## Supervised Learning Introduction & Framework

CMPUT 261: Introduction to Artificial Intelligence

P&M §7.1-7.3

# Logistics & Assignment #2

- Assignment #2 will be released today See eClass
- Due Tuesday, October 25 at 11:59pm
  - Submissions past the deadline will have late penalty applied
  - Leave yourself some margin for error when submitting!

# Recap: Uncertainty

- We represent uncertainty about the world by **probabilities** 
  - We update our knowledge by conditioning on observations
  - Observations = learning the value of a random variable
- Full, unstructured joint distributions are intractable to reason about
- Conditional independence is a kind of structure that is:
  - 1. widespread
  - 2. easy to reason about
  - 3. allows tractable inference (computing distribution of unobserved variables)
- Belief networks let us compactly represent joint distributions with a lot of conditional independence
  - Variable elimination is an algorithm for efficient inference on belief networks
- Causal models allow us to infer the consequences of interventions

# Supervised Learning, informally

- In the uncertainty section, we took the probability distribution as given
  - Our only problem was to represent and derive distributions
- Question: Where do these probabilities come from?
- Supervised learning is a way to learn probabilities from examples
  - Probability of a target feature (or label) given input features
  - i.e., condition on input features to get probability of target
- Basic idea:
  - Take a bunch of inputs (e.g., images) and "correct" outputs
  - Learn a model that correctly maps inputs to outputs

## Lecture Outline

- Recap & Logistics
- Supervised Learning Problem 2.
- Measuring Prediction Quality З.

### After this lecture, you should be able to:

- define supervised learning task, classification, regression, loss function
- represent categorical target values in multiple ways (indicator variables, indexes)
- define generalization performance lacksquare
- identify an appropriate loss function for different tasks
- explain why a separate test set estimates generalization performance
- define trivial predictors and explain why they are useful  $\bullet$

• define 0/1 error, absolute error, (log-)likelihood loss, mean squared error, worst-case error

# Supervised Learning

### **Definition:** A *supervised learning task* consists of

- A set of input features  $X_1, \ldots, X_n$
- A set of target features  $Y_1, \ldots, Y_k$
- A set of training examples, sampled randomly from some population, for which *both* input and target features are given
- A set of test examples, sampled from the same population, for which only the input features are given

The goal is to predict the values of the target features given the input features; i.e., learn a function h(x) that will map features X to a prediction of Y

- Classification: Y<sub>i</sub> are discrete
- **Regression:**  $Y_i$  are real-valued

# Supervised Learning Examples

- for the main object in the image
  - Input features: Pixel values of the image
  - *Target features:* One feature for each label (e.g., dog, plane, etc.)
- **Precision medicine:** Given examples of symptoms, test results, and treatments, output an 2. estimate of recovery time
  - Input features: symptoms, treatment indicators, test results, demographic information • Target features: recovery time, survival time, etc.
- **Natural language processing:** Given example sentences and labels representing З. "sentiment", output how positive or negative the sentence is
  - Input features: binary indicators for words or characters (\*\*!)
  - Target features: One feature per label (e.g., **positive**, **negative**)

**Computational vision:** Given example images and labels representing objects, output a label

# Regression Example

- Aim is to predict the value of target Y based on features X
- Both X and Y are real-valued
  - Exact values of both targets and features may not have been in the training set
  - $e_8$  is an interpolation problem: X is within the range of the training examples' values
  - $e_9$  is an **extrapolation** problem: X is **outside** the range of the training examples' values

Ex.	X	Y
e1	0.7	1.7
e <sub>2</sub>	1.1	2.4
e <sub>3</sub>	1.3	2.5
<b>e</b> 4	1.9	1.7
<b>e</b> 5	2.6	2.1
<b>e</b> <sub>6</sub>	3.1	2.3
e7	3.9	7

<b>e</b> 8	2.9	?
<b>e</b> 9	5.0	?

# Data Representation

- For real-valued features, we typically just record the feature values
- For **discrete** features, there are multiple options:
  - Binary features: Can code  $\{false, true\}$  as  $\{0,1\}$  or  $\{-1, +1\}$
  - Can record numeric values for each possible value
    - Cardinal values: Differences are meaningful (e.g., 1, 2, 7)
    - Ordinal values: Order is meaningful (e.g., Good, Fair, Poor)
    - Categorical values: Neither differences nor order meaningful (e.g., Red, Green, Blue)
  - Vector of **indicator variables**: One per feature value, exactly one is true (sometimes called a "one-hot" encoding) (e.g., *Red* as (1,0,0), *Green* as (0,1,0), etc.)

