CSCE 970 Lecture 3: Regularization

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Introduction

- Machine learning can generally be distilled to an optimization problem
- Choose a classifier (function, hypothesis) from a set of functions that minimizes an objective function
- Clearly we want part of this function to measure performance on the training set, but this is insufficient



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Outline

- Types of machine learning problems
- Loss functions
- Generalization performance vs training set performance
- Overfitting
- Regularization
- Estimating generalization performance



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Machine Learning Problems

- Supervised Learning: Algorithm is given labeled training data and is asked to infer a function (hypothesis) from a family of functions (e.g., set of all ANNs) that is able to predict well on new, unseen examples
 - Classification: Labels come from a finite, discrete set
 - Regression: Labels are real-valued
- Unsupervised Learning: Algorithm is given data without labels and is asked to model its structure
 - Clustering, density estimation
- Reinforcement Learning: Algorithm controls an agent that interacts with its environment and learns good actions in various situations



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Measuring Performance

 In any learning problem, need to be able to quantify performance of an algorithm

 In supervised learning, we often use a loss function (or error function) $\ensuremath{\mathcal{J}}$ for this task

 Given instance x with true label y, if the learner's prediction on x is \hat{y} , then

 $\mathcal{J}(y,\hat{y})$

is the loss on that instance



Measuring Performance Examples of Loss Functions

• **0-1 Loss:** $\mathcal{J}(y, \hat{y}) = 1$ if $y \neq \hat{y}$, 0 otherwise

• Square Loss: $\mathcal{J}(y,\hat{y}) = (y - \hat{y})^2$

• Cross-Entropy: $\mathcal{J}(y, \hat{y}) = -y \ln \hat{y} - (1 - y) \ln (1 - \hat{y})$ (y and \hat{y} are considered probabilities of a '1' label; generalizes to multi-class.)

• Hinge Loss: $\mathcal{J}(y, \hat{y}) = \max(0, 1 - y \hat{y})$ (used sometimes for large margin classifiers like SVMs)

All non-negative

Measuring Performance

 \bullet Given a loss function ${\cal J}$ and a training set ${\cal X},$ the total loss of the classifier h on \mathcal{X} is

$$\textit{error}_{\mathcal{X}}(\textit{h}) = \sum_{\textit{x} \in \mathcal{X}} \mathcal{J}(\textit{y}_{\textit{x}}, \hat{\textit{y}}_{\textit{x}}) \enspace ,$$

where y_x is x's label and \hat{y}_x is h's prediction

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Measuring Performance

• More importantly, the learner needs to **generalize** well: Given a new example drawn iid according to unknown probability distribution \mathcal{D} , we want to minimize h's expected loss:

$$error_{\mathcal{D}}(h) = \mathsf{E}_{x \sim \mathcal{D}} \left[\mathcal{J}(y_x, \hat{y}_x) \right]$$

 Is minimizing training loss the same as minimizing expected loss?

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Measuring Performance Expected vs Training Loss

- Sufficiently sophisticated learners (decision trees, multi-layer ANNs) can often achieve arbitrarily small (or zero) loss on a training set
- A hypothesis (e.g., ANN with specific parameters) h overfits the training data \mathcal{X} if there is an alternative hypothesis h' such that

$$error_{\mathcal{X}}(h) < error_{\mathcal{X}}(h')$$

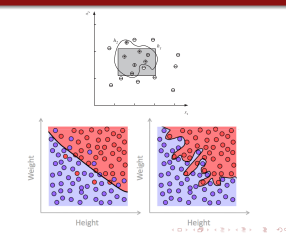
and

$$error_{\mathcal{D}}(h) > error_{\mathcal{D}}(h')$$

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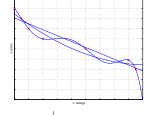
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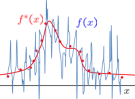
Measuring Performance Overfitting



