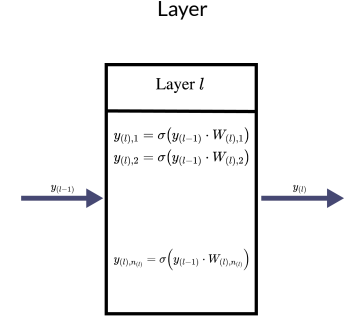
How does the NN "learn" the transformations?

The matrix \mathbf{W} contains the "patterns" that serve to recognize the synthetic features created by each layer



- $\mathbf{W}_{(l),j}$ are the weights /pattern for feature $\mathbf{y}_{(l),j}$
- How are these patterns discovered?

The answer is: exactly as we did in Classical Machine Learning

• Define a loss function that is parameterized by W:

$$\mathcal{L} = L(\hat{\mathbf{y}}, \mathbf{y}; \mathbf{W})$$

- Per example loss $\mathcal{L}^{(i)}$
- Average loss $\mathcal{L} = \frac{1}{m} \sum_{i=1}^{m} \mathcal{L}^{(i)}$
- ullet Our goal is to find $f W^*$ the "best" set of weights

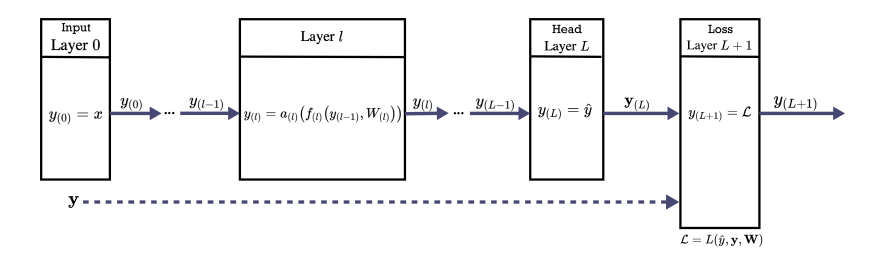
$$\mathbf{W}^* = \operatorname*{argmin}_W L(\hat{\mathbf{y}}, \mathbf{y}; \mathbf{W})$$

• Find \mathbf{W}^* using Gradient Descent!

Very much in spirit of the multi-layer architecture

• We add a new layer (L+1) to compute the loss \mathcal{L} !

Additional Loss Layer (L+1)



Gradient Descent review

Gradient Descent is an iterative method for finding the minimum of a function. <!--EdX: Omit this from EdX: can't refer to prior course

See the <u>Gradient Descent lecture (Gradient Descent.ipynb)</u> in the Classical ML part of the course for more details -->

Let's review Gradient Descent using our current notation

- ullet We start with an initial guess for f W and iteratively improve it.
- Compute the loss $\mathcal L$ given the current $\mathbf W$
 - Average loss of the m examples in the training examples
- Compute the gradient

$$rac{\partial \mathcal{L}}{\partial W}$$

- ullet Update f W in the direction of the *negative* of the gradient
- Scaled by a learning rate α

$$\mathbf{W} = \mathbf{W} - lpha * rac{\partial \mathcal{L}}{\partial W}$$

A unit change in \mathbf{W} increases \mathcal{L} by $\frac{\partial \mathcal{L}}{\partial W}$

- ullet That's why there is a negative sign: we proceed in the direction *opposite* the one that increases ${\cal L}$
- ullet We move only a fraction $lpha \leq 1$ of the (negative) of the gradient
- To avoid the possibility of over-shooting the minimum

 ${f W}$ is a multi-dimensional vector, not a scalar

- So the gradient is multi-dimensional
- We will formally discuss Matrix Gradients in a later lecture
 - lacktriangledown For now: we compute the derivative with respect to each element of f W and arrange in a matrix

We can approximate ${\cal L}$ by sampling from the m training examples

- ullet Choose a random subset (of size $m' \leq m$) of examples: $I = \{i_1, \dots, i_{m'}\}$
- ullet Approximate ${\cal L}$ on I

$$\mathcal{L} pprox rac{1}{|I|} \sum_{i \in I} \mathcal{L^{(i)}}$$

Minibatch gradient descent

The average loss $\mathcal L$ is defined over all m training examples.

- This can be expensive to compute when m is large.
- The gradient can be approximated by sampling from the m training examples
 - Choose a random subset (of size $m' \leq m$) of examples:

$$I=\{i_1,\ldots,i_{m'}\}$$

■ Approximate \mathcal{L} on I

$$\mathcal{L} pprox rac{1}{|I|} \sum_{i \in I} \mathcal{L}^{(\mathbf{i})}$$

Minibatch gradient descents divides the m examples into chunks (mini-batches) and approximates the gradient

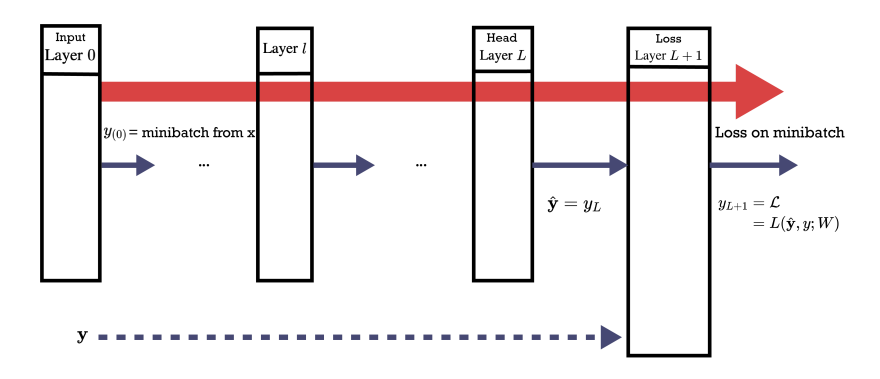
- ullet Divides the m training examples
- ullet Into b=m/m' disjoint batches of size $m'\leq m$ examples each