### Classification Example: Holiday Preferences

- An agent wants to learn a person's preference for the length of holidays
- Holiday can be for 1,2,3,4,5, or 6 days
- Two possible representations:





<b>Y</b> 1	Y <sub>2</sub>	<b>Y</b> 3	<b>Y</b> 4	<b>Y</b> 5	<b>Y</b> 6
1	0	0	0	0	0
0	0	0	0	0	1
0	0	0	0	0	1
0	1	0	0	0	0
1	0	0	0	0	0

### **Question:**

What are the advantages/ disadvantages of each representation?



## Generalization

- Question: What does it mean for a trained model to perform well?
- We want to be able to make correct predictions on **unseen** data, not just the training examples
  - We are even willing to sacrifice some training accuracy to achieve this
  - We want our learners to generalize: to go beyond the given training examples to classify new examples well
  - **Problem:** We can't measure performance on unobserved examples!
- We can estimate generalization performance by evaluating performance on the test set (Why?)
  - The learning algorithm doesn't have access to the test data, but we do

# Generalization Example

**Example:** Consider binary two classifiers, **P** and **N** 

- P classifies all the **positive examples** from the training data as true, and all others as false
- N classifies all of the **negative examples** from the training data as false, and all others as true

Question: Which classifier generalizes better?

- **Question:** Which classifier performs better on the training data?



- The hypothesis space is the set of possible hypotheses
  - "Training a model" = "Choosing a hypothesis from the hypothesis space based on data"
- A preference for one hypothesis over another is called **bias** 
  - Bias is not a bad thing in this context!
  - Preference for "simple" models is a bias
  - Which bias works best for generalization is an empirical question

### Bias

- Given training data, a hypothesis space, an error measurement, and a bias, learning can be reduced to local search
- Learning searches the hypothesis space trying to find the hypothesis that best fits the data given the bias
  - Search space is prohibitively large (typically infinite)
  - Almost all supervised learning methods are versions of local search

## Learning as Search

# Measuring Prediction Error

- We choose our hypothesis partly by measuring its performance on training data
  - **Question:** What is the other consideration?
- This is usually described as minimizing some quantitative measurement of error (or loss)
  - Question: What might error mean?

### **Definition:**

for which the prediction was not correct:

- $e \in E$
- Not appropriate for **real-valued** target features (**why**?)  $\bullet$
- Does not take into account how wrong the answer is

• e.g., 
$$1 [2 \neq 1] = 1 [6 \neq 1]$$

Most appropriate for **binary** or **categorical** target features  $\bullet$ 

### 0/1 Error

The 0/1 error for a dataset E of examples and hypothesis Y is the number of examples



## Absolute Error

### **Definition:**



- Meaningless for categorical variables
- Takes account of how wrong the predictions are
- Most appropriate for cardinal or possibly ordinal values

The absolute error for a dataset E of examples and hypothesis  $\hat{Y}$  is the sum of absolute distances between the predicted target value and the actual target value:

$$Y(e) - \hat{Y}(e)$$

# Squared Error

### **Definition:**

The squared error (or sum of squares error or mean squared error) for a dataset Eof examples and hypothesis  $\hat{Y}$  is the sum of squared distances between the predicted target value and the actual target value:



- Meaningless for **categorical** variables
- Takes account of how wrong the predictions are
  - Large errors are much more important than small errors
- Most appropriate for **cardinal** values  $\bullet$

$$(e) - \hat{Y}(e) \Big)^2$$

## Worst-Case Error

### **Definition:**

- Meaningless for **categorical** variables  $\bullet$
- Takes account of how wrong the predictions are
  - but only on one example (the one whose prediction is furthest from the true target)
- Most appropriate for cardinal values

- The worst-case error for a dataset E of examples and hypothesis  $\dot{Y}$  is the maximum absolute difference between the predicted target value and the actual target value:
  - $\max_{e \in E} \left| Y(e) \hat{Y}(e) \right|$

