Deep Learning Teaching Deep Learners to Generalize

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Summary

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What About Supervised Pre-training?

What is Model Generalization?

- ▶ In a machine learning problem, we try to generalize the known dependent variable on seen instances to unseen instances.
 - Unseen ⇒ The model did not see it during training.
 - Given training images with seen labels, try to label an unseen image.
 - Given training emails labeled as spam or nonspam, try to label an unseen email.
- The classification accuracy on instances used to train a model is usually higher than on unseen instances.
 - We only care about the accuracy on unseen data.

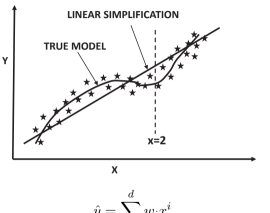
Memorization vs Generalization

- Why is the accuracy on seen data higher?
 - Trained model remembers some of the irrelevant nuances.
 - "Free Money" present on spam emails.
- When is the gap between seen and unseen accuracy likely to be high?
 - When the amount of data is limited. Suppose the model is trained on only two email messages.
 - When the model is complex (which has higher capacity to remember nuances).
 - The combination of the two is a deadly cocktail.
- ▶ A high accuracy gap between the predictions on seen and unseen data is referred to as *overfitting*.

Overfitting

Depends both on the complexity of the model and on the amount of data available.

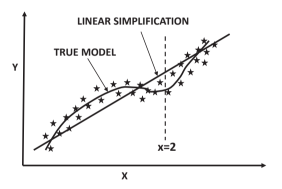
Example: Predict y from x



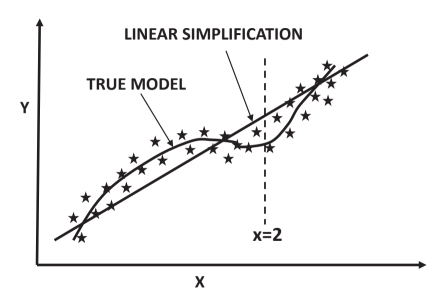
$$\hat{y} = \sum_{i=0}^{d} w_i x^i$$

$$\mathcal{L} = (y - \hat{y})^2$$
(1)

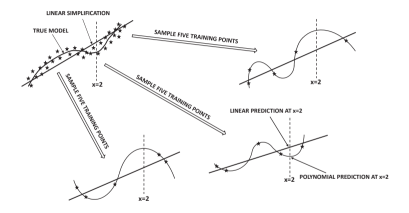
Example: Predict y from x



- ▶ First impression: Polynomial model such as $\hat{y} = w_0 + w_1 x + w_2 x^2 + w_3 x^3 + w_4 x^4$ is "better" than linear model $\hat{y} = w_0 + w_1 x$.
 - Bias-variance trade-off says: "Not necessarily! How much data do you have?"



Different Training Data Sets with Five Points



lacktriangle Zero error on training data but wildly varying predictions of x=2

Observations

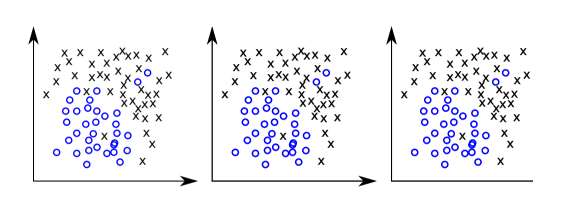
- Linear model does not change much with the training data, whereas the polynomial model changes drastically.
- ▶ The higher-order model is more complex than the linear model and has less bias.
 - But it has more parameters.
 - For a small training data set, the learned parameters will be more sensitive to the nuances of that data set.
 - Different training data sets will provide different predictions for y at a particular x.
 - This variation is referred to as model variance.
- Neural networks are inherently low-bias and high-variance learners ⇒ Need ways of handling complexity.

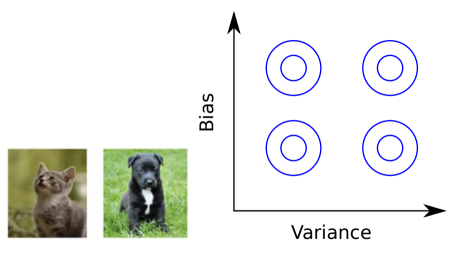
Noise Component

- Unlike bias and variance, noise is a property of the data rather than the model.
- Noise refers to unexplained variations ϵ_i of data from true model $y_i = f(x_i) + \epsilon_i$.
- ► Real-world examples:
 - Human mislabeling of test instance ⇒ Ideal model will never predict it accurately.
 - Error during collection of temperature due to sensor malfunctioning.
- Cannot do anything about it even if seeded with knowledge about true model.

