Lab 8 - Neural Networks

In this lab you'll use *neural networks* to classify images using both <u>scikit-learn (https://scikit-learn.org)</u> and <u>PyTorch (https://pytorch.org/)</u>. PyTorch 1.9 or later is assumed to be installed. The goal is for you to see:

- 1. that logistic regression is a special case of neural networks; and
- 2. how to express the same type of network in both scikit-learn and in PyTorch, both shallow (logistic regression) and deep (several layers).

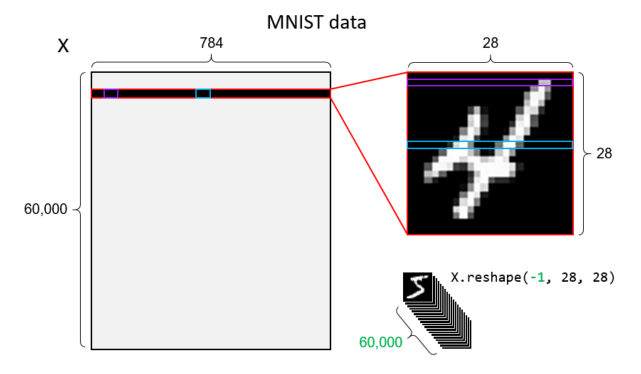
Run the code cell below to import the required packages.

```
In [1]: import numpy as np
    import matplotlib.pyplot as plt
    import sklearn
    import sklearn.preprocessing  # For StandardScaler
    import sklearn.linear_model  # For LogisticRegression
    import sklearn.neural_network  # For MLPClassifier
    import torch
    import warnings
    warnings.filterwarnings("ignore", category=sklearn.exceptions.ConvergenceWa np.set_printoptions(precision=3, suppress=True) # Print as 0.001 instead or
```

1. Digit classification with neural networks in scikit-learn

Exercise 1.1–1.8 ask you to load and train a model on the classic MNIST data set. It's so classic it has its own Wikipedia page (https://en.wikipedia.org/wiki/MNIST_database)! The MNIST data set contains 60,000 training examples and 10,000 test examples. Each example comprises a 784-dimensional feature vector \mathbf{x}_i representing 28x28 grayscale image of a hand-written digit (784 = 28x28) with a label $y_i \in \{0, ..., 9\}$.

Since there are 60,000 training cases, the matrix of training features \mathbf{X} is provided in as a 60000x784 matrix of pixel intensities. Value $X_{i,j} \in \{0,\dots,255\}$ represents the intensity (0=black, 255=white) of pixel number j in training image i. Each 784-dimensional feature vector \mathbf{x}_i can be reshaped into a 28x28 image as depicted below.



Run the code cell below to define a function that will be useful for plotting matrices.

```
In [2]: def plot matrix grid(V):
            Given an array V containing stacked matrices, plots them in a grid layo
            V should have shape (K,M,N) where V[k] is a matrix of shape (M,N).
            assert V.ndim == 3, "Expected V to have 3 dimensions, not %d" % V.ndim
            k, m, n = V.shape
            ncol = 8
                                                         # At most 8 columns
            nrow = min(4, (k + ncol - 1) // ncol)
                                                         # At most 4 rows
            V = V[:nrow*ncol]
                                                         # Focus on just the matric
            figsize = (2*ncol, max(1, 2*nrow*(m/n))) # Guess a good figure shap
            fig, axes = plt.subplots(nrow, ncol, sharex=True, sharey=True, figsize=
            vmin, vmax = np.percentile(V, [0.1, 99.9]) # Show the main range of v
            for v, ax in zip(V, axes.flat):
                img = ax.matshow(v, vmin=vmin, vmax=vmax, cmap=plt.get cmap('gray')
                ax.set xticks([])
                ax.set yticks([])
            fig.colorbar(img, cax=fig.add axes([0.92, 0.25, 0.01, .5]))
                                                                          # Add a c
```

Exercise 1.1 – Load MNIST and plot some digits

The MNIST training data has been provided to you in a file called <code>mnist_train.npz</code>. The file is located in the same directory as this Jupyter Notebook. A <code>npz</code> file is an efficient way to store multiple Numpy arrays in a file. Use Numpy's <code>load</code>

(https://docs.scipy.org/doc/numpy/reference/generated/numpy.load.html) function to open an npz file. When the file is opened, you can think of the file as being a Python dictionary where you can ask for an array by its name (its 'key'). The example below shows how to open the file and list the keys:

```
>>> with np.load("mnist_train.npz") as data:
... print(list(data.keys()))

['X', 'y']
```

(The reason we open the file using a *with*-statement is because once the *with*-statement is complete the file ("file descriptor") is automatically closed, rather than Python trying to keep the file open. This isn't important for the lab *per se*, closing files when you're done with them is just good programming practice!)

