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

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

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

for continuous outcomes Y, or to maximize

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

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 ⇒ "deep" neural net.
 - The layer-structure and the parameters θ 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

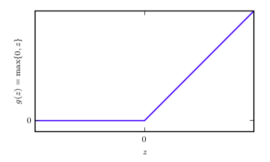
Back propagation

Stochastic gradient descent

Early stopping

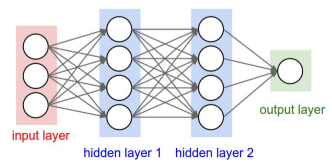
Activation functions

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



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:



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):

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

- Matrix θ^{j} :
 - Number of rows: Width of layer j 1.
 - Number of columns: Width of layer j.
- Vector x^j:
 - Number of entries: Width of layer j.
- Vector η_i :
 - Number of entries: Width of layer j.
 - Intercepts. Confusingly called "bias" in machine learning.

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.

Setup

Network design

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Stochastic gradient descent

Early stopping

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:

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

- A lot of the same terms show up in derivatives w.r.t different θ_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 $\partial x^j / \partial z^j = \mathbf{1}(z^j \ge \mathbf{0})$ (componentwise), and $\partial z^j / \partial \theta^j = x^{j-1}$

• First, forward propagation: Calculate all the z^{j} and x^{j} , starting at j = 1.

Then **backpropagation**: Iterate backward, starting at *j* = *k*:

1. Calculate and store

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

2. Calculate

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

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!).

Setup

Network design

Back propagation

Stochastic gradient descent

Early stopping

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:

$$\hat{g}_t = rac{1}{b} \sum_{i \in B_t}
abla_{ heta} m(X_i, Y_i, heta)$$
 $heta_{t+1} = heta_t - arepsilon_t \hat{g}_t.$

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 ε_t is chosen well, some convergence guarantees exist.
- The built-in randomness might help avoiding local minima.
- Extension: SGD with momentum,

$$\mathbf{v}_t = lpha \mathbf{v}_{t-1} - arepsilon_t \hat{\mathbf{g}}_t, \ \mathbf{\theta}_{t+1} = \mathbf{\theta}_t + \mathbf{v}_t.$$

• 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.

- 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 largescale machine learning. SIAM Review, 60(2):223–311