Tell-tale signs of overfitting

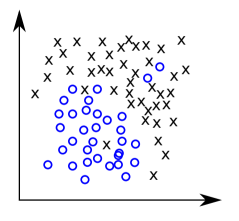
- Same test instance obtains very different predictions → Indication that the training process is memorizing the nuances of the specific training data set, rather than learning patterns that generalize to unseen test instances.
- 2. The gap between the error of predicting training instances and unseen test instances is rather large.





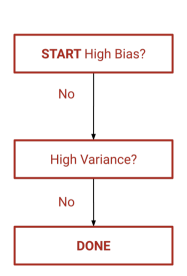
Assume human error $\approx 0\%$.

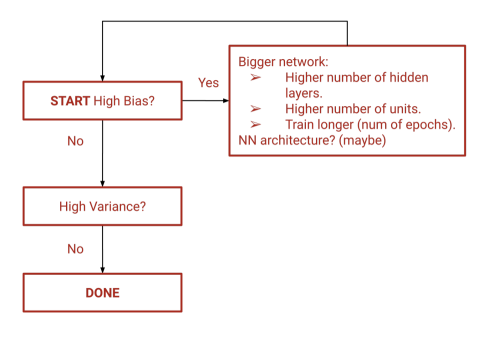
The worst of both worlds (high bias and high variance)

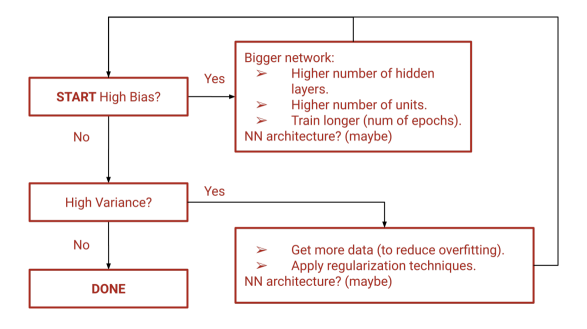


Key Takeaway of Bias-Variance Trade-Off

- ▶ A model with greater complexity might be *theoretically* more accurate (i.e., low bias).
 - But you have less control on what it might predict on a tiny training data set.
 - Different training data sets will result in widely varying predictions of same test instance.
 - Some of these must be wrong \Rightarrow Contribution of model variance.
- A more accurate model for infinite data is not a more accurate model for finite data.
 - Do not use a sledgehammer to swat a fly!







Assume

- ightharpoonup x Independent variable.
- \triangleright y Dependent variable.
- ightharpoonup f(x) Describes the true underlying dependence of y on x.
- $ightharpoonup y = f(x) + \epsilon$ The result of f(x) and random noise.
- $ightharpoonup \epsilon$ Random variable representing noise.
 - $\mathbb{E}[\epsilon] = 0$
 - $\operatorname{\mathsf{var}}[\epsilon] = \mathbb{E}[\epsilon^2] = \sigma_\epsilon^2$

Recall

Variance:

$$\begin{aligned} \operatorname{var}(X) &= \mathbb{E}\left[(X - \mathbb{E}[X])^2 \right] \\ &= \mathbb{E}\left[X^2 - 2X\mathbb{E}[X] + \mathbb{E}^2[X] \right] \\ &= \mathbb{E}\left[X^2 \right] - 2\mathbb{E}^2[X] + \mathbb{E}^2[X] \\ &= \mathbb{E}\left[X^2 \right] - \mathbb{E}^2[X] \end{aligned}$$

And also, since $\mathbb{E}[\epsilon] = 0$, we have:

$$\begin{aligned} \text{var}(\epsilon) &= \mathbb{E}\left[\epsilon^2\right] - \mathbb{E}^2\left[\epsilon\right] \\ \text{var}(\epsilon) &= \mathbb{E}\left[\epsilon^2\right] = \sigma^2 \end{aligned}$$

Goal

- Model the underlying real-life problem.
- ▶ I. e., find \hat{f} s.t. $\hat{f}(x) \approx f(x)$ by reducing MSE $= \mathbb{E}\left[\left(Y \hat{f}(X)\right)^2\right]$

Bias

Difference of average value of prediction realizations (over different realizations of training data) to the true underlying function f(x) for a given *unseen* test point, i.e.:

$$\operatorname{bias}\left[\hat{f}(x)\right] = \mathbb{E}\left[\hat{f}(x)\right] - f(x)$$

Notice that $\hat{f}(x)$ is a random variable affected by the randomness in which we obtain training data.

