Training tips/recent advances

Why now?

- 1. Advances in computational hardware (GPU, CPU, TPU)
- 2. Some algorithmic developments, help in training
- 3. Advent of big data: many more samples now than ever before

A brief history of Neural networks

- 1. McCulloch and Pitts (1943): all-or-nothing model of neurons.
- 2. Rosenblatt (1957): perceptron mechanism for learning based on Hebb (1949).
- 3. Limitations of perceptrons: Minksy and Papert (1969)
- 4. 1973: Lighthill report led to first AI winter.
- 5. Backpropagation (Werbos; 1974). Popularised by Hinton et al in 1980s.
- 6. Limitations in hardware led to 2nd AI winter late 1990s.
- 7. 2012: resurgence due to hardware and some new "tricks".
- 8. Computational biology now a big user (Angermueller et al 2016), e.g. protein folding (ALphaFold) and biomedicine (Ching et al 2018).

See Schmidhuber (2015) for further history.

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Practical matters

- Like computational biology, 80% of the work is mundane but critical (data collection, cleaning, hyper-parameter selction).
- The mathematics of the systems are clearly defined; engineering them to work is a real challenge.
- Good news: many frameworks. We will use Keras (with Tensor Flow backend).
- GPU vs CPU
- Desktop vs Cloud

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Classification: modifications

Three tweaks:

- 1. **1 HOT encoding** of training signal
- 2. **Softmax** function on output layer:

$$y_i = \frac{\exp(z_i)}{\sum_{j=1}^N \exp(z_j)}$$

Converts output layer into probability distribution.

Effectively consider another layer, with "lateral interactions" to compute sum term.

Related to logistic regression from classical statistics.

3. **Cross-entropy** loss function:

$$E = -\sum_{i=1}^{N} t_i \log(y_i)$$

for comparing two probability distributions. Minimal () when t=y. See Stone book, eq 4.45.

Some more definitions

- 1. **flattening images** 2D image structure collapsed into 1D vector
- 2. **iteration** presentation of one input vector
- 3. mini batch grouping together input vectors and then learning
- 4. **epoch** one presentation of all input vectors (in training set)
- 5. training set vs test set Used for learning vs testing.

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Additional terms

Many additions have been suggested to back-propagation to help prevent e.g. overfitting, local minima:

• Weight decay/ L2 regularization:

$$E = \frac{1}{2}(t-y)^2 + \beta \sum_{i} w_i^2$$

• Momentum: add history to weight changes:

$$\Delta w_{ij}(t+1) = -\eta \frac{\partial E}{\partial w_{ij}} + \alpha \Delta w_{ij}(t)$$

where 0 $< \alpha <$ 1.

Several other methods available (conjugate gradient, rmsprop, adam) that improve convergence.

See Ruder (2017) for overview.

Merit function

Error minimisation can also be a maximisation of a "merit function". e.g. Stone and Bray (1995) form of "self-supervised activity".

$$F = \log \frac{V}{U}$$

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Recent advances

Awareness of gradient descent

If early methods (family trees, nettalk) worked so well, why did research stop in the early 1990s?

And what made it work again?

- Not strictly a new innovation, but watching the 'vanishing' gradient, or 'exploding' gradient helped.
- Avoiding saturation of hidden units. RELU helped here.
- Weight initialisation schemes suggested to help avoid saturation (Glorot and Bengio, 2010). Unsupervised pre-learning of weights.

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GPUs

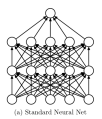
Increased volumes of training data

Compute resource is improving all the time. GPUs especially useful, given parallel nature of computation.

- We now live in an age of data... large data sets are everywhere. Large networks needs lots of data.
- Labelled data were sparse in the 1990s, but now relatively abundant. ImageNet competition being a prime example (see later).
- Remember to clean up the input data, and try to normalise it somehow, e.g. so inputs are all scaled to same distribution (zero mean, unit variance).

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Dropout (Srivastava et al 2014, fig 1)



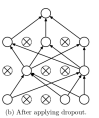


Figure 1: Dropout Neural Net Model. Left: A standard neural net with 2 hidden layers. Right: An example of a thinned net produced by applying dropout to the network on the left. Crossed units have been dropped.

- **Dropout** introduced as a way to reduce overfitting. Bank-teller analogy.
- During training, on each iteration silence some fraction of units.
- Implemented during training:
 - 1. once the activation of a layer calculated, set output of half of neurons to zero.
 - 2. Double the activation of rest of the neurons.

Hyper parameters

How do you decide on the hyper-parameters of the model?

- 1. Network architecture: how many hidden units? how many hidden layers?
- 2. Learning rates (and other parameters)
- 3. Error functions
- 4. Feature selection
- 5. Data set size

One approach is to overfit then regularize. Sophisticated methods exist, but often more pragmatic approaches taken.

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Autograd tools

- 1. We would like to find $\frac{\partial E}{\partial w_i}$ for every weight in the network.
- 2. Autograd tools can now do the hard work for us. They do not calculate the symbolic derivative. (See separate video and code example.) See also the Julia implementation:
 - https://www.youtube.com/watch?v=vAp6nUMrKYg.
- 3. R work in this area is developing, see e.g. torch package on CRAN.
- 4. https://en.wikipedia.org/wiki/Automatic_differentiation