We can approximate the gradient and update our guess of ${f W}$ on each mini-batch

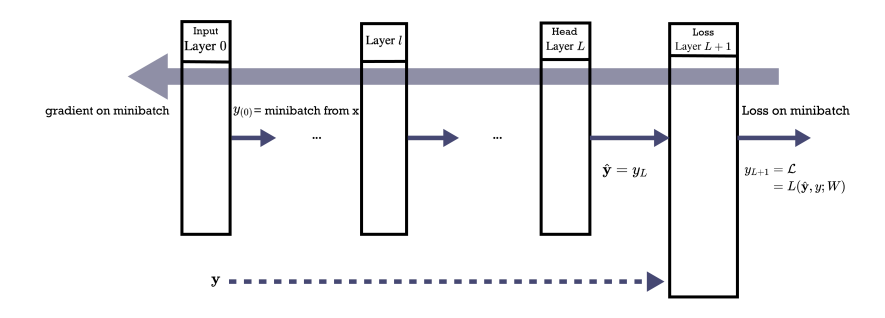
$$\mathbf{W} = \mathbf{W} - lpha * rac{\partial \mathcal{L}}{\partial W}$$

An **epoch** is defined as the processing of all m examples (using b batches of size m^\prime)

Minibatch: Forward Pass From minibatch to Loss



Minibatch: Backwards Pass From minibatch Loss to Gradient



During one epoch, the gradient gets updated b times

- ullet Contrast to the single update when there is a single batch ($m^\prime=m$)
- May be faster as updates occur more frequently

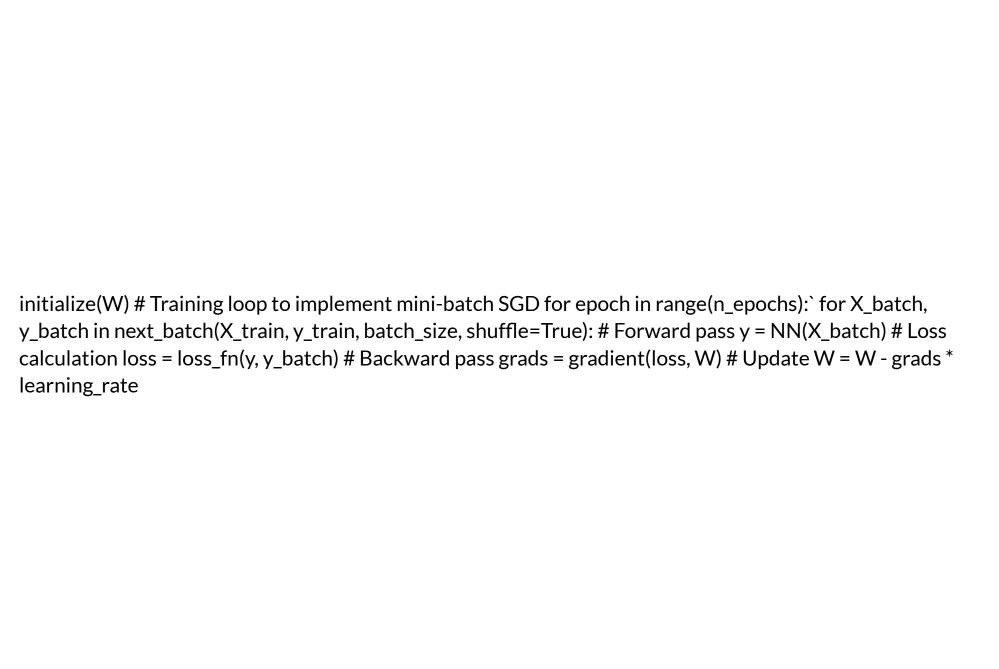
The Training loop

Gradient Descent is an iterative process

- Iterate over multiple epochs
- Within an epoch
 - Iterate over mini-batches

This iterative process is called the *training loop*.

Here is some pseudo-code:



It used to be the case that this fairly standard training loop was coded for each problem.
Just as sklearn wrapped common code into a high-level API
We will use a toolkit that hides the training loop behind a high level API

Scaling the inputs

Many times in this course we have pointed out that some models are scale sensitive.

Neural Networks are not mathematically sensitive but tend to be so in practice.

It is *highly recommended* to scale your data so their absolute values are around 1.0 or at least somewhat small.

Gradient Descent is the root of the problem:

- Two features on different scales can cause the optimizer to favor one over the other
- Activations can saturate
 - Output of dot product (Dense layer) is in the "flat" area of the activation
 - Zero derivative: no learning
- The Loss may be large in initial epochs when the target values are too different from the dot products
 - Large gradients: unstable learning
 - Weights are typically initialized to values less than 1.0, leading to small dot products

Remember: if you re-scale the inputs, you will need to invert the transformation when communicating the results.

```
In [4]: print("Done")
```

Done