Variance

Mean Squared Deviation of $\hat{f}(x)$ from its expected value $\mathbb{E}\left[\hat{f}(x)\right]$ over different realizations of training data, i. e.:

$$\operatorname{var}\left(\hat{f}(x)\right) = \mathbb{E}\left[\left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)^2\right]$$

Summary

$$\blacktriangleright \mathsf{MSE} = \mathbb{E}_x \left[\left(y - \hat{f}(x) \right)^2 \right].$$

$$ightharpoonup$$
 bias $\left[\hat{f}(x)\right] = \mathbb{E}\left[\hat{f}(x)\right] - f(x)$.

$$\qquad \qquad \mathsf{var}\left(\hat{f}(x)\right) = \mathbb{E}\left[\left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)^2\right]$$

From the Mean Squared Error (MSE):

$$\mathbb{P}\left[\left(\hat{x}_{i,j}\right)^{2}\right]$$

$$\mathbb{E}\left[\left(y=\hat{f}(x)\right)^2\right]$$

$$\mathbb{E}\left\{\left(y-f(x)\right)\right\}$$

$$\mathbb{E}\left[\left(y - \hat{f}(x)\right)^{2}\right] = \mathbb{E}\left[\left(f(x) + \epsilon - \hat{f}(x)\right)^{2}\right]$$

Since expectation is linear and ϵ and \hat{f} are independent:

 $= \mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)^{2}\right] + 2\mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)\epsilon\right] + \mathbb{E}\left[\epsilon^{2}\right]$

 $= \mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)^{2}\right] + 2\mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)\right] \underbrace{\mathbb{E}\left[\epsilon\right]}_{\epsilon} + \sigma_{\epsilon}^{2}$

 $\mathsf{MSE} = \mathbb{E}\left|\left(y - \hat{f}(x)\right)^2\right| = \mathbb{E}\left|\left(f(x) - \hat{f}(x)\right)^2\right| + \sigma_\epsilon^2$

$$\mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)^{2}\right] = \mathbb{E}\left[\left(f(x) - \mathbb{E}\left[\hat{f}(x)\right] + \mathbb{E}\left[\hat{f}(x)\right] - \hat{f}(x)\right)^{2}\right]$$

$$= I$$

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$$= \mathbb{E}\left[\left(\left(f(x) - \mathbb{E}\left[\hat{f}(x)\right]\right) - \left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)\right)^{2}\right] =$$

$$= \mathbb{E}\left[\underbrace{\left(\mathbb{E}\left[\hat{f}(x)\right] - f(x)\right)^{2}}_{\text{total loss at least total loss}} + \left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)^{2} - 2\underbrace{\left(f(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)}_{\text{total loss at least total loss at least loss at l$$

$$= \mathbb{E}\left[\underbrace{\left(\mathbb{E}\left[\hat{f}(x)\right] - f(x)\right)^{2}}_{\text{squared bias} = \text{ cte}} + \left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)^{2} - 2\underbrace{\left(f(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)}_{\text{cte}} \left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)\right]$$

$$= \underbrace{\left(\mathbb{E}\left[\hat{f}(x)\right] - f(x)\right)^{2}}_{\text{squared bias} = \text{ cte}} + \mathbb{E}\left[\left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)^{2}\right] - 2\left(f(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)\underbrace{\mathbb{E}\left[\left(\hat{f}(x) - \mathbb{E}\left[\hat{f}(x)\right]\right)\right]}_{\text{cte}}$$

