# CPSC 340: Machine Learning and Data Mining

Probabilistic Classification Fall 2020

### Admin

- Waiting list people: every should be in!
- Course webpage:
  - <a href="https://www.cs.ubc.ca/~fwood/CS340/">https://www.cs.ubc.ca/~fwood/CS340/</a>
- Homework 1 due tonight.

# Last Time: Training, Testing, and Validation

### Training step:

Input: set of 'n' training examples x; with labels y;
Output: a model that maps from arbitrary x; to a ŷ;

### • Prediction step:

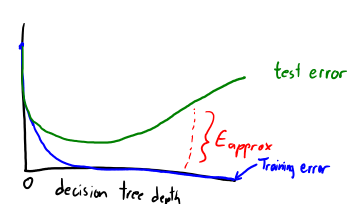
Input: set of '(') testing examples  $\hat{x}_i$  and a model. Output: predictions  $\hat{y}_i$  for the testing examples.

- What we are interested in is the test error:
  - Error made by prediction step on new data.

### Last Time: Fundamental Trade-Off

We decomposed test error to get a fundamental trade-off:

- Where  $E_{approx} = (E_{test} - E_{train})$ .



- E<sub>train</sub> goes down as model gets complicated:
  - Training error goes down as a decision tree gets deeper.
- But E<sub>approx</sub> goes up as model gets complicated:
  - Training error becomes a worse approximation of test error.

### Last Time: Validation Error

- Golden rule: we can't look at test data during training.
- But we can approximate E<sub>test</sub> with a validation error:
  - Error on a set of training examples we "hid" during training.

- Find the decision tree based on the "train" rows.
- Validation error is the error of the decision tree on the "validation" rows.
  - We typically choose "hyper-parameters" like depth to minimize the validation error.

### Overfitting to the Validation Set?

- Validation error usually has lower optimization bias than training error.
  - Might optimize over 20 values of "depth", instead of millions+ of possible trees.
- But we can still overfit to the validation error (common in practice):
  - Validation error is only an unbiased approximation if you use it once.
  - Once you start optimizing it, you start to overfit to the validation set.
- This is most important when the validation set is "small":
  - The optimization bias decreases as the number of validation examples increases.
- Remember, our goal is still to do well on the test set (new data), not the validation set (where we already know the labels).

### Scenario 1:

- "I built a model based on the data you gave me."
- "It classified your data with 98% accuracy."
- "It should get 98% accuracy on the rest of your data."

### Probably not:

- They are reporting training error.
- This might have nothing to do with test error.
- E.g., they could have fit a very deep decision tree.

### • Why 'probably'?

- If they only tried a few very simple models, the 98% might be reliable.
- E.g., they only considered decision stumps with simple 1-variable rules.

### • Scenario 2:

- "I built a model based on half of the data you gave me."
- "It classified the other half of the data with 98% accuracy."
- "It should get 98% accuracy on the rest of your data."

### Probably:

- They computed the validation error once.
- This is an unbiased approximation of the test error.
- Trust them if you believe they didn't violate the golden rule.

### • Scenario 3:

- "I built 10 models based on half of the data you gave me."
- "One of them classified the other half of the data with 98% accuracy."
- "It should get 98% accuracy on the rest of your data."

### Probably:

- They computed the validation error a small number of times.
- Maximizing over these errors is a biased approximation of test error.
- But they only maximized it over 10 models, so bias is probably small.
- They probably know about the golden rule.

### • Scenario 4:

- "I built 1 billion models based on half of the data you gave me."
- "One of them classified the other half of the data with 98% accuracy."
- "It should get 98% accuracy on the rest of your data."

### Probably not:

- They computed the validation error a huge number of times.
- They tried so many models, one of them is likely to work by chance.

### Why 'probably'?

If the 1 billion models were all extremely-simple, 98% might be reliable.

#### Scenario 5:

- "I built 1 billion models based on the first third of the data you gave me."
- "One of them classified the second third of the data with 98% accuracy."
- "It also classified the last third of the data with 98% accuracy."
- "It should get 98% accuracy on the rest of your data."

### Probably:

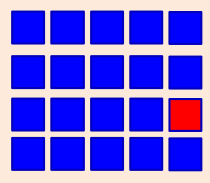
- They computed the first validation error a huge number of times.
- But they had a second validation set that they only looked at once.
- The second validation set gives unbiased test error approximation.
- This is ideal, as long as they didn't violate golden rule on the last third.
- And assuming you are using IID data in the first place.

