is an equal sign = between the word epoch and the epoch number, but no spaces. There should be a single space after the epoch number (e.g. a space after epoch=1), and a single space after the colon preceding the metric value (e.g. a space after epoch=1 likelihood(train):). Each line should be terminated by a Unix line ending \n.

2.4 Autolab Submission

You must submit a .tar file named neuralnet .tar containing neuralnet . $\{py \mid m \mid java \mid cpp\}$. You can create that file by running:

```
tar -cvf neuralnet.tar neuralnet.{py|m|java|cpp}
```

from the directory containing your code.

Some additional tips: **DO NOT** compress your files; you are just creating a tarball. Do not use tar -czvf. **DO NOT** put the above files in a folder and then tar the folder. Autolab is case sensitive, so observe that all your files should be named in **lowercase**. You must submit this file to the corresponding homework link on Autolab. The autograder for Autolab prints out some additional information about the tests that it ran. You can view this output by selecting "Handin History" from the menu and then clicking one of the scores you received for a submission. For example on this assignment, among other things, the autograder will print out which language it detects (e.g. Python, Octave, C++, Java).

Python3 Users: Please include a blank file called python3.txt (case-sensitive) in your tar submission and we will execute your submitted program using Python 3 instead of Python 2.7.

Note: For this assignment, you may make up to 10 submissions to Autolab before the deadline, but only your last submission will be graded.