 $=\mathbb{E}[\hat{f}(x)]-\mathbb{E}[\hat{f}(x)]=0$

$$\mathsf{bias}^2[\hat{f}(x)]$$
 $\mathsf{var}[\hat{f}(x)]$

$$\mathbb{E}\left[\left(f(x)-\hat{f}(x)\right)^2\right]=\mathsf{bias}^2\left[\hat{f}(x)\right]+\mathsf{var}\left(\hat{f}(x)\right)$$

$$\left|\mathbb{E}\left[\left(f(x)-\hat{f}(x)
ight)^2
ight]=\mathsf{bias}^2\left[\hat{f}(x)
ight]+\mathsf{var}\left(\hat{f}(x)
ight)
ight|$$

Since

hence, we have

 $\mathsf{MSE} = \mathbb{E}\left[\left(y - \hat{f}(x)\right)^2\right] = \mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)^2\right] + \sigma_\epsilon^2$

 $\mathbb{E}\left[\left(f(x) - \hat{f}(x)\right)^{2}\right] = \mathsf{bias}^{2}\left[\hat{f}(x)\right] + \mathsf{var}\left(\hat{f}(x)\right)$

 $\mathsf{MSE} = \mathsf{bias}^2 \left[\hat{f}(x) \right] + \mathsf{var} \left(\hat{f}(x) \right) + \sigma_\epsilon^2$



- and we've just shown that

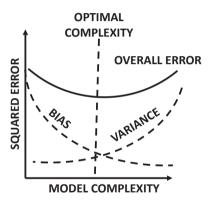
Bias-Variance Equation

Let E[MSE] be the expected mean-squared error of the fixed set of test instances over different samples of training data sets.

$$E[MSE] = Bias^2 + Variance + Noise$$
 (2)

- In linear models, the bias component will contribute more to ${\cal E}[MSE].$
- In polynomial models, the variance component will contribute more to $E[MSE]. \label{eq:energy}$
- We have a trade-off, when it comes to choosing model complexity!

The Bias-Variance Trade-Off



▶ Optimal point of model complexity is somewhere in middle.

Bias-Variance Trade-off: Setup

- ightharpoonup Imagine you are given the true distribution $\mathcal B$ of training data (including labels).
- You have a principled way of sampling data sets $\mathcal{D} \sim \mathcal{B}$ from the training distribution.
- Imagine you create an infinite number of training data sets (and trained models) by repeated sampling.
- You have a fixed set T of unlabeled test instances.
 - The test set ${\mathcal T}$ does not change over different training data sets.
 - Compute prediction of each instance in $\ensuremath{\mathcal{T}}$ for each trained model.

Informal Definition of Bias

- ▶ Compute averaged prediction of each test instance x over different training models $g(x, \mathcal{D})$.
- New Averaged prediction of test instance will be different from true (unknown) model f(x).
- ▶ Difference between (averaged) $g(x, \mathcal{D})$ and f(x) caused by erroneous assumptions/simplifications in modeling \Rightarrow Bias
 - **Example:** Linear simplification to polynomial model causes bias.
 - If the true (unknown) model f(x) were an order-4 polynomial, and we used any polynomial of order-4 or greater in $g(x,\mathcal{D})$, bias would be 0.

Informal Definition of Variance

- ▶ The value $g(x, \mathcal{D})$ will vary with \mathcal{D} for fixed x.
 - The prediction of the same test instance will be different over different trained models.
- ► All these predictions cannot be simultaneously correct ⇒ Variation contributes to error
- ▶ Variance of $g(x, \mathcal{D})$ over different training data sets \Rightarrow Model Variance
 - Example: Linear model will have low variance.
 - Higher-order model will have high variance.

Bias-Variance Equation

Let E[MSE] be the expected mean-squared error of the fixed set of test instances over different samples of training data sets.

$$E[MSE] = \mathsf{Bias}^2 + \mathsf{Variance} + \mathsf{Noise} \tag{3}$$

- In linear models, the bias component will contribute more to ${\cal E}[MSE].$
- In polynomial models, the variance component will contribute more to $E[MSE]. \label{eq:energy}$
- We have a trade-off, when it comes to choosing model complexity!