Write a few lines of code to load the training data from $mnist_tnin.npz$ and create two global variables X_trn and y_trn to refer to the data you loaded.

```
In [6]: # Your code here. Aim for 3 lines.
with np.load("mnist_train.npz") as data:
    X_trn = data['X']
    y_trn = data['y']
```

Inspect the data by printing information about the arrays.

[252 30 22 119 197 241 253 252 251

- 1. Print the shape and dtype of both your X_{trn} and y_{trn} arrays.
- 2. Print the first five training samples from X_{trn} and y_{trn} arrays.

Since your X_trn array is big, and because most of the first/last pixels in each image are 0 (black), to see any patterns in the features try printing a slice of values taken from the "middle" of each image. For example, pixels 400:415 are roughly from the middle row of each image (similar to blue rectangle in the diagram earlier), so try printing a slice of just those pixels. You should see $\begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix}$

```
In [7]: # Your code for printing shape and dtype here. Aim for 2 lines.
        print("X:", X trn.shape, X trn.dtype)
        print("y:", y_trn.shape, y_trn.dtype)
        # Your code for printing sample values. Aim for 2 lines.
        print(X trn[0:5,400:415])
        print(y_trn[0:5])
        X: (60000, 784) uint8
        y: (60000,) int32
        0 ]]
                                81 240 253 253 119
                                                     25
                                                          0
                                                                       0]
                0
                     0
                         0
                             0
                                                               0
                                                                   0
         [253 190
                     0
                         0
                             0
                                 0
                                     0
                                          0
                                              0
                                                  0
                                                      0
                                                          0 255 253 1961
               47 49 116 144 150 241 243 234 179 241 252
                                                              40
            0
                                                                       01
                     0
                         0
                            80 240 251 193 23
                                                  0
                                                          0
                                                               0
                                                                       01
```

Plot a few digits to see what they look like. Use the *plot_matrix_grid* function defined earlier. To do this, you'll need to reshape the array referred to by your *X_trn* variable so that the plotting code knows the images have shape 28x28 rather than being just 784-dimensional vectors.

77

0

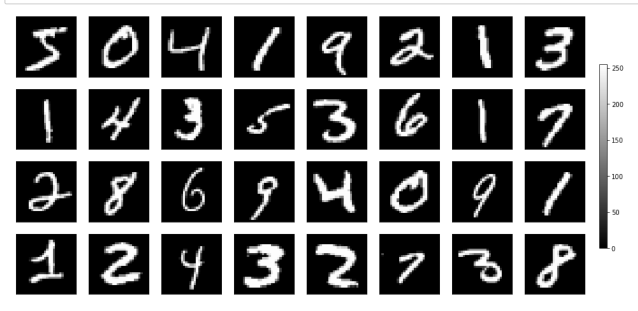
0

0

0]]

[5 0 4 1 9]

```
In [8]: # Your code here. Aim for 1-2 lines.
plot_matrix_grid(X_trn.reshape(-1, 28, 28))
```



Look at the patterns you printed when inspecting the X_{trn} variable earlier, and make sure you see where they come from in the first five images plotted above.

If you want to see more of the MNIST training digits, rather than just the first few, you can try plotting different "slices" of the X_trn variable, such as $X_trn[100:]$ to start plotting at the 101st training example. (You still have to reshape the resulting array, of course.)

Finally, **load the MNIST test data** from the file $mnist_test.npz$, just like you did for the training data. Create global variables X_tst and y_tst to refer to the arrays that you loaded. These arrays will be used to evaluate test-time accuracy later on.

```
In [9]: # Your code here. Aim for 3 lines.
with np.load("mnist_test.npz") as data:
    X_tst = data['X']
    y_tst = data['y']
```

Exericise 1.2 - Preprocess the MNIST data

Certain models trained on MNIST work better when the features are normalized. Use scikit-learn to normalize the MNIST data using scaling, such as the *StandardScaler*. (You can just treat the pixels as independent features, nothing fancy.)

