

Very convenient!

$$P(Y = 1 | X = \langle X_1, ... X_n \rangle) = \frac{1}{1 + exp(w_0 + \sum_i w_i X_i)}$$
 implies

$$P(Y = 0|X = < X_1, ...X_n >) = \frac{exp(w_0 + \sum_i w_i X_i)}{1 + exp(w_0 + \sum_i w_i X_i)}$$

implies

P(Y = 0|X)
$$P(Y = 1|X) = exp(w_0 + \sum_i w_i X_i)$$
take the In

take the In implies
$$0=\ln 1 \quad \ln \frac{P(Y=0|X)}{P(Y=1|X)} = w_0 + \sum_i w_i X_i \stackrel{?}{\nearrow} 0$$

classification

Logistic regression v. Naïve Bayes



- Consider learning f: X → Y, where
 - □ X is a vector of real-valued features, < X1 ... Xn >
 - ☐ Y is boolean
- Could use a Gaussian Naïve Bayes classifier
 - □ assume all X_i are conditionally independent given Y

 - □ model $P(X_i | Y = y_k)$ as Gaussian $N(\mu_{ik}, \sigma_i)$ □ model P(Y) as $Bernoulli(\theta, 1-\theta)$ [additional for all classes]
- What does that imply about the form of P(Y|X)?

$$P(Y = 1|X = < X_1, ...X_n >) = \frac{1}{1 + exp(w_0 + \sum_i w_i X_i)}$$

Cool!!!!

Derive form for P(Y|X) for continuous
$$X_i$$

By as fact

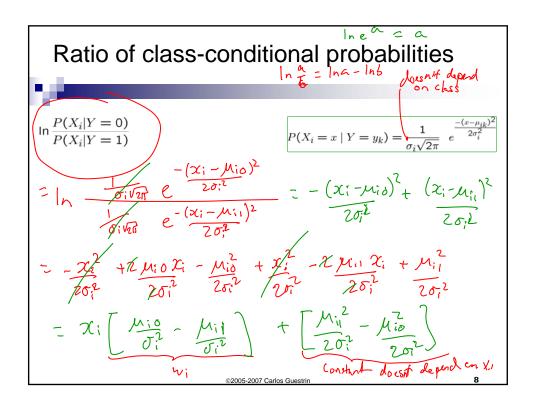
$$P(Y = 1|X) = \frac{P(Y = 1)P(X|Y = 1)}{P(Y = 1)P(X|Y = 1) + P(Y = 0)P(X|Y = 0)}$$

$$= \frac{1}{1 + \frac{P(Y = 0)P(X|Y = 0)}{P(Y = 1)P(X|Y = 1)}}$$

$$= \frac{1}{1 + \exp(\ln \frac{P(Y = 0)P(X|Y = 0)}{P(Y = 1)P(X|Y = 1)})}$$

$$= \frac{1}{1 + \exp(\ln \frac{1 - \theta}{\theta}) + \sum_{i} \ln \frac{P(X_i|Y = 0)}{P(X_i|Y = 1)})}$$
Constant

We show that X_i is X_i in X_i



Derive form for P(Y|X) for continuous X_i

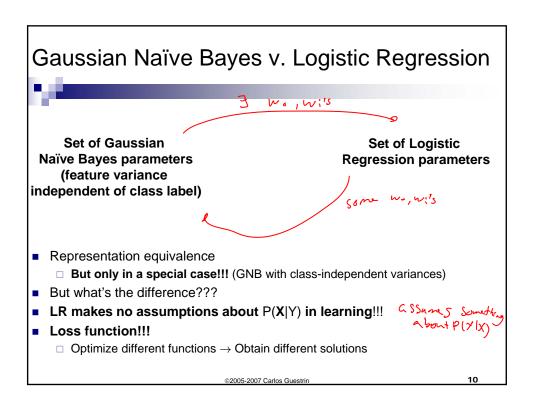
$$P(Y = 1|X) = \frac{P(Y = 1)P(X|Y = 1)}{P(Y = 1)P(X|Y = 1) + P(Y = 0)P(X|Y = 0)}$$

$$= \frac{1}{1 + \exp(\left(\ln\frac{1-\theta}{\theta}\right) + \sum_{i}\ln\frac{P(X_{i}|Y = 0)}{P(X_{i}|Y = 1)})}$$

$$\sum_{i} \frac{\left(\frac{\mu_{i0} - \mu_{i1}}{\sigma_{i}^{2}}X_{i} + \frac{\mu_{i1}^{2} - \mu_{i0}^{2}}{2\sigma_{i}^{2}}\right)}{1 + \exp(w_{0} + \sum_{i=1}^{n} w_{i}X_{i})}$$

$$w_{6} \geq \frac{1}{\theta} + \sum_{i} \frac{\mu_{i}^{2} - \mu_{i0}^{2}}{2\sigma_{i}^{2}}$$

$$w_{6} \geq \frac{1}{\theta} + \sum_{i} \frac{\mu_{i}^{2} - \mu_{i0}^{2}}{2\sigma_{i}^{2}}$$



