# Train, Validate, Test



Recap of Basic Neural Networks (and some Deep Network Fundamentals)

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- Supervised Learning learn to predict an output when given an input vector
- Unsupervised Learning discover a good internal representation of the input
- Reinforcement Learning learn to select an action to maximize the expectation of future rewards (payoff)
- Self-supervised Learning learn with targets induced by a prior on the unlabelled training data
- Semi-supervised Learning learn with few labelled examples and many unlabelled ones

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#### Two Types of Supervised Learning

- Regression: The machine is asked predict k numerical values given some input. The machine is a function  $f : \mathbb{R}^n \to \mathbb{R}^k$ .
- Classification: The machine is asked to specify which of k categories some input belongs to.
  - Multiclass classification target is one of the k classes
  - Multilabel classification target is some number of the k classes
  - In both cases, the machine is a function f : ℝ<sup>n</sup> → {1, ..., k} (although it is most common for the learning algorithm to actually learn f̂ : ℝ<sup>n</sup> → ℝ<sup>k</sup>).
- Note that there are lots of exceptions in the form the inputs (and outputs) can take though! We'll see lots of variations in the coming weeks.

- Start by choosing a model-class: ŷ = f(x; W) where the model-class f is a way of using some numerical parameters, W, to map each input vector x to a predicted output ŷ.
- Learning means adjusting the parameters to reduce the discrepancy between the true target output y on each training case and the output ŷ, predicted by the model.



where f and g are activation functions.

- Identity
- Sigmoid (aka Logistic)
- Hyperbolic Tangent (tanh)
- Rectified Linear Unit (ReLU) (aka Threshold Linear)

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### Final layer activations

#### $\hat{y} = g(\boldsymbol{W}^{(2)}f(\boldsymbol{W}^{(1)}\boldsymbol{x}))$

- What form should the final layer function g take?
- It depends on the task (and on the chosen loss function)...
  - For regression it is typically linear (e.g. identity), but you might choose others if you say wanted to clamp the range of the network.
  - For binary classification (MLP has a single output), one would choose Sigmoid
  - For multilabel classification, typically one would choose Sigmoid
  - For multiclass classification, typically you would use the Softmax function

#### Softmax

The softmax is an activation function used at the output layer of a neural network that forces the outputs to sum to 1 so that they can represent a probability distribution across a discrete mutually exclusive alternatives.

$$\operatorname{softmax}(\boldsymbol{z})_i = \frac{e^{z_i}}{\sum_{j=1}^{K} e^{z_j}} \qquad \forall i = 1, 2, \dots, K$$

- Note that unlike the other activation functions you've seen, softmax makes reference to all the elements in the output.
- The output of a softmax layer is a set of positive numbers which sum up to 1 and can be thought of as a probability distribution.
- Note:

$$\frac{\partial \operatorname{softmax}(\boldsymbol{z})_i}{\partial z_i} = \operatorname{softmax}(z_i)(1 - \operatorname{softmax}(z_i))$$
$$\frac{\partial \operatorname{softmax}(\boldsymbol{z})_i}{\partial z_j} = \operatorname{softmax}(z_i)(1(i = j) - \operatorname{softmax}(z_j))$$
$$= \operatorname{softmax}(z_i)(\delta_{ij} - \operatorname{softmax}(z_j))$$

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#### Ok, so let's talk loss functions

- The choice of loss function depends on the task (e.g. classification/regression/something else)
- The choice also depends on the activation function of the last layer
  - For numerical reasons (see Log-Sum-Exp in a few slides) many times the activation is computed directly within the loss rather than being part of the model
  - Some classification losses require *raw outputs* (e.g. a linear layer) of the network as their input
    - These are often called unnormalised log probabilities or logits
    - An example would be hinge-loss used to create a Support Vector Machine that maximises the margin — e.g.:
       ℓ<sub>hinge</sub>(ŷ, y) = max(0, 1 - y · ŷ) with a true label, y ∈ {-1,1}, for binary classification.
- There are many different loss functions we might encounter (MSE, Cross-Entropy, KL-Divergence, huber, L1 (MAE), CTC, Triplet, ...) for different tasks.