Write a few lines of code to normalize both you X_tm and X_t variables. You can just over-write those variables with the new (normalized) feature arrays, and discard the original unscaled data.

```
In [10]: # Your code here. Aim for 3-4 lines.
    scaler = sklearn.preprocessing.StandardScaler()
    scaler.fit(X_trn)
    X_trn = scaler.transform(X_trn)
    X_tst = scaler.transform(X_tst)
```

Plot the rescaled training digits using the *plot_matrix_grid* function.

```
In [12]: # Your code here. Aim for 1-2 lines.
plot_matrix_grid(X_trn.reshape(-1, 28, 28))
```



Exericise 1.3 – Train multinomial logistic regression on MNIST

Train a LogisticRegression (https://scikit-

you understand why?

<u>learn.org/stable/modules/generated/sklearn.linear_model.LogisticRegression.html</u>) object to classify MNIST digits. Use *random_state=*0 and default settings otherwise.

```
In [13]: # Your code here. Aim for 2-3 lines.
lr = sklearn.linear_model.LogisticRegression(C=0.01, random_state=0) # C=0
lr.fit(X_trn, y_trn);
```

You can use the score (https://scikit-

<u>learn.org/stable/modules/generated/sklearn.linear_model.LogisticRegression.html#sklearn.line</u> method of the *LogisticRegression* object to compute the accuracy as a number in the range [0.0, 1.0]. Figure out how to convert that number (e.g., 0.934) into an error rate percentage (e.g., 6.6%).

Print the training *error rate* and testing *error rate* of your logistic regression model on the MNIST data set. Your output should be in the form:

```
X.XX% training error
X.XX% testing error
```

```
In [14]: # Your code here. Aim for 2-4 lines.
print("%.2f%% training error" % (100*(1-lr.score(X_trn, y_trn))))
print("%.2f%% testing error" % (100*(1-lr.score(X_tst, y_tst))))
6.40% training error
7.40% testing error
```

How does the testing error rate you see compare to some of the error rates mentioned on the MNIST Wikipedia page (https://en.wikipedia.org/wiki/MNIST_database)?

Print the predicted class probabities of the **first five examples** in the training set. Use the *predict_proba* method of your *LogisticRegression* object. The first row of output should look something like:

```
[0.001 0. 0. 0.203 0. 0.796 0. 0. 0. ]
```

From the above probabilities we can see that the model thinks the first digit in the training set is *probably* digit "5" but *might also* be digit "3".

```
In [15]: # Your code here. Aim for 1-2 lines.
         print(lr.predict_proba(X_trn[:5]))
                         0.001 0.231 0.
          [[0.001 0.
                                             0.766 0.
                                                          0.001 0.
                                                                       0.
                                                                            ]
           [1.
                                0.
                                      0.
                                             0.
                                                   0.
                                                          0.
                                                                0.
                                                                       0.
           [0.
                  0.001 0.009 0.086 0.873 0.
                                                   0.
                                                          0.025 0.001 0.006]
           [0.
                  0.965 0.019 0.002 0.
                                                                0.013 0.
                                             0.
                                                   0.
                                                          0.015 0.002 0.965]]
           [0.
                                0.
                                      0.018 0.
                                                   0.
```

Exericise 1.4 – Visualize the weights of your logistic regression model

The logisitic regression model you trained in Exercise 1.3 has a $coef_{-}$ attribute. This attribute is the array of weights \mathbf{W} seen in Lecture 4 (e.g. slide 28). For the MNIST data, this matrix has shape (10, 784), because there are 10 output classes and 784=28x28 pixels. Weight $W_{k,j}$ is the weight with which of pixel j contributes to output class k.

You are asked to visualize the weights using *plot_matrix_grid*. You may need to reshape the weight matrix to do this. The first two outputs, corresponding to predictin digit "0" and predicting digit "1" should look something like this:

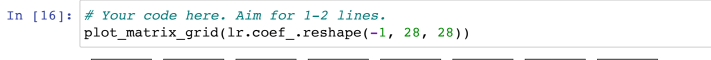


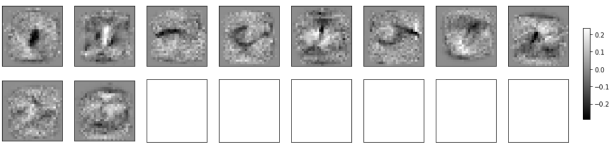


Notice how the pattern for "0" is has strong negative weights in the center: that's because if there are white pixels in the center, it's unlikely that the image represents digit "0"!