Cross-Fold Validation

Model Generalization in Neural Networks

- The recent success of neural networks is made possible by increased data.
 - Large data sets help in generalization.
- In a neural network, increasing the number of hidden units in intermediate layers tends to increase complexity.
- Increasing depth often helps in reducing the number of units in hidden layers.
- ▶ Proper design choices can reduce overfitting in complex models ⇒ Better to use complex models with appropriate design choices

How to Detect Overfitting

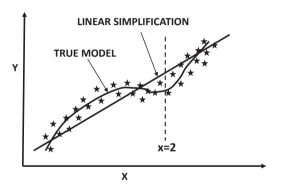
- The error on test data might be caused by several reasons.
 - Other reasons might be bias (underfitting), noise, and poor convergence.
- Overfitting shows up as a large gap between in-sample and out-of-sample accuracy.
- First solution is to collect more data.
 - More data might not always be available!

Penalty-Based Regularization

- ► Key techniques to improve generalization in NNs:
 - Penalty-based regularization.
 - Constraints like shared parameters.
 - Using ensemble methods like Dropout.
 - Adding noise and stochasticity to input or hidden units.

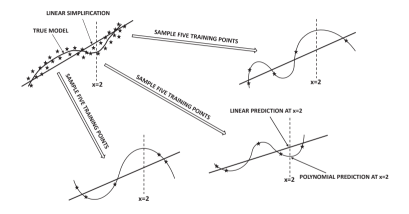
Penalty-Based Regularization

Revisiting Example: Predict y from x



- ▶ First impression: Polynomial model such as $y = w_0 + w_1 x + w_2 x^2 + w_3 x^3 + w_4 x^4$ is "better" than linear model $y = w_0 + w_1 x$.
 - However, with less data, using the linear model is better.

Penalty-Based Regularization



lacktriangle Zero error on training data but wildly varying predictions of x=2

Economy in Parameters

- A lower-order model has economy in parameters.
 - A linear model uses two parameters, whereas an order-4 model uses five parameters.
 - Economy in parameters discourages overfitting.
- Choosing a neural network with fewer units per layer enforces economy.
- Reducing the number of parameters is a hard penalty.
- ▶ We can also penalize parameters in a *soft* way.

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What About Supervised Pre-training

L_2 Regularization and Soft Economy vs Hard Economy

- Fixing the architecture up front is an inflexible solution.
- ▶ A softer solution uses a larger model but imposes a (tunable) penalty on parameter use.

$$\hat{y} = \sum_{i=0}^{d} w_i x^i \tag{4}$$

- ▶ Loss function: $L = \sum_{(x,y) \in \mathcal{D}} (y \hat{y})^2 + \underbrace{\lambda \cdot \sum_{i=0}^d w_i^2}_{L_2 \text{Regularization}}$
- ▶ The (tuned) value of λ decides the level of regularization.
- ► Softer approach with a complex model performs better!

Effect on Updates

For learning rate α , effect on update is to multiply parameter with $(1 - \alpha \lambda) \in (0, 1)$.

$$w_i \Leftarrow w_i (1 - \alpha \lambda) - \alpha \frac{\partial L}{\partial w_i}$$

- Interpretation: Decay-based forgetting!
- Unless a parameter is important, it will have small absolute value.
 - Model decides what is important.
 - Better than inflexibly deciding up front.
- ▶ A forgetting mechanism prevents a model from *memorizing* the training data, because only significant and repeated updates will be reflected in the weights.

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What About Supervised Pre-training?

L_1 -Regularization

In L₁-regularization, an L₁-penalty is imposed on the loss function.

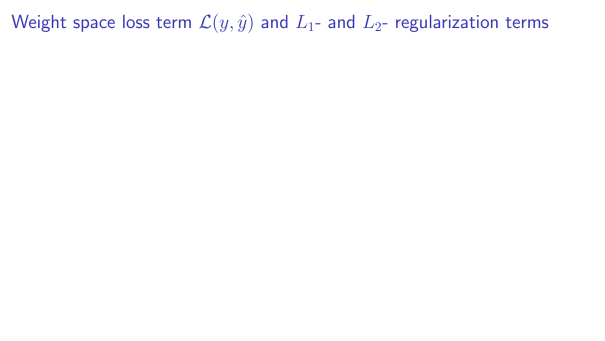
$$L = \sum_{(x,y)\in\mathcal{D}} (y - \hat{y})^2 + \lambda \cdot \sum_{i=0}^{d} |w_i|_1$$

▶ Update has slightly different form (define the update equation at least for the case when $w_i \neq 0$:

$$w_i \Leftarrow w_i - \alpha \lambda s_i - \alpha \frac{\partial L}{\partial w_i}$$

▶ The value of s_i is the partial derivative of $|w_i|$ w.r.t. w_i :

$$s_i = \begin{cases} -1 & w_i < 0 \\ +1 & w_i > 0 \end{cases}$$



L_1 - or L_2 -Regularization?

