

Foundations of machine learning  
Deep Neural Nets

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# Outline

- What are neural nets?
- Network design:
  - Activation functions,
  - network architecture,
  - output layers.
- Calculating gradients for optimization:
  - Backpropagation,
  - stochastic gradient descent.
- Regularization using early stopping.

## Takeaways for this part of class

- Deep learning is regression with complicated functional forms.
- Design considerations in feedforward networks include depth, width, and the connections between layers.
- Optimization is difficult in deep learning because of
  1. lots of data
  2. and even more parameters
  3. in a highly non-linear model.
- $\Rightarrow$  Specially developed optimization methods.
- Cross-validation for penalization is computationally costly, as well.
- A popular alternative is sample-splitting and early stopping.

# Deep Neural Nets

## Setup

- Deep learning is (regularized) maximum likelihood, for regressions with complicated functional forms.
- We want, for instance, to find  $\theta$  to minimize

$$E \left[ (Y - f(X, \theta))^2 \right]$$

for continuous outcomes  $Y$ , or to maximize

$$E \left[ \sum_y \mathbf{1}(Y = y) \cdot \log(f^y(X, \theta)) \right]$$

for discrete outcomes  $Y$ .

# What's deep about that?

## Feedforward nets

- Functions  $f$  used for deep (feedforward) nets can be written as

$$f(\mathbf{x}, \theta) = f^k(f^{k-1}(\dots f^1(\mathbf{x}, \theta^1), \theta^2), \dots, \theta^k).$$

- Biological analogy:
  - Each value of a component of  $f^j$  corresponds to the “activation” of a “neuron.”
  - Each  $f^j$  corresponds to a layer of the net.  
Many layers  $\Rightarrow$  “deep” neural net.
  - The layer-structure and the parameters  $\theta$  determine how these neurons are connected.
- Inspired by biology, but practice moved away from biological models.
- Best to think of as a class of nonlinear functions for regression.

## So what's new?

- Very non-linear functional forms  $f$ . Crucial when
  - mapping pixel colors into an answer to “Is this a cat?”
  - or when mapping English sentences to Mandarin sentences.
  - Probably less relevant when running Mincer-regressions.
- Often more parameters than observations.
  - Not identified in the usual sense.  
But we care about predictions, not parameters.
  - Overparametrization helps optimization:  
Less likely to get stuck in local minima.
- Lots of computational challenges.
  1. Calculating gradients:  
Backpropagation, stochastic gradient descent.
  2. Searching for optima.
  3. Tuning: Penalization, early stopping.

Setup

Network design

Back propagation

Stochastic gradient descent

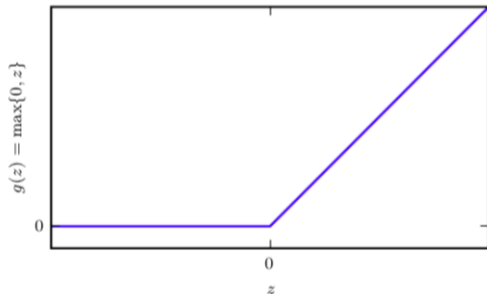
Early stopping

References

# Network design

## Activation functions

- Basic unit of a net:  
A neuron  $i$  in layer  $j$ .
- Receives input vector  $\mathbf{x}_i^j$   
(output of other neurons).
- Produces output  $\mathbf{g}(\mathbf{x}_i^j \theta_i^j + \eta_i^j)$ .
- Activation function  $\mathbf{g}(\cdot)$ :
  - Older nets: Sigmoid function  
(biologically inspired).
  - Modern nets: "Rectified linear  
units:"  $\mathbf{g}(z) = \max(\mathbf{0}, z)$ .  
More convenient for getting  
gradients.