If your weight patterns appear **noisier** than above, try repeating Exercise 1.3 but weaken *LogisticRegression*'s L2 penalty by a factor of 100 from its default. Take note of any change in training/test accuracy, too.

Write a few lines of code to plot the weights and see what patterns they contain. You should see ten patterns. (Don't worry if the last few grid entries are just white boxes.)





When an input image (of a hand-written digit) causes one of these patterns to have a large positive response (strong activation), then the corresponing class $\{0, 1, 2, \dots, 9\}$ will be given a high probability by the final softmax operation.

Exericise 1.5 – Train a neural network on MNIST with zero hidden layers

Train a neural network on MNIST using the <u>sklearn.neural network.MLPClassifier (https://scikitlearn.org/stable/modules/generated/sklearn.neural network.MLPClassifier.html)</u> class.

A neural network has *many* more hyperparameters to configure. Configure your neural network as follows:

- Ask for no hidden layers. You can do this by specifying an empty tuple () for the hidden_layer_sizes argument. This will create a neural network where the 784 input features are directly 'connected' to the 10 output predictions, which in this case corresponds to the multinomial logistic regression you did in Exercise 1.4.
- Use the sqd solver. This means stochastic gradient descent that we saw in Lecture 1.
- Use a batch size of 100. This means that at each step of SGD the gradient will be computed from only 100 of the 60,000 training cases. This is also callsed a "mini-batch". The SGD algorithm works by starting with the firs 100, then the next 100, and then it gets to the last 100 in the training set it starts from the beginning again.

- Use *max_iter*=10. This causes the training to stop after SGD has passed over all 60,000 training cases exactly 10 times.
- Use *learning_rate_init*=0.01, which determines the step size for SGD once it has computed a gradient.
- Use *momentum*=0.9, which speeds up training.
- Use random_state=0 for reproducibility
- Use *verbose*=True to see progress printed out. Each time it prints "Iteration X" it means SGD has made another pass over all 60,000 training examples.

```
In [17]: # Your code here. Aim for 1-2 lines, plus whatever line wrapping you need f
         mlp = sklearn.neural_network.MLPClassifier(hidden_layer_sizes=(),
                                                     solver='sgd',
                                                     batch size=100,
                                                     max_iter=10,
                                                     learning rate init=.01,
                                                     momentum=0.9,
                                                     verbose=True,
                                                     random state=0)
         mlp.fit(X_trn, y_trn);
         Iteration 1, loss = 0.40770306
         Iteration 2, loss = 0.30859873
         Iteration 3, loss = 0.29337169
         Iteration 4, loss = 0.28490254
         Iteration 5, loss = 0.27543585
         Iteration 6, loss = 0.26958339
         Iteration 7, loss = 0.26545986
         Iteration 8, loss = 0.26427636
         Iteration 9, loss = 0.26236903
         Iteration 10, loss = 0.25963997
```

Print the training error rate and test error rate of your neural network classifier, just like you did for logistic regression.

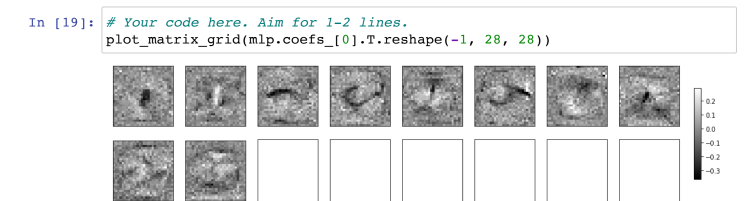
```
In [18]: # Your code here. Aim for 2-4 lines.
print("%.2f%% training error" % (100*(1-mlp.score(X_trn, y_trn))))
print("%.2f%% testing error" % (100*(1-mlp.score(X_tst, y_tst))))
6.84% training error
7.68% testing error
```

Exericise 1.6 – Visualize the weights of a neural network (no hidden layers)

The MLPClassifier object has a coefs_ attribute that works just like the coef_ attribute that contained coefficient matrix \mathbf{W} of LogisticRegression, except that for a neural network there are two differences:

- 1. $coefs_i$ is a *list* of coefficient matrices, so $coefs_i[0]$ is $\mathbf{W}^{(1)}$, the coefficient matrix of the *first layer*. Since the neural network you trained in Exercise 1.5 has no hidden layers, this $\mathbf{W}^{(1)}$ matrix holds the same weights as the \mathbf{W} matrix for LogisticRegression.
- 2. The weight matrix for *MLPClassifier* has a different layout: it is 784x10 rather than 10x784. Do you now how to account for this?