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Measuring Performance





To generalize well, need to balance training accuracy with simplicity

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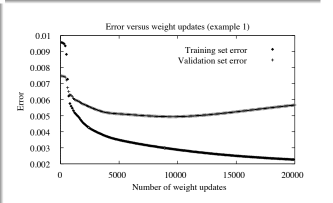
Regularization Causes of Overfitting

ullet Generally, if the set of functions ${\cal H}$ the learner has to choose from is complex relative to what is required for correctly predicting the labels of X, there's a larger chance of overfitting due to the large number of "wrong" choices in ${\cal H}$

• Could be due to an overly sophisticated set of functions

- E.g., can fit any set of *n* real-valued points with an (n-1)-degree polynomial, but perhaps only degree 2 is needed
- E.g., using an ANN with 5 hidden layers to solve the logical AND problem
- Could be due to training an ANN too long
 - Over-training an ANN often leads to weights deviating far from zero
 - Makes the function more non-linear, and more complex
- Often, a larger data set mitigates the problem

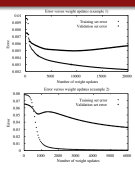
Regularization Causes of Overfitting: Overtraining



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Regularization Early Stopping



- Danger of stopping too soon
 - "Patience" parameter in Algorithm 7.1
- Can re-start with small, random weights (Algorithm 7.2)

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Regularization Parameter Norm Penalties

• Still want to minimize training loss, but balance it against a complexity penalty on the parameters used:

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

• $\alpha \in [0, \infty)$ weights loss $\mathcal J$ against penalty Ω

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Regularization

Parameter Norm Penalties: L² Norm

- $\Omega(\theta) = (1/2) \|\theta\|_2^2$, i.e., sum of squares of network's weights
- Since $\theta = w$, this becomes

$$\tilde{\mathcal{J}}(\mathbf{w}; \mathcal{X}, \mathbf{y}) = (\alpha/2)\mathbf{w}^{\top}\mathbf{w} + \mathcal{J}(\mathbf{w}; \mathcal{X}, \mathbf{y})$$

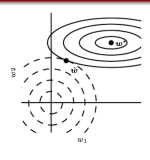
 As weights deviate from zero, activation functions become more nonlinear, which is higher risk of overfitting

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Regularization

Parameter Norm Penalties: L2 Norm



- w^* is optimal for \mathcal{J} , 0 optimal for regularizer
- \mathcal{J} less sensitive to w_1 , so \tilde{w} (optimal for $\tilde{\mathcal{J}}$) closer to 0 than w_2

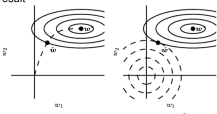
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Parameter Norm Penalties: L² Norm

Related to early stopping: For linear model and square loss, get trajectory of weight updates that, on average, lands in a similar result



Length τ of trajectory related to weight α in L^2 regularizer

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Parameter Norm Penalties: L1 Norm

• $\Omega(\theta) = \|\theta\|_1$, i.e., sum of absolute values of network's weights

$$\tilde{\mathcal{J}}(\mathbf{w}; \mathcal{X}, \mathbf{y}) = \alpha \|\mathbf{w}\|_1 + \mathcal{J}(\mathbf{w}; \mathcal{X}, \mathbf{y})$$

- As with L^2 regularization, penalizes large weights
- Unlike L^2 regularization, can drive some weights to zero
 - Sparse solution
 - Sometimes used in feature selection (e.g., LASSO algorithm)



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Regularization Data Augmentation

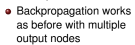
- If \mathcal{H} is powerful and \mathcal{X} is small, then a learner can choose some $h \in \mathcal{H}$ that fits the idiosyncrasies or noise in the data
- Deep ANNs would like to have at least thousands or tens of thousands of data points
- In classification of high-dimensional data (e.g., image classification), expect the classifier to tolerate transformations and noise
 - ⇒ Can artificially enlarge data set by duplicating existing instances and applying transformations
 - Translating, rotating, scaling
 - Be careful to to change the class, e.g., "b" vs "d" or "6"
 - ⇒ Can also apply noise injection to input or even hidden lavers