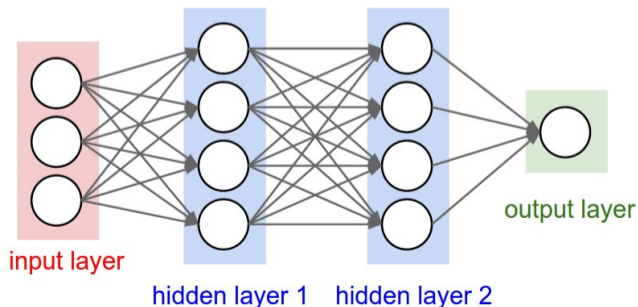




# Network design

## Architecture

- These neurons are connected, usually structured by layers.  
Number of layers: Depth. Number of neurons in a layer: Width.
- Input layer: Regressors.
- Output layer: Outcome variables.
- A typical example:



# Network design

## Architecture

- Suppose each layer is fully connected to the next, and we are using RELU activation functions.
- Then we can write in matrix notation (using componentwise  $\max$ ):

$$\mathbf{x}^j = f^j(\mathbf{x}^{j-1}, \theta^j) = \max(0, \mathbf{x}^{j-1} \cdot \theta^j + \eta_j)$$

- Matrix  $\theta^j$ :
  - Number of rows: Width of layer  $j - 1$ .
  - Number of columns: Width of layer  $j$ .
- Vector  $\mathbf{x}^j$ :
  - Number of entries: Width of layer  $j$ .
- Vector  $\eta_j$ :
  - Number of entries: Width of layer  $j$ .
  - Intercepts. Confusingly called “bias” in machine learning.

# Network design

## Output layer

- Last layer is special: Maps into predictions.
- Leading cases:
  1. Linear predictions for continuous outcome variables,

$$f^k(x^{k-1}, \theta^k) = x^{k-1} \cdot \theta^k.$$

2. Multinomial logit (aka “softmax”) predictions for discrete variables,

$$f^{ky_j}(x^{k-1}, \theta^k) = \frac{\exp(x_j^{k-1} \cdot \theta_j^k)}{\sum_{j'} \exp(x_{j'}^{k-1} \cdot \theta_{j'}^k)}$$

- Network with only output layer: Just run OLS / multinomial logit.

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# The gradient of the likelihood

## Practice problem

Consider a fully connected feedforward net with rectified linear unit activation functions.

1. Write out the derivative of its likelihood, for  $n$  observations, with respect to any parameter.
2. Are there terms that show up repeatedly, for different parameters?
3. In what sequence would you calculate the derivatives, in order to minimize repeat calculations?
4. Could you parallelize the calculation of derivatives?

# Backpropagation

## The chain rule

- In order to maximize the (penalized) likelihood, we need its gradient.
- Recall  $f(\mathbf{x}, \theta) = f^k(f^{k-1}(\dots f^1(\mathbf{x}, \theta^1), \theta^2), \dots, \theta^k)$ .
- By the **chain rule**:

$$\frac{\partial f(\mathbf{x}, \theta)}{\partial \theta_i^j} = \left( \prod_{j'=j+1}^k \frac{\partial f^{j'}(\mathbf{x}^{j'}, \theta^{j'})}{\partial \mathbf{x}^{j'-1}} \right) \cdot \frac{\partial f^j(\mathbf{x}^{j-1}, \theta^j)}{\partial \theta_i^j}.$$

- A lot of the same terms show up in derivatives w.r.t different  $\theta_i^j$ :
  - $\mathbf{x}^{j'}$  (values of layer  $j'$ ),
  - $\frac{\partial f^{j'}(\mathbf{x}^{j'}, \theta^{j'})}{\partial \mathbf{x}^{j'-1}}$  (intermediate layer derivatives w.r.t.  $\mathbf{x}^{j'-1}$ ).

# Backpropagation

- Denote  $\mathbf{z}^j = \mathbf{x}^{j-1}\theta^j + \eta^j$ . Recall  $\mathbf{x}^j = \max(\mathbf{0}, \mathbf{z}^j)$ .
- Note  $\partial \mathbf{x}^j / \partial \mathbf{z}^j = \mathbf{1}(\mathbf{z}^j \geq \mathbf{0})$  (componentwise), and  $\partial \mathbf{z}^j / \partial \theta^j = \mathbf{x}^{j-1}$
- First, **forward propagation**:  
Calculate all the  $\mathbf{z}^j$  and  $\mathbf{x}^j$ , starting at  $j = 1$ .
- Then **backpropagation**:  
Iterate backward, starting at  $j = k$ :
  1. Calculate and store

$$\frac{\partial f(\mathbf{x}, \theta)}{\partial \mathbf{x}^{j-1}} = \frac{\partial f(\mathbf{x}, \theta)}{\partial \mathbf{x}^j} \cdot \mathbf{1}(\mathbf{z}^j \geq \mathbf{0}) \cdot \theta^j.$$