Write a few lines of code to repeat Exercise 1.4 but this time with the neural network weights.



If your patterns look streaky then you may need to try transposing your weight matrix to account for the different layout.

Exericise 1.7 – Train and visualize the weights of a neural network with 1 hidden layer

Here you're asked to train a neural network like you did in Exercise 1.5, but this time **add a hidden layer with 16 'tanh' hidden units** to your neural network. Then you'll visualize the weights of this network.

Read the documentation for MLPClassifier (https://scikit-

<u>learn.org/stable/modules/generated/sklearn.neural_network.MLPClassifier.html)</u> to learn how to do specify a hidden layer. (*Note:* In Python if you want to create a *tuple* object with only one item in it, you can use (*item,*) with an extra comma, rather than (*item*), which Python interprets to just be regular parentheses.) All the other hyperparameters can stay the same as Exercise 1.5.

Write a few lines of code to train a new neural network, this time with 16 *tanh* hidden units. In other words, this will be a 784-16-10 neural network where the hidden layer uses *tanh* activations.

```
In [20]: # Your code here. Aim for 1-2 lines, plus whatever line wrapping you need f
         mlp h16 = sklearn.neural network.MLPClassifier(hidden layer sizes=(16,),
                                                         activation='tanh',
                                                         solver='sgd',
                                                         batch size=100,
                                                         max iter=10,
                                                         learning_rate_init=.01,
                                                         momentum=0.9,
                                                         verbose=True,
                                                         random_state=0)
         mlp_h16.fit(X_train, y_train);
         Iteration 1, loss = 1.90614999
         Iteration 2, loss = 1.65236888
         Iteration 3, loss = 1.55716868
         Iteration 4, loss = 1.39333193
         Iteration 5, loss = 1.43590724
         Iteration 6, loss = 1.42916529
         Iteration 7, loss = 1.31703452
         Iteration 8, loss = 1.34322581
         Iteration 9, loss = 1.28388686
         Iteration 10, loss = 1.20374607
```

Print the training error rate and test error rate of your neural network classifier, just like you did for logistic regression. How does your error rate compare to multinomial logistic regression? (Exercises 1.3 and 1.5)

```
In [21]: # Your code here. Aim for 2-4 lines.
print("%.2f%% training error" % (100*(1-mlp_h16.score(X_trn, y_trn))))
print("%.2f%% testing error" % (100*(1-mlp_h16.score(X_tst, y_tst))))
41.44% training error
41.61% testing error
```

Plot the first-layer weights $\mathbf{W}^{(1)}$ of your neural network using the *plot_matrix_grid* function, just in Exercise 1.6.

```
In [22]: # Your code here. Aim for 1-2 lines.
plot_matrix_grid(mlp_h16.coefs_[0].T.reshape(-1, 28, 28))
```

Notice that there are now 16 patterns, not 10, and they no longer seem to correspond to the digits $\{0, 1, \dots, 9\}$ in any particular order. Do you understand why?

Plot the second-layer weights $W^{(2)}$ of your neural network using the plot_matrix_grid function.

However, this time if you inspect the shape of the second weight matrix, <code>coefs_[1]</code>, you'll see that it has shape (16, 10), and so it cannot be reshaped into a 28x28 pattern. In fact the second layer has only dimension: the "hidden layer" is just a vector of 16 values (the 16 tanh-transformed activations of the first-layer patterns). Each of the 10 output units has 16 weights contributing to it, rather than 784 weights like in Exercise 1.6.

Figure out how to reshape the weight matrix so that when you call *plot_matrix_grid* you see a grid of 1x16 weight vectors, like the two examples below:

```
In [23]: # Your code here. Aim for 1-2 lines.
plot_matrix_grid(mlp_h16.coefs_[1].T.reshape(-1, 1, 16))
```

Exericise 1.8 – Train a neural network with lots of hidden units

Repeat Exercise 1.7 but with two hidden layers having **100 and 50 hidden units** respectively. This time use *relu* activations. All other hyperparameters can stay the same.