## Probabilistic Predictors

- Rather than predicting exactly what a target value will be, many common algorithms predict a **probability distribution** over possible values
  - Especially for **classification** tasks
- scheme:

  - Predicted target values are probabilities that sum to 1

Vectors of indicator variables are the most common data representation for this

• Target features of training examples have a single 1 for the true value

## Probabilistic Predictions Example

### Training examples



Output on test example

X	Ŷcat	Ŷdog	Ŷpanda
	0.5	0.45	0.05

## Likelihood

algorithm

### **Definition:**

The likelihood for a dataset E of examples and hypothesis  $\hat{Y}$  is the probability of independently observing the examples according to the probabilities assigned by the hypothesis:

 $Pr(E \mid \hat{Y})$ 

- This has a clear Bayesian interpretation
- **Numerical stability issues:** product of probabilities shrinks **exponentially**!
  - *Example:* Probability of any sequence of 5000 coin tosses has probability  $2^{-5000}$ !
  - Floating point underflows almost immediately (double-precision floating point can't represent anything smaller than  $2^{-1021}$ )

• For probabilistic predictions, we can use likelihood to measure the performance of a learning

$$) = \prod_{e \in E} \hat{Y}_{Y(e)}(e).$$

# Log-Likelihood

### **Definition:**

The log-likelihood for a dataset E of examples and hypothesis  $\dot{Y}$  is the log-probability of independently observing the examples according to the probabilities assigned by the hypothesis:

 $\log \Pr(E \mid \hat{Y})$ 

- Taking log of the likelihood fixes the underflow issue (**why**?)
- $\bullet$ maximizing likelihood:

$$\left(\Pr(E \mid \hat{Y}_1) > \Pr(E \mid \hat{Y}_2)\right) \iff \left(\log\Pr(E \mid \hat{Y}_1) > \log\Pr(E \mid \hat{Y}_2)\right)$$

$$= \log \prod_{e \in E} \hat{Y}_{Y(e)}(e)$$
$$= \sum_{e \in E} \log \hat{Y}_{Y(e)}(e)$$

The log function grows monotonically, so maximizing log-likelihood is the same thing as

## Trivial Predictors

- **same value** *v* for any example
- **Question:** Why would we every want to think about these? ullet

### • The simplest possible predictor **ignores all input features** and just predicts the

### Optimal Trivial Predictors for Binary Data

- Suppose we are predicting a binary target
- *n*<sub>0</sub> **negative** examples
- *n*<sub>1</sub> **positive** examples
- **Question:** What is the optimal single prediction?

Measure	<b>Optimal Prediction</b>
0/1 error	0 if $n_0 > n_1$ else 1
absolute error	0 if $n_0 > n_1$ else 1
squared error	$\frac{n_1}{n_0 + n_1}$
worst case	$\begin{cases} 0 & \text{if } n_1 = 0 \\ 1 & \text{if } n_0 = 0 \\ 0.5 & \text{otherwise} \end{cases}$
likelihood	$\frac{n_1}{n_0 + n_1}$
log-likelihood	$\frac{n_1}{n_0 + n_1}$

### **Optimal Trivial Predictor Derivations**

0/1 error 0 if  $n_0 > n_1 \text{ else } 1$ 

log-likelihood	$\frac{n_1}{n_0 + n_1}$

 $L(v) = vn_1 + (1 - v)n_0$ 

$$L(v) = n_1 \log v + n_0 \log(1 - v)$$
  
$$\frac{d}{dv}L(v) = 0$$
  
$$0 = \frac{n_1}{v} - \frac{n_0}{1 - v}$$
  
$$\frac{n_0}{1 - v} = \frac{n_1}{v}$$

$$\frac{v}{-v} = \frac{n_1}{n_0} \wedge (0 < v < 1) \implies v = \frac{n_1}{n_0 + n_1}$$

# Summary

- Supervised learning is learning a hypothesis function from training examples
  - Maps from input features to target features
  - Classification: Discrete target features
  - Regression: Real-valued target features
- Preferences among hypotheses are called bias
- Choice of error measurement (loss) is an important design decision
- Different losses have different optimal trivial predictors
  - Trivial predictors are a baseline: your real model better outperform the trivial predictor