- $ightharpoonup L_1$ -regularization leads to sparse parameter learning.
 - Zero values of w_i can be dropped.
 - Equivalent to dropping edges from neural network.
- $ightharpoonup L_2$ -regularization generally provides better performance.
- $ightharpoonup L_2$ is differentiable and can be used in different techniques.

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What About Supervised Pre-training

Connections with Noise Injection

- ▶ L_2 -regularization with parameter λ is equivalent to adding Gaussian noise with variance λ to input.
 - Intuition: Bad effect of noise will be minimized with simpler models (smaller parameters).
 - Proof in book.
- Result is only true for single layer network (linear regression).
 - Main value of result is in providing general intuition.
- Similar results can be shown for denoising autoencoders.

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What About Supervised Pre-training

Penalizing Hidden Units

- One can also penalize hidden units (activations).
- ▶ Applying L_1 -penalty leads to sparse activations.
- Straightforward modification of backpropagation.
 - Penalty contributions from hidden units are picked up in backward phase.

$$L' = L + \lambda \sum_{i=1}^{M} |h_i| \tag{5}$$

Where:

- M is the total number of units in the network.
- \blacktriangleright h_i is the value of the *i*th hidden unit.
- \triangleright λ is the regularization parameter.

Ensemble Methods

- ► Inspired in Bias-Variance trade-off.
- Try to reduce either the bias or the variance without affecting the other component.
- Ensemble methods are commonly used in Machine Learning.
- ► Two examples:
 - Bagging Variance reduction.
 - Boosting Bias reduction.

Ensemble Methods

- ▶ Most ensemble methods in NNs are focused on variance reduction.
- ► This is because neural networks are valued for their ability to build arbitrarily complex models with relatively low bias.
- But arbitrarily complex models lead to high variance:

OVERFITTING

Therefore, the goal of most ensemble methods in NN is variance reduction.

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What About Supervised Pre-training

- ▶ If a sufficient number of samples is available, after all, the variance of most types of statistical estimates can be asymptotically reduced to zero.
- ▶ One approach: Predict one instance repeatedly using different training data sets.
- ▶ With a sufficient large number of training data sets is used, the variance can be reduced to zero (infinite source of data).
- ▶ Although we don't have infinite number of instances, an imperfect simulation of the aforementioned methodology has still better variance characteristics than a single execution of the model on the entire training data set.

- ► The predictions on a particular test instance, which are obtained from the models built with different training sets, are then averaged to create the final prediction.
- ▶ One can average either the real-valued prediction (e.g., probability estimates of class labels) or the discrete predictions.
- In the case of real-valued predictions, better results are sometimes obtained by using the median of the values.

Bagging:

- Sample with replacement.
- Sample size: s (common s=m, in which case we have sample size equals training data size).
- ▶ When s=m, we'll have duplicates and about a fraction $(1-1/m)^m \approx 1/e$ of the original data set will not be included.
- Repeat k times. Apply each of the k models to a given test instance.
- Average the results to yield a single robust prediction.
- ▶ Common: s = m; but best results with $s \ll m$.

Sub-sampling:

- ► Sample without replacement.
- Predictions are averaged.
- Essential to choose s < m, since s = m would imply same training data set and identical results.

- ▶ All the variance cannot be removed by using bagging or sub-sampling.
- ▶ Predictions of test instances from different samples will be positively correlated.
- ► The average of a set of random variables that are positively correlated will always have a variance that is proportional to the level of correlation.

Bagging and Sub-sampling are imperfect simulations of drawing the training data from a base distribution.

Nevertheless, the variance of this approach is still lower than that of constructing a single model on the entire training data set.

- ightharpoonup Main challenge is to construct multiple training models ightarrow highly inefficient.
- ightharpoonup But it can be fully parallelized ightharpoonup use with GPUs.