#### The Cost Function (measure of discrepancy)

Recall from Foundations of Machine Learning:

- Mean Squared Error (MSE) loss for a single data point (here assumed to be a vector, but equally applicable to a scalar) is given by *ℓ*<sub>MSE</sub>(ŷ, y) = ∑<sub>i</sub>(ŷ<sub>i</sub> - y<sub>i</sub>)<sup>2</sup> = (ŷ - y)<sup>T</sup>(ŷ - y)
- We often multiply this by a constant factor of <sup>1</sup>/<sub>2</sub> can anyone guess/remember why?
- $\ell_{MSE}(\hat{y}, y)$  is the predominant choice for regression problems with linear activation in the last layer
- For a classification problem with Softmax or Sigmoidal (or really anything non-linear) activations, MSE can cause slow learning, especially if the predictions are very far off the targets
  - Gradients of  $\ell_{MSE}$  are proportional to the difference in target and predicted multiplied by the gradient of the activation function<sup>1</sup>
  - The Cross-Entropy loss function is generally a better choice in this case

<sup>1</sup>http://neuralnetworksanddeeplearning.com/chap3.html Jonathon Hare COMP6258

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### Binary Cross-Entropy

For the binary classification case:

$$\ell_{BCE}(\hat{y}, y) = -y \log(\hat{y}) - (1-y) \log(1-\hat{y})$$

- The cross-entropy cost function is non-negative,  $\ell_{BCE} > 0$
- $\ell_{BCE} \approx 0$  when the prediction and targets are equal (i.e. y = 0 and  $\hat{y} = 0$  or when y = 1 and  $\hat{y} = 1$ )
- With Sigmoidal final layer,  $\frac{\partial \ell_{BCE}}{\partial W_i^{(2)}}$  is proportional to just the error in the output  $(\hat{y} y)$  and therefore, the larger the error, the faster the network will learn!
- Note that the BCE is the negative log likelihood of the Bernoulli Distribution

- The cross-entropy can be thought of as a measure of surprise.
- Given some input x<sub>i</sub>, we can think of ŷ<sub>i</sub> as the estimated probability that x<sub>i</sub> belongs to class 1, and 1 − ŷ<sub>i</sub> is the estimated probability that it belongs to class 0.
- Note the extreme case of infinite cross-entropy, if your model believes that a class has 0 probability of occurrence, and yet the class appears in the data, the 'surprise' of your model will be infinitely great.

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### Binary Cross-Entropy for multiple labels

In the case of multi-label classification with a network with multiple sigmoidal outputs you just sum the BCE over the outputs:

$$\ell_{BCE} = -\sum_{k=1}^{K} [y_k \log(\hat{y}_k) + (1 - y_k) \log(1 - \hat{y}_k)]$$

where K is the number of classes of the classification problem,  $\hat{y} \in \mathbb{R}^{K}$ .

#### Numerical Stability: The Log-Sum-Exp trick

$$\ell_{BCE}(\hat{y}, y) = -y \log(\hat{y}) - (1-y) \log(1-\hat{y})$$

- Consider what might happen early in training when the model might confidently predict a positive example as negative
  - $\hat{y} = \sigma(z) \approx 0 \implies z \ll 0$
  - if  $\hat{y}$  is small enough, it will become 0 due to limited precision of floating-point representations
  - but then  $log(\hat{y}) = -inf$ , and everything will break!
- To tackle this problem implementations usually combine the sigmoid computation and BCE into a single loss function that you would apply to a network with linear outputs (e.g. BCEWithLogitsLoss).
- Internally, a trick called 'log-sum-exp' is used to *shift* the centre of an exponential sum so that only numerical underflow can potentially happen, rather than overflow<sup>2</sup>.
  - Ultimately this means you'll always get a numerically reasonable result (and will avoid NaNs and Infs originating from this point).

<sup>2</sup>https://www.xarg.org/2016/06/the-log-sum-exp-trick-in-machine-learning/ Jonathon Hare COMP6258

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#### Multiclass classification with Softmax Outputs

- Softmax can be thought of making the *K* outputs of the network mimic a probability distribution.
- The target label y could also be represented as a distribution with a single 1 and zeros everywhere else.
  - e.g. they are "one-hot encoded".
- In such a case, the obvious loss function is the negative log likelihood of the Categorical distribution (aka Multinoulli, Generalised Bernoulli, Multinomial with one sample)<sup>3</sup>: ℓ<sub>NLL</sub> = -∑<sup>K</sup><sub>k=1</sub> y<sub>k</sub> log ŷ<sub>k</sub>
  - Note that in practice as  $y_k$  is zero for all but one class you don't actually do this summation, and if y is an integer class index you can write  $\ell_{NLL} = -\log \hat{y}_y$ .
- Analogously to what we saw for BCE, Log-Sum-Exp can be used for better numerical stability.
  - PyTorch combines LogSoftmax with NLL in one loss and calls this "Categorical Cross-Entropy" (so you would use this with a *linear* output layer)
- <sup>3</sup>Note: Keras calls this function 'Categorical Cross-Entropy'; you would need to have a Softmax output layer to use this