Write a few lines of code to train the model here.

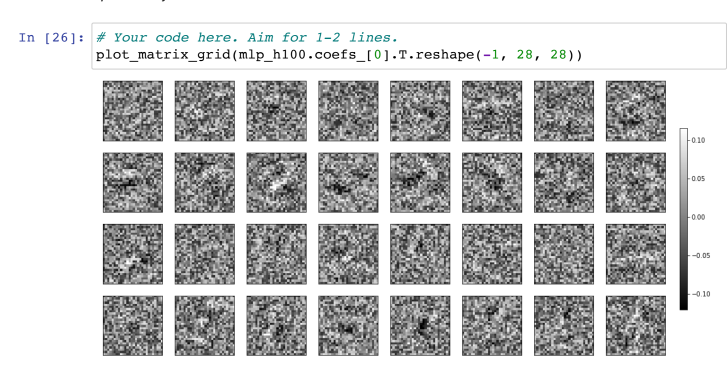
```
# Your code here. Aim for 1-2 lines, plus whatever line wrapping you need f
In [24]:
         mlp h100 = sklearn.neural network.MLPClassifier(hidden layer sizes=(100,50)
                                                          activation='relu',
                                                          solver='sgd',
                                                          batch size=100,
                                                          max iter=10,
                                                          learning rate init=.001,
                                                          momentum=0.9,
                                                          verbose=True,
                                                          random state=0)
         mlp h100.fit(X trn, y trn);
         Iteration 1, loss = 0.84550457
         Iteration 2, loss = 0.32680824
         Iteration 3, loss = 0.25102323
         Iteration 4, loss = 0.21264192
         Iteration 5, loss = 0.18728786
         Iteration 6, loss = 0.16855373
         Iteration 7, loss = 0.15370994
         Iteration 8, loss = 0.14172767
         Iteration 9, loss = 0.13166153
         Iteration 10, loss = 0.12283612
```

Print the training and testing error rates here. How do they compare to earlier models?

```
In [25]: # Your code here. Aim for 2-4 lines.
print("%.2f%% training error" % (100*(1-mlp_h100.score(X_trn, y_trn))))
print("%.2f%% testing error" % (100*(1-mlp_h100.score(X_tst, y_tst))))

3.31% training error
4.50% testing error
```

Plot the first-layer weights $\mathbf{W}^{(1)}$ of your neural network here. Are the pattern detectors here qualitatively different than for earlier models?



Don't bother plotting the 2nd and 3rd layer weights, they are high-dimensional and hard to interpret.

2. Neural networks in PyTorch

Exercise 2.1–2.3 ask you to train a simple neural network in PyTorch
(https://pytorch.org/docs/stable/index.html). Here you'll use PyTorch to train an MNIST classifier using the same MNIST data that you already preprocess in Part 1. The goal is just to get you familiar with PyTorch basics and how they compare to scikit-learn.

PyTorch is a deep learning framework like TensorFlow. PyTorch tends to be popular with deep learning researchers because it's very flexible for trying new ideas. TensorFlow is also flexible but is designed in such a way that it's more popular for companies trying to deploy high-performance models (in the cloud etc). Both can be used for research, of course!

Exericise 2.1 – Convert MNIST from Numpy arrays to PyTorch tensors

PyTorch has its own Numpy-like array class, called *Tensor*. In order to train a PyTorch model, you must first convert the Numpy arrays. PyTorch understands Numpy arrays, so this is easy. The only tricky part is that, in order to be fast and not waste memory, PyTorch tends to be more picky about the *dtype* of the arrays you give it.