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Parametric Model Selection and Averaging

- ► Problems:
 - Large number of hyper-parameters and configurations.
 - Sensitivity to some choices such as activation functions.
- ▶ Strategy: Hold out a portion of the training data.
- Select the model out of the pool providing highest performance.

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What About Supervised Pre-training

Basic Dropout Training Procedure

- For each training instance do:
 - Sample each node in the network in each layer (except output layer) with probability p.
 - Keep only edges for which both ends are included in network.
 - Perform forward propagation and backpropagation only on sampled network.
- Note that weights are shared between different sampled networks.
- ▶ A different neural network is used for every small mini-batch of training examples.
- The number of NNs is rather large in Dropout.

Basic Dropout Testing Procedures

- First procedure:
 - Perform repeated sampling (like training) and average results.
 - Geometric averaging for probabilistic outputs (averaging log-likelihood)
- ▶ Second procedure with *weight scaling inference rule* (more common):
 - Multiply weight of each outgoing edge of a sampled node i with its sampling probability p_i .
 - Perform single inference on full network with down-scaled weights.

Why Does Dropout Help?

- By dropping nodes, we are forcing the network to learn without the presence of some inputs (in each layer).
- ▶ Will resist co-adaptation, unless the features are truly synergistic.
- Will create many (smaller) groups of self-sufficient predictors.
- ▶ Many groups of self-sufficient predictors will have a model-averaging effect.

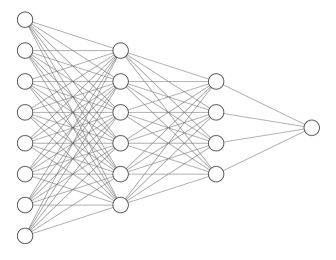
The Regularization Perspective

- One can view the dropping of a node as the same process as adding masking noise.
 - Noise is added to both input and hidden layers.
- ► Adding noise is equivalent to regularization.
- ▶ Forces the weights to become more spread out.
 - Updates are distributed across weights based on sampling.

Practical Aspects of Dropout

- ▶ Typical dropout rate (i.e., probability of exclusion) is somewhere between 20% to 50%.
- Better to use a larger network with Dropout to enable learning of independent representations.
- Dropout is applied to both input layers and hidden layers.
- Large learning rate with decay and large momentum.
- Impose a max-norm constraint on the size of network weights.
 - Norm of input weights to a node upper bounded by constant $\emph{c}.$

Feature Co-Adaptation



 $\mbox{Input Layer} \in \mathbb{R}^8 \qquad \mbox{Hidden Layer} \in \mathbb{R}^6 \quad \mbox{Hidden Layer} \in \mathbb{R}^4 \quad \mbox{Output Layer} \in \mathbb{R}^1$

One-Way Adaptation

- Consider a single-hidden layer neural network.
 - All edges into and out of half the hidden nodes are fixed to random values.
 - Only the other half are updated during backpropagation.
- Half the features will adapt to the other half (random features).
- ► Feature co-adaptation is natural in neural networks where rate of training varies across different parts of network over time.
 - Partially a manifestation of training inefficiency (over and above true synergy).

Why is Feature Co-Adaptation Bad?

- We want features working together only when essential for prediction.
 - We do not want features adjusting to each other because of inefficiencies in training.
 - Does not generalize well to new test data.
- ▶ We want many groups of minimally essential features for robust prediction ⇒ Better redundancies.
- ▶ We do not want a *few* large and inefficiently created groups of co-adapted features.

Feature Co-Adaptation

- ► The process of training a neural network often leads to a high level of dependence among features.
- Different parts of the network train at different rates:
 - Causes some parts of the network to adapt to others.
- This is referred to as feature co-adaptation.
- ► Uninformative dependencies are sensitive to nuances of specific training data ⇒ **OVERFITTING**

Summary

Introduction

The Bias-Variance Trade-Off

Generalization Issues in Model Tuning and Evaluation

Penalty-Based Regularization

 L_2 Regularization and Soft Economy \emph{vs} Hard Economy L_1 -Regularization

Penalizing Hidden Units: Learning Sparse Representations

Ensemble Methods

Bagging and Sub-sampling Parametric Model Selection and Averagin Dropout

Data Perturbation Ensembles

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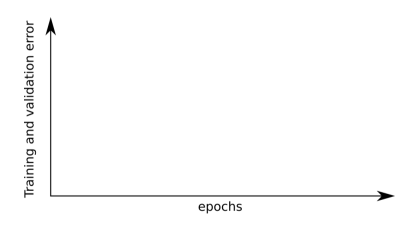
Data Perturbation Ensembles