Logistic regression for more than 2 classes

Logistic regression in more general case, where $Y \in \{Y_1 \dots Y_R\}$: learn R-1 sets of weights $Y \in \{Y_1 \dots Y_R\}$

$$P(Y=1|X) \propto e^{w_{01} + \sum_{i} w_{i}} X_{i}$$

$$P(Y=1|X) \sim e^{w_{01} + \sum_{i} w_{i}} X_{i}$$

$$P(Y=1|X) \sim e^{w_{01} + \sum_{i} w_{i}} X_{i}$$

$$P(Y=1|X) \sim e^{w_{01} + \sum_{i} w_{i}} X_{i}$$

Logistic regression more generally

Logistic regression in more general case, where $Y \in \{Y_1 \dots Y_R\}$: learn R-I sets of weights $\{Y_1 \dots Y_R\}$: $\{(R-I) \cdot (I+I)\}$

especially li takes

for
$$k < R$$

$$P(Y = y_k | X) = \frac{\exp(w_{k0} + \sum_{i=1}^n w_{ki} X_i)}{1 + \sum_{j=1}^{R-1} \exp(w_{j0} + \sum_{i=1}^n w_{ji} X_i)}$$

 $^{\sim}$ fixed for k=R (normalization, so no weights for this class)

((= 1X)**[**

 $P(Y = y_R | X) = \frac{1}{1 + \sum_{j=1}^{R-1} \exp(w_{j0} + \sum_{i=1}^{n} w_{ji} X_i)}$

Features can be discrete or continuous!

Announcements

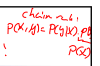


■ Don't forget recitation tomorrow

Still 5:30 pm on Thursdays

And start the homework early

Loss functions: Likelihood v. Conditional Likelihood





Generative (Naïve Bayes) Loss function:

Data likelihood
$$\ln P(\mathcal{D} \mid \mathbf{w}) = \sum_{j=1}^{N} \ln P(\mathbf{x}^{j}, y^{j} \mid \mathbf{w})$$

$$= \sum_{j=1}^{N} \ln P(y^{j} \mid \mathbf{x}^{j}, \mathbf{w}) + \sum_{j=1}^{N} \ln P(\mathbf{x}^{j} \mid \mathbf{w})$$
 Discriminative models cannot compute $\Pr[\mathbf{x}^{j} \mid \mathbf{w}]$

- But, discriminative (logistic regression) loss function:

Conditional Data Likelihood

$$\ln P(\mathcal{D}_Y \mid \mathcal{D}_X, \mathbf{w}) = \sum_{j=1}^{N} \ln P(y^j \mid \mathbf{x}^j, \mathbf{w})$$

 $\hfill\Box$ Doesn't waste effort learning P(X) – focuses on P(Y|X) all that matters for classification

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Expressing Conditional Log Likelihood

$$l(\mathbf{w}) \equiv \sum_{j} \ln P(y^{j} | \mathbf{x}^{j}, \mathbf{w})$$

$$P(Y = 0 | \mathbf{X}, \mathbf{w}) = \frac{1}{1 + exp(w_{0} + \sum_{i} w_{i} X_{i})}$$

$$P(Y = 1 | \mathbf{X}, \mathbf{w}) = \frac{exp(w_{0} + \sum_{i} w_{i} X_{i})}{1 + exp(w_{0} + \sum_{i} w_{i} X_{i})}$$

$$l(\mathbf{w}) = \sum_{j} y^{j} \ln P(y^{j} = 1 | \mathbf{x}^{j}, \mathbf{w}) + (1 - y^{j}) \ln P(y^{j} = 0 | \mathbf{x}^{j}, \mathbf{w})$$

$$\ln P(y = 1 | \mathbf{x}^{j}, \mathbf{w}) = w_{o} + \sum_{i} w_{i} \mathbf{x}_{i} - \ln \left[1 + \ell^{w_{o} + \sum_{i} w_{i}} \mathbf{x}_{i} \right]$$

$$\ln P(y = 0 | \mathbf{x}^{j}, \mathbf{w}) = | \mathbf{x}^{j} - \ln \left[1 + \ell^{w_{o} + \sum_{i} w_{i}} \mathbf{x}_{i} \right]$$