- Define total loss as  $\mathcal{L} = \sum_{(x,y)\in D} \ell(g(x,\theta), y)$  for some loss function  $\ell$ , dataset **D** and model g with learnable parameters  $\theta$ .
- Define how many passes over the data to make (each one known as an Epoch)
- Define a learning rate  $\eta$

Gradient Descent updates the parameters  $\theta$  by moving them in the direction of the negative gradient with respect to the **total loss**  $\mathcal{L}$  by the learning rate  $\eta$  multiplied by the gradient:

for each Epoch: $oldsymbol{ heta} \leftarrow oldsymbol{ heta} - \eta 
abla_{oldsymbol{ heta}} \mathcal{L}$ 

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#### Reminder: Stochastic Gradient Descent

- Define loss function  $\ell$ , dataset **D** and model g with learnable parameters  $\theta$ .
- Define how many passes over the data to make (each one known as an Epoch)
- Define a learning rate  $\eta$

Stochastic Gradient Descent updates the parameters  $\theta$  by moving them in the direction of the negative gradient with respect to the loss of a **single item**  $\ell$  by the learning rate  $\eta$  multiplied by the gradient:

```
for each Epoch:
for each (m{x},m{y})\inm{D}:
m{	heta}\leftarrowm{	heta}-\eta
abla_{m{	heta}}\ell
```

### A Quick Introduction to Tensors

Broadly speaking a tensor is defined as a linear mapping between sets of algebraic objects<sup>4</sup>.

A tensor T can be thought of as a generalization of scalars, vectors and matrices to a single algebraic object.

We can just think of this as a multidimensional array<sup>5</sup>.

- A 0*D* tensor is a scalar
- A 1D tensor is a vector
- A 2D tensor is a matrix
- A 3D tensor can be thought of as a vector of identically sized matrices
- A 4D tensor can be thought of as a matrix of identically sized matrices or a sequence of 3D tensors

• . . .

<sup>4</sup>This statement is always entirely true

<sup>5</sup>This statement will upset mathematicians and physicists because its not always true for them (but it is for us!).

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#### Aside: Tensor Decompositions

- Just in the same way a matrix can be decomposed into a product of matrices (EVD, SVD, QR, LU, Cholesky, ...), there are tensor decompositions:
  - PARAFAC / Canonical polyadic / HO-SVD / Tucker
  - These have found their way into some deep learning models as a form of structural regularisation or weight reduction

- PyTorch lets you do all the standard matrix operations on 2D tensors
  - including important things you might not yet have seen like the hadamard product of two  $N \times M$  matrices:  $A \odot B$ )
- You can do element-wise add/divide/subtract/multiply to ND-tensors
  - and even apply scalar functions element-wise (log, sin, exp, ...)
- you can slice, reshape, and even index a single element (generally don't do that!)
- PyTorch often lets you broadcast operations (just like in numpy)
  - if a PyTorch operation supports broadcast, then its Tensor arguments can be automatically expanded to be of equal sizes (without making copies of the data).<sup>6</sup>

<sup>6</sup>Important - read and understand this after the lab next week:

https://pytorch.org/docs/stable/notes/broadcasting.html
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#### Tensors, batches and vectorisation

- The reality of training a model is that we neither use gradient descent or stochastic gradient descent; we do something in-between called mini-batch SGD.
- This works on batches of data (e.g. small subsets of the training set)
- These batches are assembled into a tensor
- Broadcasting is used to apply operations/functions to all the samples in the batch tensor *in parallel* to compute a loss vector
- the loss vector is summed/averaged using a vectorised method (e.g. .sum())

## Tensor implementation

It's important to understand something about how tensors are implemented in software and particularly how memory copies can be avoided...

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We are siamese...

An important and clever trick:

#### PyTorch Tensor 101: https://colab.research.google.com/gist/jonhare/ d98813b2224dddbb234d2031510878e1/notebook.ipynb

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