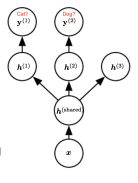
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Regularization Multitask Learning

 If multiple tasks share generic parameters. initially process inputs via shared nodes, then do final processing via task-specific nodes



 Serves as a regularizer since parameter tuning of shared nodes is based on backpropagated error from multiple tasks



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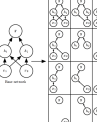
Regularization Dropout

 Imagine if, for a network, we could average over all networks with each subset of nodes deleted

 Analogous to bagging, where we average over ANNs trained on random samples of \mathcal{X}

• In each training iteration, sample a random bit vector μ , which determines which nodes are used (e.g.,

 $P(\mu_i = 1) = 0.8$ for input unit, 0.5 for hidden unit)



 Make predictions by sampling new vectors μ and averaging



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Regularization Other Approaches

• Parameter Tying: If two learners are learning the same task but different distributions, can tie their parameters together

• If $w^{(A)}$ are weights for task A and $w^{(B)}$ are weights for task B, then can use regularization term $\Omega(\mathbf{w}^{(A)}, \mathbf{w}^{(B)}) = \|\mathbf{w}^{(A)} - \mathbf{w}^{(B)}\|_2^2$

• Parameter Sharing: When detecting objects in an image, the same recognizer should apply invariant to translation

• Can train a single detector (subnetwork) for the object (e.g., cat) by training full network on multiple images with translated cats, where the cat detector subnetworks share parameters (single copy, used multiple times)

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Regularization Other Approaches (cont'd)

 Sparse Representations: Instead of penalizing large weights, penalize large outputs of hidden nodes:

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

where h is the vector of hidden unit outputs

Estimating Generalization Performance

 Before setting up an experiment, need to understand exactly what the goal is

- · Estimate the generalization performance of a hypothesis
- Tuning a learning algorithm's parameters
- Comparing two learning algorithms on a specific task
- Will never be able to answer the question with 100% certainty
 - Due to variances in training set selection, test set selection, etc.
 - Will choose an estimator for the quantity in question, determine the probability distribution of the estimator, and bound the probability that the estimator is way off
 - Estimator needs to work regardless of distribution of training/testing data

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Estimating Generalization Performance

 Need to note that, in addition to statistical variations, what we determine is limited to the application that we are studying

- E.g., if ANN₁ better than ANN₂ on speech recognition, that means nothing about video analysis
- In planning experiments, need to ensure that training data not used for evaluation
 - I.e., don't test on the training set!
 - Will bias the performance estimator
 - Also holds for validation set used for early stopping, tuning parameters, etc.
 - · Validation set serves as part of training set, but not used for model building



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Confidence Intervals

ntroduction

Let $error_{\mathcal{D}}(h)$ be 0-1 loss of h on instances drawn according to distribution \mathcal{D} . If

- V contains N examples, drawn independently of h and each other
- N > 30

Then with approximately 95% probability, $error_{\mathcal{D}}(h)$ lies in

$$error_{\mathcal{V}}(h) \pm 1.96 \sqrt{\frac{error_{\mathcal{V}}(h)(1-error_{\mathcal{V}}(h))}{N}}$$

E.g. hypothesis h misclassifies 12 of the 40 examples in test set \mathcal{V} :

$$error_{\mathcal{V}}(h) = \frac{12}{40} = 0.30$$

Then with approx. 95% confidence, $error_{\mathcal{D}}(h) \in [0.158, 0.442]$

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Confidence Intervals (cont'd)

Let $error_{\mathcal{D}}(h)$ be 0-1 loss of h on instances drawn according to distribution \mathcal{D} . If

- V contains N examples, drawn independently of h and each other
- N > 30

Then with approximately c% probability, $error_{\mathcal{D}}(h)$ lies in

$$error_{\mathcal{V}}(h) \pm z_c \sqrt{\frac{error_{\mathcal{V}}(h)(1 - error_{\mathcal{V}}(h))}{N}}$$

	50%						
z_c :	0.67	1.00	1.28	1.64	1.96	2.33	2.58

Why?