### Validation Error and Optimization Bias

- Optimization bias is small if you only compare a few models:
  - Best decision tree on the training set among depths 1, 2, 3,..., 10.
  - Risk of overfitting to validation set is low if we try 10 things.
- Optimization bias is large if you compare a lot of models:
  - All possible decision trees of depth 10 or less.
  - Here we're using the validation set to pick between a billion+ models:
    - Risk of overfitting to validation set is high: could have low validation error by chance.
  - If you did this, you might want a second validation set to detect overfitting.
- And optimization bias shrinks as you grow size of validation set.

### Aside: Optimization Bias leads to Publication Bias

Suppose that 20 researchers perform the exact same experiment:

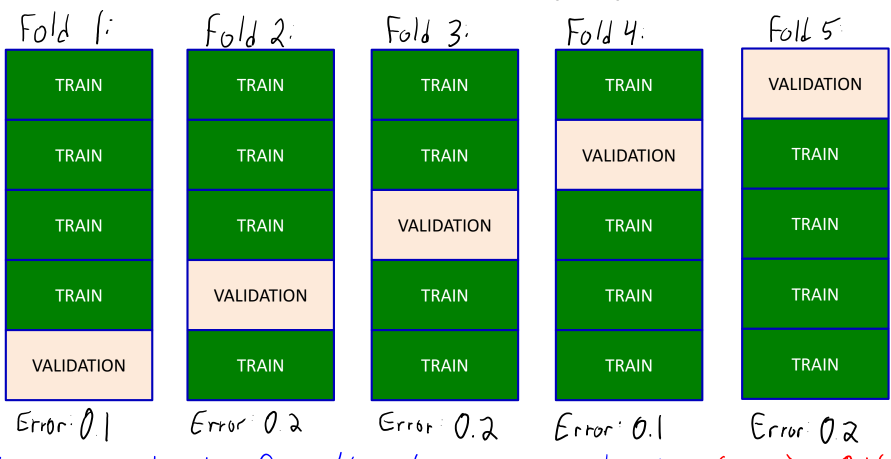


- They each test whether their effect is "significant" (p < 0.05).
  - 19/20 find that it is not significant.
  - But the 1 group finding it's significant publishes a paper about the effect.
- This is again optimization bias, contributing to publication bias.
  - A contributing factor to many reported effects being wrong.

# Cross-Validation (CV)

- Isn't it wasteful to only use part of your data?
- 5-fold cross-validation:
  - Train on 80% of the data, validate on the other 20%.
  - Repeat this 5 more times with different splits, and average the score.

# Cross-Validation (CV)



CV error estimate for this hyper-purameterimean(errors) = 016

### Cross-Validation Pseudo-Code

# To choose depth for depth in 1:20 compute cross-validations core return depth with highest score

To compute 5-fold cross-validation score:

for fold in 1:5

train 80% that doesn't include fold
test on fold
return average test error

### Notes:

- This fits 100 models!
  (20 depths times 5 folds)
- We get one (average) Score for each of the 20 depths.
- Use this score to pick depth

# Cross-Validation (CV)

- You can take this idea further ("k-fold cross-validation"):
  - 10-fold cross-validation: train on 90% of data and validate on 10%.
    - Repeat 10 times and average (test on fold 1, then fold 2,..., then fold 10),
  - Leave-one-out cross-validation: train on all but one training example.
    - Repeat n times and average.
- Gets more accurate but more expensive with more folds.
  - To choose depth we compute the cross-validation score for each depth.
- As before, if data is ordered then folds should be random splits.
  - Randomize first, then split into fixed folds.

(pause)

# The "Best" Machine Learning Model

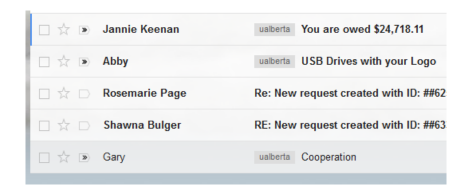
- Decision trees are not always most accurate on test error.
- What is the "best" machine learning model?
- An alternative measure of performance is the generalization error:
  - Average error over all x<sub>i</sub> vectors that are not seen in the training set.
  - "How well we expect to do for a completely unseen feature vector".
- No free lunch theorem (proof in bonus slides):
  - There is **no** "best" model achieving the best generalization error for every problem.
  - If model A generalizes better to new data than model B on one dataset,
     there is another dataset where model B works better.
- This question is like asking which is "best" among "rock", "paper", and "scissors".