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Maximizing Conditional Log Likelihood

$$l(\mathbf{w}) \equiv \ln \prod_{j} P(y^{j} | \mathbf{x}^{j}, \mathbf{w})$$

$$= \sum_{j} \sqrt{y^{j}(w_{0} + \sum_{i} w_{i}x_{i}^{j})} - \ln(1 + exp(w_{0} + \sum_{i} w_{i}x_{i}^{j}))$$

Good news: $l(\mathbf{w})$ is concave function of $\mathbf{w} \to \mathsf{no}$ locally optimal solutions

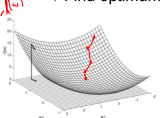
Bad news: no closed-form solution to maximize $l(\mathbf{w})$

Good news: concave functions easy to optimize

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Optimizing concave function – Gradient ascent

■ Conditional likelihood for Logistic Regression is concave → Find optimum with gradient ascent



Gradient:
$$\nabla_{\mathbf{w}} l(\mathbf{w}) = [\frac{\partial l(\mathbf{w})}{\partial w_0}, \dots, \frac{\partial l(\mathbf{w})}{\partial w_n}]'$$

Update rule:
$$\Delta \mathbf{w} = \eta \nabla_{\mathbf{w}} l(\mathbf{w})$$

$$w_i^{(t+1)} \leftarrow w_i^{(t)} + \eta \frac{\partial l(\mathbf{w})}{\partial w_i}$$

- Gradient ascent is simplest of optimization approaches
 - □ e.g., Conjugate gradient ascent much better (see reading)

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Maximize Conditional Log Likelihood:

Gradient ascent
$$\frac{1}{2\pi} | \frac{\ln f(x)}{f(x)} = \frac{\sqrt{2\pi}}{f(x)}$$

$$l(w) = \sum_{j} y^{j} (w_{0} + \sum_{i}^{n} w_{i} x_{i}^{j}) - \ln(1 + \exp(w_{0} + \sum_{i}^{n} w_{i} x_{i}^{j}))$$

$$\frac{1}{2\pi} | \frac{1}{2\pi} | \frac{$$

Gradient Descent for LR

Gradient ascent algorithm: iterate until change < ε

$$\begin{split} w_0^{(t+1)} \leftarrow w_0^{(t)} + \eta \sum_j [y^j - \hat{P}(Y^j = 1 \mid \mathbf{x}^j, \mathbf{w})] \\ \text{For } i = 1 \dots n, \\ w_i^{(t+1)} \leftarrow w_i^{(t)} + \eta \sum_j x_i^j [y^j - \hat{P}(Y^j = 1 \mid \mathbf{x}^j, \mathbf{w})] \end{split}$$

repeat

That's all M(C)LE. How about MAP?



$$p(\mathbf{w} \mid Y, \mathbf{X}) \propto P(Y \mid \mathbf{X}, \mathbf{w}) p(\mathbf{w})$$

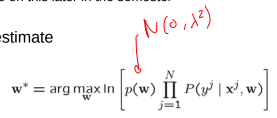


- One common approach is to define priors on w
 - □ Normal distribution, zero mean, identity covariance ⊋(ω)
 - □ "Pushes" parameters towards zero



- Corresponds to Regularization
 - □ Helps avoid very large weights and overfitting
 - ☐ More on this later in the semester

MAP estimate



M(C)AP as Regularization

$$\ln \left[p(\mathbf{w}) \prod_{j=1}^{N} P(y^{j} \mid \mathbf{x}^{j}, \mathbf{w}) \right] \qquad p(\mathbf{w}) = \prod_{i} \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_{i}^{2}}{2\kappa^{2}}}$$

$$= \ln p(\omega) + \ln \prod_{j>i} p(y^{j} \mid \chi^{j}, \omega)$$

$$= \lim_{i \to \infty} p(\omega) + \lim_{j \to i} p(y^{j} \mid \chi^{j}, \omega)$$

$$= \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) = \lim_{i \to \infty} \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_{i}^{2}}{2\kappa^{2}}}$$

$$= \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) = \lim_{i \to \infty} \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_{i}^{2}}{2\kappa^{2}}}$$

$$= \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) + \lim_{i \to \infty} p(\omega) = \lim_{i \to \infty} \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_{i}^{2}}{2\kappa^{2}}}$$