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$error_{\mathcal{V}}(h)$ is a Random Variable

Repeatedly run the experiment, each with different randomly drawn V (each of size N)

Probability of observing r misclassified examples:

Binomial distribution for
$$n = 40, p = 0.3$$

0.14

0.12

0.1

0.08

0.06

0.04

0.02

0 5 10 15 20 25 30 35 40

$$P(r) = \binom{N}{r} error_{\mathcal{D}}(h)^r (1 - error_{\mathcal{D}}(h))^{N-r}$$

I.e., let $error_{\mathcal{D}}(h)$ be probability of heads in biased coin, then P(r) = prob. of getting r heads out of N flips

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Binomial Probability Distribution

 $P(r) = \binom{N}{r} p^r (1-p)^{N-r} = \frac{N!}{r!(N-r)!} p^r (1-p)^{N-r}$

Probability P(r) of r heads in N coin flips, if p = Pr(heads)

• Expected, or mean value of X, E[X] (= # heads on Nflips = # mistakes on N test exs), is

$$E[X] \equiv \sum_{i=0}^{N} iP(i) = Np = N \cdot error_{\mathcal{D}}(h)$$

Variance of X is

$$Var(X) \equiv E[(X - E[X])^2] = Np(1 - p)$$

• Standard deviation of X, σ_X , is

$$\sigma_X \equiv \sqrt{E[(X - E[X])^2]} = \sqrt{Np(1 - p)}$$

Approximate Binomial Dist. with Normal

CSCE 970 Lecture 3: $error_{\mathcal{V}}(h) = r/N$ is binomially distributed, with

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• mean $\mu_{error_{\mathcal{V}}(h)} = error_{\mathcal{D}}(h)$ (i.e., unbiased est.) • standard deviation $\sigma_{error_{\mathcal{V}}(h)}$

standard deviation
$$\sigma_{error_{\mathcal{V}}(h)}$$

$$\sigma_{error_{\mathcal{V}}(h)} = \sqrt{\frac{error_{\mathcal{D}}(h)(1-error_{\mathcal{D}}(h))}{N}}$$

(increasing N decreases variance)

Want to compute confidence interval = interval centered at $error_D(h)$ containing c% of the weight under the distribution

Approximate binomial by normal (Gaussian) dist:

- mean $\mu_{error_{\mathcal{V}}(h)} = error_{\mathcal{D}}(h)$
- standard deviation $\sigma_{error_V(h)}$

$$\sigma_{error_{\mathcal{V}}(h)} \approx \sqrt{\frac{error_{\mathcal{V}}(h)(1 - error_{\mathcal{V}}(h))}{N_{\bullet} \cdot M_{\bullet} \cdot \frac{1}{2} \cdot \frac{1}{2}}} \approx \sqrt{2} \cdot \sqrt{2}$$

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Normal Probability Distribution

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Introduction

Machine Learning Problems

Measuring Performance

Estimating Generalization Performance

Comparing Learning Algorithms

$$p(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \, \exp\left(-\frac{1}{2} \left(\frac{x-\mu}{\sigma}\right)^2\right)$$

- The probability that *X* will fall into the interval (a, b) is given by $\int_a^b p(x) dx$
- Expected, or mean value of X, E[X], is $E[X] = \mu$
- Variance is $Var(X) = \sigma^2$, standard deviation is $\sigma_X = \sigma$

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Normal Probability Distribution (cont'd)

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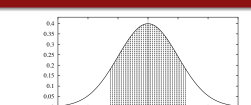
Introduction Outline

Machine Learning Problems

Measuring Performance Regularization

Estimating Generalization Performance Setting Goals

Comparing Learning Algorithms Other/52



80% of area (probability) lies in $\mu \pm 1.28\sigma$ c% of area (probability) lies in $\mu \pm z_c \sigma$

c%:	50%	68%	80%	90%	95%	98%	99%
z_c :	0.67	68% 1.00	1.28	1.64	1.96	2.33	2.58

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Normal Probability Distribution (cont'd)