Write a few lines of code to create four global variables: *X_trn_torch*, *y_trn_torch*, *X_tst_torch*, *y_tst_torch* that are PyTorch versions of your preprocessed MNIST training data from Part 1. The *X* tensors should have *dtype* float32, and the *y* tensors should have *dtype* int64.

```
In [27]: # Your code here. Aim for 2-4 lines.
X_trn_torch = torch.tensor(X_trn.astype(np.float32))
y_trn_torch = torch.tensor(y_trn.astype(np.int64))
X_tst_torch = torch.tensor(X_tst.astype(np.float32))
y_tst_torch = torch.tensor(y_tst.astype(np.int64))
```

Run the code cell below to check your answer.

```
In [28]: assert 'X_trn_torch' in globals(), "You didn't declare a X_trn_torch variab
          assert 'y_trn_torch' in globals(), "You didn't declare a y_trn_torch variab
assert 'X_tst_torch' in globals(), "You didn't declare a X_tst_torch variab
          assert 'y_tst_torch' in globals(), "You didn't declare a y_tst_torch variab
          assert isinstance(X trn torch, torch.Tensor)
          assert isinstance(y trn torch, torch.Tensor)
          assert isinstance(X tst torch, torch.Tensor)
          assert isinstance(y_tst_torch, torch.Tensor)
          assert X trn torch.dtype == torch.float32
          assert y trn torch.dtype == torch.int64
          assert X trn torch.shape == (60000,784)
          assert y trn torch.shape == (60000,)
          assert X tst torch.dtype == torch.float32
          assert y tst torch.dtype == torch.int64
          assert X tst torch.shape == (10000,784)
          assert y tst torch.shape == (10000,)
          print("Correct!")
```

Correct!

Exericise 2.2 – Train a PyTorch neural network *without* hidden layers

This exercise only asks you to **run existing code** so that you learn how PyTorch works. The code in this cell defines a simple logistic model, and then you are asked to modify the code to add hidden layers to the network.

Useful documentation for understanding the code that you see:

- torch.nn (https://pytorch.org/docs/stable/nn.html) (neural network)
- torch.optim (https://pytorch.org/docs/stable/optim.html) (optimizers such as SGD)

Here are some comments to help you understand the "starter code" below:

- A neural network is a sequence of non-linear transformations, so PyTorch provides a
 <u>Sequential</u>
 (https://pytorch.org/docs/stable/generated/torch.nn.Sequential.html#torch.nn.Sequential)
 class that accepts a list of desired transformations.
- In a standard neural network, the transformations are just linear, i.e. $\mathbf{W}\mathbf{x} + \mathbf{b}$, and in PyTorch this is implemented by a <u>Linear</u> (https://pytorch.org/docs/stable/generated/torch.nn.Linear.html#torch.nn.Linear) class where constructing one of these objects with Linear(D, M) tells the new object that it should be expecting an D-dimensional input and transform it into a M-dimensional output. To do this, the Linear object will create its own parameter matrix $\mathbf{W} \in \mathbb{R}^{M \times D}$ and bias vector $\mathbf{b} \in \mathbb{R}^{M}$.
- In PyTorch, the <u>CrossEntropyLoss</u>
 (https://pytorch.org/docs/stable/generated/torch.nn.CrossEntropyLoss.html) class conveniently combines applying a softmax and then computing the negative log likelihood, so you don't explicitly apply softmax while training. Once you have a *CrossEntropyLoss* object, you can call it with your predictions and targets (both vectors), and it will compute the negative log likelihood, which is just one number (a scalar).

Run the code cell below to define a simple 784-10 neural network (i.e. logistic regression).

Run the code cell below to define some objects and variables needed for training the neural network.

```
In [30]: # Create an object that can compute "negative log likelihood of a softmax"
loss = torch.nn.CrossEntropyLoss()

# Use stochastic gradient descent to train the model
optimizer = torch.optim.SGD(model.parameters(), lr=0.01, momentum=0.9)

# Use 100 training samples at a time to compute the gradient.
batch_size = 100

# Make 10 passes over the training data, each time using batch_size samples
num_epoch = 10
next_epoch = 1
```