# The "Best" Machine Learning Model

- Implications of the lack of a "best" model:
  - We need to learn about and try out multiple models.
- So which ones to study in CPSC 340?
  - We'll usually motivate each method by a specific application.
  - But we're focusing on models that have been effective in many applications.
- Caveat of no free lunch (NFL) theorem:
  - The world is very structured.
  - Some datasets are more likely than others.
  - Model A really could be better than model B on every real dataset in practice.
- Machine learning research:
  - Large focus on models that are useful across many applications.

### Application: E-mail Spam Filtering

- Want a build a system that detects spam e-mails.
  - Context: spam used to be a big problem.





Can we formulate as supervised learning?

# Spam Filtering as Supervised Learning

• Collect a large number of e-mails, gets users to label them.

\$	Hi	CPSC	340	Vicodin	Offer		Spam?
1	1	0	0	1	0		1
0	0	0	0	1	1		1
0	1	1	1	0	0		0
							•••

- We can use  $(y_i = 1)$  if e-mail 'i' is spam,  $(y_i = 0)$  if e-mail is not spam.
- Extract features of each e-mail (like bag of words).
  - $-(x_{ij} = 1)$  if word/phrase 'j' is in e-mail 'i',  $(x_{ij} = 0)$  if it is not.

### Feature Representation for Spam

- Are there better features than bag of words?
  - We add bigrams (sets of two words):
    - "CPSC 340", "wait list", "special deal".
  - Or trigrams (sets of three words):
    - "Limited time offer", "course registration deadline", "you're a winner".
  - We might include the sender domain:
    - <sender domain == "mail.com">.
  - We might include regular expressions:
    - <your first and last name>.

# Review of Supervised Learning Notation

• We have been using the notation 'X' and 'y' for supervised learning:

	\$	Hi	CPSC	340	Vicodin	Offer			Spam?	
X=	1	1	0	0	1	0		19X26	1	)
	0	0	0	0	1	1		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	1	
	0	1	1	1	0	0		1 1-1	(0)-	7 1/3
	)									
							_	73		_

- X is matrix of all features, y is vector of all labels.
  - We use y<sub>i</sub> for the label of example 'i' (element 'i' of 'y').
  - We use x<sub>ii</sub> for feature 'j' of example 'i'.
  - We use  $x_i$  as the list of features of example 'i' (row 'i' of 'X').
    - So in the above  $x_3 = [0 \ 1 \ 1 \ 1 \ 0 \ 0 \ ...].$
    - In practice, only store list of non-zero features for each x<sub>i</sub> (small memory requirement).

### **Probabilistic Classifiers**

- For years, best spam filtering methods used naïve Bayes.
  - A probabilistic classifier based on Bayes rule.
  - It tends to work well with bag of words.
  - Recently shown to improve on state of the art for CRISPR "gene editing" (link).
- Probabilistic classifiers model the conditional probability,  $p(y_i \mid x_i)$ .
  - "If a message has words x<sub>i</sub>, what is probability that message is spam?"
- Classify it as spam if probability of spam is higher than not spam:
  - If  $p(y_i = \text{"spam"} \mid x_i) > p(y_i = \text{"not spam"} \mid x_i)$ 
    - return "spam".
  - Else
    - return "not spam".

To model conditional probability, naïve Bayes uses Bayes rule:

$$\rho(y_i = ||span''||x_i) = \frac{\rho(x_i | y_i = ||span''|)\rho(y_i = ||span''|)}{\rho(x_i)}$$

- So we need to figure out three types of terms:
  - Marginal probability  $p(y_i)$  that an e-mail is spam.
  - Marginal probability  $p(x_i)$  that an e-mail has the set of words  $x_i$ .
  - Conditional probability  $p(x_i | y_i)$  that a spam e-mail has the words  $x_i$ .
    - And the same for non-spam e-mails.

$$\rho(y_i = ||span''||x_i) = \frac{\rho(x_i | y_i = ||span''|)\rho(y_i = ||span''|)}{\rho(x_i)}$$

What do these terms mean?