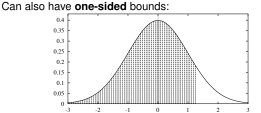
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Introduction Outline

Machine Learning Problems Measuring

Regularization
Estimating
Generalization
Performance
Setting Goals

Comparing Learning Algorithms Other/52



c% of area lies $<\mu+z_c'\,\sigma$ or $>\mu-z_c'\sigma,$ where $z_c'=z_{100-(100-c)/2}$

<i>c</i> %:	50%	68%	80%	90%	95%	98%	99%
z_c' :	0.0	0.47	0.84	1.28	1.64	2.05	2.33

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Confidence Intervals Revisited

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Introduction Outline

Machine Learning Problems

Measuring Performance

Regularization Estimating Generalization Performance

Confidence Interest Comparing Learning Algorithms Ottos/52 If V contains $N \ge 30$ examples, indep. of h and each other Then with approximately 95% probability, $error_V(h)$ lies in

$$\mathit{error}_{\mathcal{D}}(h) \pm 1.96 \sqrt{\frac{\mathit{error}_{\mathcal{D}}(h)(1-\mathit{error}_{\mathcal{D}}(h))}{N}}$$

Equivalently, $error_{\mathcal{D}}(h)$ lies in

$$error_{\mathcal{V}}(h) \pm 1.96 \sqrt{\frac{error_{\mathcal{D}}(h)(1 - error_{\mathcal{D}}(h))}{N}}$$

which is approximately

$$error_{\mathcal{V}}(h) \pm 1.96 \sqrt{\frac{error_{\mathcal{V}}(h)(1-error_{\mathcal{V}}(h))}{N}}$$

(One-sided bounds yield upper or lower error bounds)

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Central Limit Theorem

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Machine Learning Problems

Measuring Performance Regularization

Estimating Generalization Performance Setting Goals

Comparing Learning Algorithms How can we justify approximation?

Consider set of iid random variables Y_1, \ldots, Y_N , all from **arbitrary** probability distribution with mean μ and finite variance σ^2 . Define sample mean $\bar{Y} \equiv (1/N) \sum_{i=1}^n Y_i$

 \bar{Y} is itself a random variable, i.e., result of an experiment (e.g., $error_S(h)=r/N)$

Central Limit Theorem: As $N\to\infty$, the distribution governing \bar{Y} approaches normal distribution with mean μ and variance σ^2/N

Thus the distribution of $error_S(h)$ is approximately normal for large N, and its expected value is $error_D(h)$

(**Rule of thumb:** $N \ge 30$ when estimator's distribution is binomial; might need to be larger for other distributions)

Calculating Confidence Intervals

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Introduction

Machine Learning

Measuring Performance

Estimating Generalization Performance

Comparing Learning Algorithms • Pick parameter to estimate: $error_{\mathcal{D}}(h)$ (0-1 loss on distribution \mathcal{D})

② Choose an estimator: $error_{\mathcal{V}}(h)$ (0-1 loss on independent test set \mathcal{V})

Oetermine probability distribution that governs estimator: error_V(h) governed by binomial distribution, approximated by normal when N ≥ 30

lacktriangledown Find interval (L,U) such that c% of probability mass falls in the interval

ullet Could have $L=-\infty$ or $U=\infty$

• Use table of z_c or z'_c values (if distribution normal)



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Comparing Learning Algorithms

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Introduction Outline

Machine Learning Problems

Measuring Performance

Estimating Generalization

Comparing Learning

K-Fold CV Student's r Distribution

- What if we want to compare two learning algorithms L¹ and L² (e.g., two ANN architectures, two regularizers, etc.) on a specific application?
- Insufficient to simply compare error rates on a single test set
- Use K-fold cross validation and a paired t test



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K-Fold Cross Validation

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Introduction Outline

Problems

Measuring

Performance Regularization

Estimating Generalizatio Performance Comparing

K-Fold CV
Student's r
Distribution
39/52

- A-1 Old C1055 Valldation
 - Partition data set \mathcal{X} into K equal-sized subsets $\mathcal{X}_1, \mathcal{X}_2, \dots, \mathcal{X}_K$, where $|\mathcal{X}_i| \geq 30$
- Provium 1 to K, do (Use \mathcal{X}_i for testing, and rest for training)