2. Calculate

$$\frac{\partial f(\mathbf{x}, \theta)}{\partial \theta^j} = \frac{\partial f(\mathbf{x}, \theta)}{\partial \mathbf{x}^j} \cdot \mathbf{1}(\mathbf{z}^j \geq \mathbf{0}) \cdot \mathbf{x}^{j-1}.$$

# Backpropagation

## Advantages

- Backpropagation improves efficiency by **storing** intermediate derivatives, **rather than recomputing** them.
- Number of computations grows only linearly in number of parameters.
- The algorithm is easily generalized to more complicated network architectures and activation functions.
- Parallelizable across observations in the data (one gradient for each observation!).



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## Stochastic gradient descent

- Gradient descent updates parameter estimates in the direction of steepest descent:

$$g_t = \frac{1}{n} \sum_{i=1}^n \nabla_{\theta} m(X_i, Y_i, \theta)$$

$$\theta_{t+1} = \theta_t - \varepsilon_t g_t.$$

- Stochastic gradient descent (SGD) does the same, but instead uses just a random subsample  $B_t = \{i_1^t, \dots, i_b^t\}$  (changing across  $t$ ) of the data:

$$\hat{g}_t = \frac{1}{b} \sum_{i \in B_t} \nabla_{\theta} m(X_i, Y_i, \theta)$$

$$\theta_{t+1} = \theta_t - \varepsilon_t \hat{g}_t.$$

# Stochastic gradient descent

- We can do this because the full gradient is a sum of gradients for each observation.
- Typically, the batches  $B_t$  cycle through the full dataset.
- If the learning rate  $\varepsilon_t$  is chosen well, some convergence guarantees exist.
- The built-in randomness might help avoiding local minima.
- Extension: SGD with **momentum**,

$$v_t = \alpha v_{t-1} - \varepsilon_t \hat{g}_t,$$
$$\theta_{t+1} = \theta_t + v_t.$$

- Initialization matters. Often start from previously trained networks.

## Why SGD makes sense

- The key observation that motivates SGD is that in an (i.i.d.) sampling context, further observations become more and more redundant.
- Formally, the standard error of a gradient estimate based on  $b$  observations is of order  $1/\sqrt{b}$ .
- But the computation time is of order  $b$ .
- Think of a very large data-set. Then it would take forever to just calculate one gradient, and do one updating step.
- During the same time, SGD might have made many steps and come considerably closer to the truth.
- Bottou et al. (2018) formalize these arguments.

## Excursion: Data vs. computation as binding constraint

- This is a good point to clarify some distinctions between the approaches of statisticians and computer scientists.
- Consider a regularized m-estimation problem.
- Suppose you are constrained by
  1. a finite data set,
  2. a finite computational budget.
- Then the difference between any estimate and the estimand has three components:
  1. Sampling error (variance),
  2. approximation error (bias),
  3. optimization error (failing to find the global optimum of your regularized objective function).

# Statistics and computer science

- Statistical decision theory focuses on the trade-off between variance and bias.
- This makes sense if data-sets are small relative to computational capacity, so that optimization error can be neglected.
- Theory in computer science often focuses on optimization error.
- This makes sense if data-sets are large relative to computational capacity, so that sampling error can be neglected.
- Which results are relevant depends on context!
- More generally, I believe there is space for interesting theory that explicitly trades off all three components of error.

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# Regularization for neural nets

- To get good predictive performance, neural nets need to be regularized.
- As before, this can be done using **penalties** such as  $\lambda \|\theta\|_2^2$  (“Ridge”) or  $\lambda \|\theta\|_1$  (“Lasso”).
- Problem: Tuning using cross-validation is often computationally too costly for deep nets.
- An alternative regularization method is **early stopping**:
  - Split the data into a training and a validation sample.
  - Run gradient-based optimization method on the training sample.
  - At each iteration, calculate prediction loss in the validation sample.
  - Stop optimization algorithm when this prediction loss starts increasing.



## References

- *Goodfellow, I., Bengio, Y., and Courville, A. (2016). Deep learning. MIT Press, chapters 6-8.*
- *Bottou, L., Curtis, F. E., and Nocedal, J. (2018). Optimization methods for large-scale machine learning. SIAM Review, 60(2):223–311*