**ALL E-MAILS** 

(including duplicates)

$$\rho(y_i = ||spam|| ||x_i|) = \frac{\rho(x_i | y_i = ||spam||)\rho(y_i = ||spam||)}{\rho(x_i)}$$

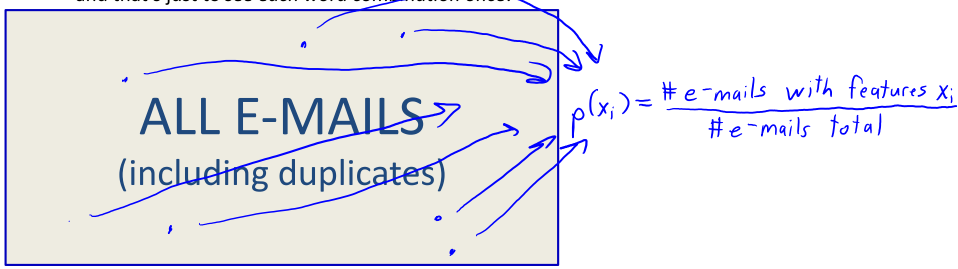
- $p(y_i = "spam")$  is probability that a random e-mail is spam.
  - This is easy to approximate from data: use the proportion in your data.



This is an "estimate" of the true probability. In particular, this formula is a "maximum likelihood estimate" (MLE). We will cover likelihoods and MLEs later in the course.

$$\rho(y_i = ||spam''||x_i) = \frac{\rho(x_i | y_i = ||spam'')\rho(y_i = ||spam'')}{\rho(x_i)}$$

- $p(x_i)$  is probability that a random e-mail has features  $x_i$ :
  - Hard to approximate: with 'd' words we need to collect 2<sup>d</sup> "coupons",
     and that's just to see each word combination once.



$$p(y_i = ||span|| ||x_i||) = \frac{p(x_i | y_i = ||span||)}{p(x_i)}$$

- $p(x_i)$  is probability that a random e-mail has features  $x_i$ :
  - Hard to approximate: with 'd' words we need to collect 2<sup>d</sup> "coupons", but it turns out we can ignore it:

Naive Bayes returns "spam" if 
$$p(y_i = "spam" \mid x_i) > p(y_i = "nd spam" \mid x_i)$$
.

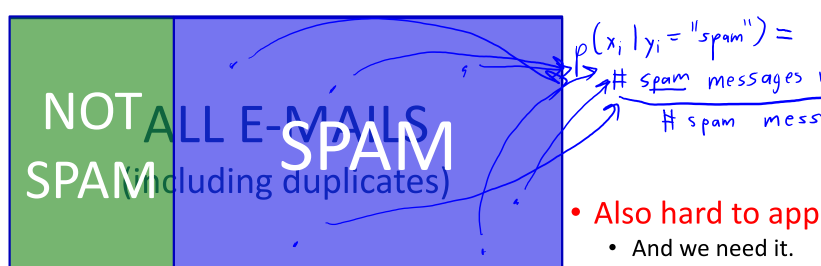
By Bayes rule this means  $p(x_i \mid y_i = "spam")p(y_i = "spam") > p(x_i \mid y_i = "not spam")dy_i = "not spam")dy_i = "not spam" p(x_i)

Multiply both sides by  $p(x_i)$ :$ 

$$p(x_i | y_i = "spam") p(y_i = "spam") > p(x_i | y_i = "not span") dy_i = "not span") dy_i = "not span"$$

$$\rho(y_i = ||spam''||x_i) = \frac{\rho(x_i | y_i = ||spam'')\rho(y_i = ||spam'')}{\rho(x_i)}$$

•  $p(x_i | y_i = "spam")$  is probability that spam has features  $x_i$ .



- Also hard to approximate.

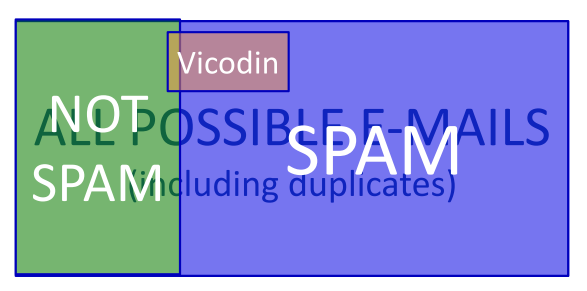
### Naïve Bayes

Naïve Bayes makes a big assumption to make things easier:

- We assume all features  $x_i$  are conditionally independent give label  $y_i$ .
  - Once you know it's spam, probability of "vicodin" doesn't depend on "340".
  - Definitely not true, but sometimes a good approximation.
- And now we only need easy quantities like  $p("vicodin" = 0 | y_i = "spam")$ .