 - $\mathcal{T}_i = \mathcal{X} \setminus \mathcal{X}_i$
 - **3** Train learning algorithm L^1 on \mathcal{T}_i to get h_i^1
 - Train learning algorithm L^2 on \mathcal{T}_i to get h_i^2
 - **5** Let p_i^j be error of h_i^j on test set \mathcal{V}_i
 - $p_i = p_i^1 p_i^2$
- **1** Error difference estimate $p = (1/K) \sum_{i}^{K} p_i$



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K-Fold Cross Validation (cont'd)

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Introduction

Machine Learning Problems

Measuring Performance

Generalization
Performance
Comparing

K-Fold CV
Student's r
Distribution

- \Rightarrow Confidence that L^1 is better than L^2 on learning task
- \bullet Use one-sided test, with confidence derived from student's t distribution with K-1 degrees of freedom
- \bullet With approximately c% probability, true difference of expected error between L^1 and L^2 is at most

$$p + t_{c,K-1} s_p$$

where

$$s_p \equiv \sqrt{\frac{1}{K(K-1)} \sum_{i=1}^{K} (p_i - p)^2}$$

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Student's *t* Distribution (One-Sided Test)

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Introduction
Outline
Machine

Problems
Measuring
Performance

Regularization
Estimating
Generalization
Performance
Comparing
Learning

Algorithms

K-Fold CV

Student's r

Distribution

 df
 0.600
 0.700
 0.800
 0.900
 0.950
 0.975
 0.990
 0.995

 1
 0.325
 0.727
 1.376
 3.078
 6.314
 12.706
 31.821
 63.657

 2
 0.289
 0.617
 1.061
 1.886
 2.920
 4.303
 6.965
 9.925

 3
 0.277
 0.584
 0.978
 1.638
 2.353
 3.182
 4.541
 5.841

 4
 0.271
 0.569
 0.941
 1.533
 2.132
 2.776
 3.747
 4.604

 5
 0.267
 0.559
 0.920
 1.476
 2.015
 2.571
 3.365
 4.032

 6
 0.265
 0.553
 0.906
 1.440
 1.943
 2.447
 3.143
 3.707

 7
 0.263
 0.549
 0.896
 1.415
 1.895
 2.365
 2.998
 3.499

 8
 0.262
 0.546
 0.889
 1.397
 1.860
 2.306
 2.891
 3.259

 9
 0.

If $p + t_{c,K-1} s_p < 0$ our assertion that L^1 has less error than L^2 is supported with confidence c

So if K-fold CV used, compute p, look up $t_{c,K-1}$ and check if $p < -t_{c,K-1} \, s_p$

One-sided test; says nothing about L^2 over L^1

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CSCE 970 Lecture 3:

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Learning Problems

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K-Fold CV
Student's r
Distribution

- Say you want to show that learning algorithm L¹ performs better than algorithms L², L³, L⁴, L⁵
- If you use K-fold CV to show superior performance of L^1 over each of L^2, \ldots, L^5 at 95% confidence, there's a 5% chance each one is wrong
- ⇒ There's an over 18.5% chance that at least one is wrong
- ⇒ Our overall confidence is only just over 81%
- Need to account for this, or use more appropriate test

More Specific Performance Measures

 So far, we've looked at a single error rate to compare hypotheses/learning algorithms/etc.

- This may not tell the whole story:
 - 1000 test examples: 20 positive, 980 negative
 - h¹ gets 2/20 pos correct, 965/980 neg correct, for accuracy of (2+965)/(20+980) = 0.967
 - Pretty impressive, except that always predicting
 - $\begin{array}{l} {\rm negative\ yields\ accuracy}=0.980 \\ \bullet \ \ {\rm Would\ we\ rather\ have}\ h^2, \ {\rm which\ gets\ 19/20\ pos\ correct} \end{array}$ and 930/980 neg, for accuracy = 0.949?
 - Depends on how important the positives are, i.e., frequency in practice and/or cost (e.g., cancer diagnosis)



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Confusion Matrices

Break down error into type: true positive, etc.