Run the code cell below to train the neural network using stochastic gradient descent (SGD). Note that if you re-run this code cell multiple times it will "continue" training from the current parameters, and if you want to "reset" the model you need to re-run the earlier code cell that defined the model!

```
In [31]: for epoch in range(next_epoch, next_epoch+num_epoch):
             \# Make an entire pass (an 'epoch') over the training data in batch size
             for i in range(0, len(X_trn), batch_size):
                 X = X trn torch[i:i+batch size] # Slice out a mini-batch of fea
                 y = y_trn_torch[i:i+batch_size] # Slice out a mini-batch of tar
                                                    # Make predictions (final-layer
                 y pred = model(X)
                 l = loss(y pred, y)
                                                   # Compute loss with respect to
                 model.zero grad()
                                                   # Reset all gradient accumulato
                                                   # Compute gradient of loss wrt
                 1.backward()
                 optimizer.step()
                                                    # Use the gradients to take a s
             print("Epoch %2d: loss on final training batch: %.4f" % (epoch, l.item(
         print("Epoch %2d: loss on test set: %.4f" % (epoch, loss(model(X_tst_torch))
         next epoch = epoch+1
         Epoch 1: loss on final training batch: 0.7097
         Epoch 2: loss on final training batch: 0.8752
         Epoch 3: loss on final training batch: 0.3313
         Epoch 4: loss on final training batch: 0.3312
         Epoch 5: loss on final training batch: 0.3239
         Epoch 6: loss on final training batch: 0.3168
         Epoch 7: loss on final training batch: 0.3128
         Epoch 8: loss on final training batch: 0.3074
         Epoch 9: loss on final training batch: 0.3044
```

Run the code cell below to retrieve the PyTorch model's parameters, convert them back to Numpy, and plot them like before.

Epoch 10: loss on final training batch: 0.2990

Epoch 10: loss on test set: 0.3211

```
In [32]: W, b, *_ = model.parameters()
W = W.detach().numpy()
plot_matrix_grid(W.reshape(-1, 28, 28))
```

Exericise 2.3 – Train a PyTorch neural network *with* hidden layers

Using Exercise 2.2 as a starting point, write new code to **implement a 784-100-50-10 neural network** with *relu* **activations** just like you did in Exercise 1.8, but now implemented with PyTorch.

To do this, you will need to:

- Create a new model object that has more sequential steps to it, including the Linear and ReLU (https://pytorch.org/docs/stable/generated/torch.nn.ReLU.html) objects.
- 2. Create a new *optimizer* object that knows about your new model's parameters.

If you succeed, you should be able to get the training loss to go to zero, especially if you run the training loop code cell extra times (i.e. more than 10 epochs total). But what happens with the test set loss, as you continue training?

We will do more PyTorch in the next lab, with convolutional neural networks.

```
In [33]: # Your PyTorch to create the model and optimizer here.
torch.manual_seed(0) # Ensure model weights initialized with same random nu

# Create an object that can evaluate a neural network.
model = torch.nn.Sequential(
    torch.nn.Linear(28*28, 100),
    torch.nn.ReLU(),
    torch.nn.Linear(100, 50),
    torch.nn.ReLU(),
    torch.nn.ReLU(),
    torch.nn.Linear(50, 10),
)

optimizer = torch.optim.SGD(model.parameters(), lr=0.01, momentum=0.9)
```

```
In [34]: # Your PyTorch training loop here.
for epoch in range(num_epoch):
    for i in range(0, len(X_trn), batch_size):
        X = X_trn_torch[i:i+batch_size]
        y = y_trn_torch[i:i+batch_size]
        y_pred = model(X)
        1 = loss(y_pred, y)
        model.zero_grad()
        l.backward()
        optimizer.step()
    print("Epoch %d final minibatch had loss %.4f" % (epoch+1, l.item()))
```

```
Epoch 1 final minibatch had loss 0.4997
Epoch 2 final minibatch had loss 0.1830
Epoch 3 final minibatch had loss 0.3667
Epoch 4 final minibatch had loss 0.0848
Epoch 5 final minibatch had loss 0.0846
Epoch 6 final minibatch had loss 0.0646
Epoch 7 final minibatch had loss 0.0538
Epoch 8 final minibatch had loss 0.0349
Epoch 9 final minibatch had loss 0.0103
Epoch 10 final minibatch had loss 0.0075
```

Finally, use the *named_parameters* method, available on all PyTorch <u>Module</u> (https://pytorch.org/docs/stable/generated/torch.nn.Module.html) objects, to print the name and shape of each parameter tensor in the neural network. Your output should look something like:

```
0.weight torch.Size([?])
0.bias torch.Size([?])
...

In [39]: # Your code here. Aim for 2-3 lines.
for name, param in model.named_parameters():
    print( "%s %s" % (name, param.size()) )

0.weight torch.Size([100], 784])
0.bias torch.Size([100])
2.weight torch.Size([50], 100])
2.bias torch.Size([50])
4.weight torch.Size([10, 50])
4.bias torch.Size([10])
```