Regularization

1

Reading

- Deep Learning: chapter 7
	- [https://www.deeplearningbook.org/contents/regularization](https://www.deeplearningbook.org/contents/regularization.html) [.html](https://www.deeplearningbook.org/contents/regularization.html)
- Regularization overview, with a deep learning bias

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- Occam's Razor
	- Try to use simplest model family possible
- Neural nets can easily overfit any dataset we have come up with
	- Regularization adds constraints to keep models wellbehaved
- A bit of a funky concept
	- We want to minimize the loss, but we also want to minimize it the right way!
	- Comes to indicate that our losses could be improved

- One of the most popular regularizations
- Suppose original loss is $J(\theta; X, y)$
- Come up with an extra term $\Omega(\boldsymbol{\theta})$ that penalizes the parameters
- Final loss becomes

 $\tilde{J} = J(\boldsymbol{\theta}; X, y) + \alpha \Omega(\boldsymbol{\theta})$

- where α is a (small) hyper-parameter

• Often reduces variance at the expense of some bias

L₂ Penalty

• Most standard penalty

$$
\Omega(\boldsymbol{\theta}) = \boldsymbol{w}^T \boldsymbol{w}
$$

 $-$ Recall that $\boldsymbol{\theta} = [\boldsymbol{w}, \boldsymbol{b}]$

- Usually applied only to weights, not to biases
	- Regularizing biases leads to underfitting without major variance benefits
- Also known as weight decay
	- Recall that weights are updated as follows $w' = w - \epsilon \nabla \tilde{J}(\boldsymbol{\theta}; X, y)$
	- With L_2 penalty, the update is

$$
w' = w - \epsilon(2\alpha w + \nabla J(\boldsymbol{\theta}; \boldsymbol{X}, \boldsymbol{y}))
$$

= (1 - 2\alpha \epsilon)w - \epsilon \nabla J(\boldsymbol{\theta}; \boldsymbol{X}, \boldsymbol{y})

Penalty, cont'd

- If true loss is quadratic, L_2 penalty penalizes learning in directions where the loss isn't affected
	- Prevents learning spurious functionality due to overparameterization (proof in book)
- Same idea in general keep weights small unless necessary – Simplifies models and improves robustness
- In linear regression, makes fitting more robust to variance $-2Xy + 2XX^T w + 2\alpha w = 0$
- Then $\boldsymbol{w}^* = \left(\boldsymbol{X}\boldsymbol{X}^T + \alpha \boldsymbol{I}\right)^{-1}$ χ
- I have also used L_2 penalty in my research

L₁ Penalty

• A slightly less standard penalty

$$
\Omega(\boldsymbol{\theta}) = ||\boldsymbol{w}||_1 = \sum_i |w_i|
$$

- Note that the derivative of $\Omega(\boldsymbol{\theta}) = sign(\boldsymbol{w})$
- So the weight update is now

$$
w' = w - \epsilon\big(\alpha sign(w) + \nabla J(\theta; X, y)\big)
$$

- i.e., a constant factor along the direction of the 1-norm
- Might lead to sparser weight matrices (more 0s)
- Hard to derive nice mathematical formulae
- Overall regularization effect is similar to L_2

• Instead of penalizing bigger weights, we can impose an explicit constraint

$$
\boldsymbol{\theta}^* = argmin_{\boldsymbol{\theta}} J(\boldsymbol{\theta}; \boldsymbol{X}, \boldsymbol{y})
$$

subject to $\Omega(\boldsymbol{\theta}) < k$

- Explicit constraints may stabilize the learning process in certain cases (since the loss is simplified)
	- However, it may hurt in others
	- Constrained non-convex optimization is a hard problem
- Can also reformulate the problem as $\theta^* = argmin_{\theta} max_{\theta}$ α $J(\boldsymbol{\theta}; X, y) + \alpha(\Omega(\boldsymbol{\theta}) - k)$
	- Same idea, optimization algorithm slightly different
	- Dual formulation of the constrained problem above

- Using fake data for training is not always a good idea! – But sometimes OK…
- In image classification, we can usually generate "new" data from a given dataset
	- Rotate, translate, add white noise to images
	- Useful because it discourages learning spurious relationships (similar to regularization)
	- You can overdo it, though. Thoughts?