- ▶ Most of the ensemble techniques discussed so far are either sampling-based ensembles of model-centric ensembles.
- Dropout can be considered an ensemble that adds noise to the data in an indirect way.
- ightharpoonup Simplest case ightharpoonup noise is added to input data and weights are trained on the disturbed scenario.
- Repeat the process and average results.
- This is the generci ensemble method, not specific to neural networks.
- If one wants to add noise to hidden-layers, it must be carefully calibrated.
- Dropout indirectly adds noise to hidden layers by randomly dropping nodes.

Data Perturbation Ensembles

- ▶ Data augmentation can often greatly improve the accuracy of a learner by increasing its generalization power.
- ▶ But they are not perturbation schemes because the augmented examples are created with a calibrated procedure and understanding of the domain at hand.

Early Stopping

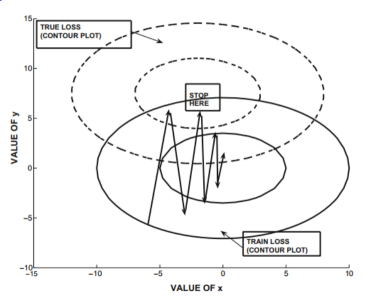


Early Stopping

Motivation:

- ▶ We use optimization to train NNs until convergence.
- Optimizes the loss on training data.
- ▶ Not necessarily on the out-of-sample test data.
- Final steps cause overfitting and generalization problems.
- Almost always used.

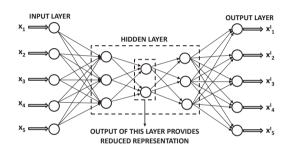
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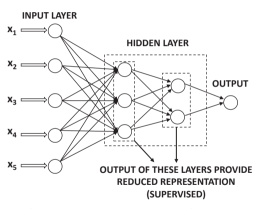


Unsupervised Pre-training – Importance of Initialization

- Bad initializations can lead to unstable convergence.
- ▶ Typical approach is to initialize to a Gaussian with variance 1/r, where r is the indegree of the neuron.
 - Xavier initialization uses both indegree and outdegree.
- ▶ Pretraining goes beyond these simple initializations *by using the training data*.

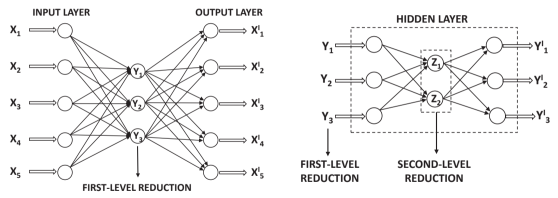
Types of Base Applications





▶ Both the two neural architectures use almost the same pretraining procedure

Layer-Wise Pretraining a Deep Autoencoder



Pretraining deep autoencoder helps in convergence issues

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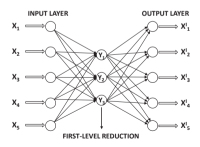
Types of Pretraining

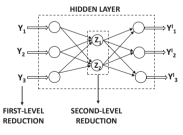
- Unsupervised pretraining: Use training data without labels for initialization.
 - Improves convergence behavior.
 - Regularization effect.
- ► Supervised pretraining: Use training data with labels for initialization.
 - Improves convergence but might overfit.
- Focus on unsupervised pretraining.

Pretraining a Supervised Learner

For a supervised learner with k hidden layers:

- Remove output layer and create an autoencoder with (2k-1) hidden layers.
- Pretrain autoencoder as discussed in previous slide.
- Keep only weights from encoder portion and cap with output layer.
- Pretrain only output layer.
- Fine-tune all layers.





Why Does Pretraining Work?

- Pretraining already brings the activations of the neural network to the manifold of the data distribution.
- Features correspond to repeated patterns in the data.
- ► Fine-tuning learns to combine/modify relevant ones for inference.
 - Pretraining initializes the problem closer to the basin of global optima.
 - Hinton: "To recognize shapes, first learn to generate images."

Thank you! tvieira@ic.ufal.br