	Predicted Class					
True Class	Positive	Negative	Total			
Positive	tp: true positive	fn: false negative	p			
Negative	fp: false positive	tn: true negative	n			
Total	p'	n'	N			

Generalizes to multiple classes

Allows one to quickly assess which classes are missed the most, and into what other class

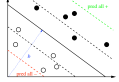


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ROC Curves

• Consider classification via ANN + linear threshold unit

- Normally threshold f(x; w, b) at 0, but what if we changed it?
- Keeping w fixed while changing threshold = fixing hyperplane's slope while moving along its normal vector



- Get a set of classifiers, one per labeling of test set
- Similar situation with any classifier with confidence value, e.g., probability-based



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ROC Curves

Plotting tp versus fp

• Consider the "always -" hyp. What is fp? What is tp? What about the "always +" hyp?

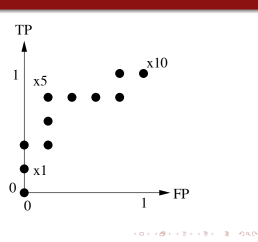
 In between the extremes, we plot TP versus FP by sorting the test examples by the confidence values

Ex	Confidence	label	Ex	Confidence	label
x_1	169.752	+	<i>x</i> ₆	-12.640	_
x_2	109.200	+	<i>x</i> ₇	-29.124	_
<i>x</i> ₃	19.210	_	<i>x</i> ₈	-83.222	_
x_4	1.905	+	<i>x</i> ₉	-91.554	+
<i>x</i> ₅	-2.75	+	x ₁₀	-128.212	_



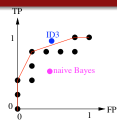
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ROC Curves Plotting *tp* versus *fp* (cont'd)



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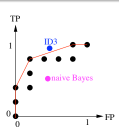
ROC Curves Convex Hull



- The convex hull of the ROC curve yields a collection of classifiers, each optimal under different conditions
 - If FP cost = FN cost, then draw a line with slope |N|/|P|at (0,1) and drag it towards convex hull until you touch it; that's your operating point
 - Can use as a classifier any part of the hull since can randomly select between two classifiers



ROC Curves Convex Hull



- Can also compare curves against "single-point" classifiers when no curves
 - In plot, ID3 better than our SVM iff negatives scarce; nB never better





ROC Curves

• What is the worst possible ROC curve?

 One metric for measuring a curve's goodness: area under curve (AUC):

$$\frac{\sum_{x_{+} \in P} \sum_{x_{-} \in N} I(h(x_{+}) > h(x_{-}))}{|P| \, |N|}$$

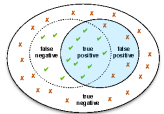
i.e., rank all examples by confidence in "+" prediction, count the number of times a positively-labeled example (from P) is ranked above a negatively-labeled one (from N), then normalize

- What is the best value?
- Distribution approximately normal if |P|, |N| > 10, so can find confidence intervals
- Catching on as a better scalar measure of performance than error rate
- ROC analysis possible (though tricky) with multi-class problems 4 D > 4 D > 4 E > 4 E > E 990

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Precision-Recall Curves

Consider information retrieval task, e.g., web search



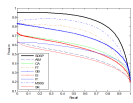
O All documents ✓ relevant 🗶 not relevant O retrieved **precision** = tp/p' = fraction of retrieved that are positive **recall** = tp/p = fraction of positives retrieved



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Precision-Recall Curves (cont'd)

As with ROC, can vary threshold to trade off precision against recall



Can compare curves based on containment

Use F_{β} -measure to combine at a specific point, where β weights precision vs recall:

$$F_{\beta} \equiv (1 + \beta^2) \frac{precision \cdot recall}{(\beta^2 \cdot precision) + recall}$$