- Can also apply noise to the weights
	- E.g., in Bayesian neural networks every weight is drawn from a Gaussian with learned parameters
	- Pushes weights to region where model is less sensitive to perturbations
- Can also do it at the output layer
	- "Soft" labels aka label smoothing
		- E.g., one-hot labels are not (0,1) but maybe (0.1, 0.9)
	- Discourages the NN from learning very big weights in trying to approximate the hard 0/1 outputs

- A lot of deep learning has to do with learning representations of the training data that are separated in some embedding space
- What if we learn the embedding separately from the classifier?
	- i.e., learn a generative model of the data first
	- An active research area, improves robustness a great deal
- Many types of generative models, such as generative adversarial networks (GANs), variational autoencoders (VAEs) and others

Early Stopping

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- Often, training loss keeps decreasing while validation loss starts increasing
	- A sign of overfitting

- Can stop training as soon as this happens
	- (or save trained weights frequently and go back to that checkpoint)
	- Technique called **early stopping**

Early Stopping, cont'd

- Simple and effective
- Essentially another hyper-parameter
- Periodically storing weights is not a major overhead
- However, if early stopping is necessary, then you're violating Occam's Razor
	- If your model overfits drastically, you should consider using a simpler model
- Related to L_2 penalty

Ensemble Methods

- A very old and effective idea in ML
- Train multiple models on the same task and average their outputs
- Very effective if done well (i.e., models make independent errors)
- Suppose you have k models, each makes error ϵ_i (where ϵ_i is a zero-mean random variable)
	- Ensemble method error is $\frac{1}{\mu}$ $\frac{1}{k} \sum_i \epsilon_i$
	- Suppose they have same variance $\mathbb{E}\big[\epsilon_i^2\big] = \nu$, and covariances are $\mathbb{E}[\epsilon_i \epsilon_j] = c$

Ensemble Methods, cont'd

- Suppose $\mathbb{E}\big[\epsilon_i^2\big] = v$ and $\mathbb{E}\big[\epsilon_i \epsilon_j\big] = c$
- Expected squared error is

$$
\mathbb{E}\left[\left(\frac{1}{k}\sum_{i}\epsilon_{i}\right)^{2}\right]=\frac{1}{k^{2}}\mathbb{E}\left[\sum_{i}\left(\epsilon_{i}^{2}+\sum_{i\neq j}\epsilon_{i}\epsilon_{j}\right)\right]
$$

$$
=\frac{1}{k}v+\frac{k-1}{k}c
$$

- If perfect correlation, $v = c$, average doesn't help
- If no correlation, $c = 0$, squared error inversely proportional to number of models

Ensemble Methods, cont'd

- Suppose you resample original dataset and train a different model each time
	- Models learn different important features
	- More robust overall since spurious features averaged out
	- This is the idea of boosting

Dropout

- Generated a lot of attention a few years ago
- A computationally cheap way to approximate ensemble of methods
- The "ensemble" is the set of all subnetworks of a given NN
	- To eliminate a neuron, just multiply its output by 0
- Slightly different from classic ensembles since data is the same

Dropout, cont'd

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- During training, select a random bit mask μ for each iteration of gradient descent
	- Enumerating all subnetworks is intractable
	- E.g., keep input neurons with a probability of 0.8 and hidden neurons with probability of 0.5
	- $-$ Once you have determined μ , train as before using backprop

- Suppose we have trained our model with dropout
- How do we predict the label given a new example?
- Ideally, we enumerate all subgraphs and compute the mean of all subnetwork outputs
	- What's the challenge with this?
		- Exponentially many subnetworks
- One option is to sample a number of masks and average over those (a reasonably good estimate of the true average) – Not deterministic
- Better idea: output expected value of each neuron (how?)
	- multiply all weights by the keep probability, $(1 p)$
	- binary variable with parameter $1-p$

- One idea that works very well in practice is to multiply all weights by the dropout probability, p
	- Most common choice
- Another idea is to sample masks and compute geometric mean

$$
\mathbb{P}_{ensemble}(y \mid x) = \sqrt[2^d]{\prod_{\mu} \mathbb{P}_{dropout}(y \mid x, \mu)}
$$

- Normalize over classes (doesn't sum up to 1 otherwise)
- For some architectures, this is the same as multiplying the weights by p
	- E.g., networks with one layer and a softmax output
	- See proof in book

- Neural networks can perfectly overfit any existing dataset – Even if you randomly shuffle the labels
- In some sense, not clear why NNs perform as well as they do
- Always a good idea to use as small a model as possible
	- If your training accuracy is significantly higher than test accuracy, then likely you need to regularize or reduce your model
- Deep learning is a powerful tool but it requires a strong understanding of statistics

Zhang, Chiyuan, Samy Bengio, Moritz Hardt, Benjamin Recht, and Oriol Vinyals. "Understanding deep learning (still) requires rethinking generalization." *Communications of the ACM* 64, no. 3 (2021): 107-115.

Summary

- Many, many ways to regularize
- Usually trial and error is the best approach
- If you set up everything well (right model, right features, etc.), you may not even need much regularization
- An L_2 regularization will typically get you a long way