### Naïve Bayes

• p("vicodin" = 1 | "spam" = 1) is probability of seeing "vicodin" in spam.



Easy to estimate:

Again, this is a "maximum likelihood estimate" (MLE). We will cover how to derive this later.

### Summary

- Optimization bias: using a validation set too much overfits.
- Cross-validation: allows better use of data to estimate test error.
- No free lunch theorem: there is no "best" ML model.
- Probabilistic classifiers: try to estimate  $p(y_i \mid x_i)$ .
- Naïve Bayes: simple probabilistic classifier based on counting.
  - Uses conditional independence assumptions to make training practical.
- Next time:
  - A "best" machine learning model as 'n' goes to ∞.

### **Back to Decision Trees**

- Instead of validation set, you can use CV to select tree depth.
- But you can also use these to decide whether to split:
  - Don't split if validation/CV error doesn't improve.
  - Different parts of the tree will have different depths.
- Or fit deep decision tree and use [cross-]validation to prune:
  - Remove leaf nodes that don't improve CV error.
- Popular implementations that have these tricks and others.

### Random Subsamples

- Instead of splitting into k-folds, consider "random subsample" method:
  - At each "round", choose a random set of size 'm'.
    - Train on all examples except these 'm' examples.
    - Compute validation error on these 'm' examples.
- Advantages:
  - Still an unbiased estimator of error.
  - Number of "rounds" does not need to be related to "n".
- Disadvantage:
  - Examples that are sampled more often get more "weight".

### **Cross-Validation Theory**

- Does CV give unbiased estimate of test error?
  - Yes!
    - Since each data point is only used once in validation, expected validation error on each data point is test error.
  - But again, if you use CV to select among models then it is no longer unbiased.
- What about variance of CV?
  - Hard to characterize.
  - CV variance on 'n' data points is worse than with a validation set of size 'n'.
    - But we believe it is close.
- Does cross-validation remove optimization bias?
  - No, but the bias might be smaller since you have more "test" points.

# **Handling Data Sparsity**

- Do we need to store the full bag of words 0/1 variables?
  - No: only need list of non-zero features for each e-mail.

\$	Hi	CPSC	340	Vicodin	Offer	
1	1	0	0	1	0	
0	0	0	0	1	1	
0	1	1	1	0	0	
1	1	0	0	0	1	•••

V5.

Non-Zeroes				
{1,2,5,}				
{5,6,}				
{2,3,4,}				
{1,2,6,}				

Math/model doesn't change, but more efficient storage.

### Proof of No Free Lunch Theorem

- Let's show the "no free lunch" theorem in a simple setting:
  - The  $x^i$  and  $y^i$  are binary, and  $y^i$  being a deterministic function of  $x^i$ .
- With 'd' features, each "learning problem" is a map from each of the  $2^d$  feature combinations to 0 or 1:  $\{0,1\}^d \rightarrow \{0,1\}$

Feature 1	Feature 2	Feature 3
0	0	0
0	0	1
0	1	0

Map 1	Map 2	Map 3	
0	1	0	
0	0	1	
0	0	0	

- Let's pick one of these maps ("learning problems") and:
  - Generate a set training set of 'n' IID samples.
  - Fit model A (convolutional neural network) and model B (naïve Bayes).

### Proof of No Free Lunch Theorem

- Define the "unseen" examples as the  $(2^d n)$  not seen in training.
  - Assuming no repetitions of  $x^i$  values, and  $n < 2^d$ .
  - Generalization error is the average error on these "unseen" examples.
- Suppose that model A got 1% error and model B got 60% error.
  - We want to show model B beats model A on another "learning problem".
- Among our set of "learning problems" find the one where:
  - The labels y<sup>i</sup> agree on all training examples.
  - The labels y<sub>i</sub> disagree on all "unseen" examples.
- On this other "learning problem":
  - Model A gets 99% error and model B gets 40% error.

### Proof of No Free Lunch Theorem

- Further, across all "learning problems" with these 'n' examples:
  - Average generalization error of every model is 50% on unseen examples.
    - It's right on each unseen example in exactly half the learning problems.
  - With 'k' classes, the average error is (k-1)/k (random guessing).
- This is kind of depressing:
  - For general problems, no "machine learning" is better than "predict 0".
- But the proof also reveals the problem with the NFL theorem:
  - Assumes every "learning problem" is equally likely.
  - World encourages patterns like "similar features implies similar labels".