$$= \lim_{i \to \infty} p(\omega) + \lim_{i \to$$

Penalizes high weights, also applicable in linear regression

Gradient of M(C)AP

$$\frac{\partial}{\partial w_i} \ln \left[p(\mathbf{w}) \prod_{j=1}^N P(y^j \mid \mathbf{x}^j, \mathbf{w}) \right] \qquad p(\mathbf{w}) = \prod_i \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_i^2}{2\kappa^2}}$$

$$= \frac{\partial}{\partial w_i} - \sum_i \frac{w_i^2}{2\kappa^2} + \frac{\partial}{\partial w_i} \mathcal{N}(\omega)$$

$$p(\mathbf{w}) = \prod_{i} \frac{1}{\kappa \sqrt{2\pi}} e^{\frac{-w_i^2}{2\kappa^2}}$$

MLE vs MAP

Maximum conditional likelihood estimate

$$\begin{aligned} \mathbf{w}^* &= \arg\max_{\mathbf{w}} \ln \left[\prod_{j=1}^N P(y^j \mid \mathbf{x}^j, \mathbf{w}) \right] \\ w_i^{(t+1)} &\leftarrow w_i^{(t)} + \eta \sum_j x_i^j [y^j - \hat{P}(Y^j = 1 \mid \mathbf{x}^j, \mathbf{w})] \end{aligned}$$

Maximum conditional a posteriori estimate

$$\mathbf{w}^* = \arg\max_{\mathbf{w}} \ln\left[p(\mathbf{w}) \prod_{j=1}^{N} P(y^j \mid \mathbf{x}^j, \mathbf{w})\right]$$

$$\begin{bmatrix} w_i^{(t+1)} \leftarrow w_i^{(t)} + \eta \left\{ -\lambda w_i^{(t)} + \sum_j x_i^j [y^j - \hat{P}(Y^j = 1 \mid \mathbf{x}^j, \mathbf{w})] \right\} \end{bmatrix}$$

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Naïve Bayes vs Logistic Regression

Consider Y boolean, X_i continuous, $X=< X_1 ... X_n>$

Number of parameters:

- NB: 4n +1
- LR: n+1

Estimation method:

- NB parameter estimates are uncoupled
- LR parameter estimates are coupled

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G. Naïve Bayes vs. Logistic Regression 1 [Ng & Jordan, 2002] Generative and Discriminative classifiers

- Asymptotic comparison (# training examples → infinity)
 - □ when model correct
 - GNB, LR produce identical classifiers
 - when model incorrect
 - LR is less biased does not assume conditional independence

 □ therefore LR expected to outperform GNB

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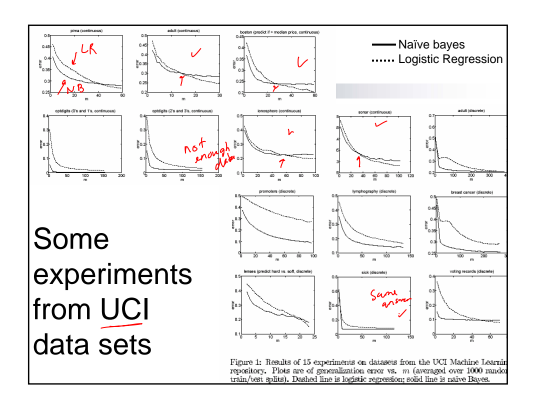
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G. Naïve Bayes vs. Logistic Regression 2

[Ng & Jordan, 2002]

- Generative and Discriminative classifiers
- Non-asymptotic analysis
 - □ convergence rate of parameter estimates, n = # of attributes in X
 - Size of training data to get close to infinite data solution
 - GNB needs O(log n) samples
 - LR needs O(n) samples
 - GNB converges more quickly to its (perhaps less helpful) asymptotic estimates

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What you should know about Logistic Regression (LR)

- Gaussian Naïve Bayes with class-independent variances representationally equivalent to LR
 - □ Solution differs because of objective (loss) function
- In general, NB and LR make different assumptions
 - \square NB: Features independent given class \rightarrow assumption on P(X|Y)
 - \square LR: Functional form of P(Y|X), no assumption on P(X|Y)
- LR is a linear classifier
 - □ decision rule is a hyperplane
- LR optimized by conditional likelihood
 - □ no closed-form solution
 - $\hfill\Box$ concave \to global optimum with gradient ascent
 - ☐ Maximum conditional a posteriori corresponds to regularization
- Convergence rates
 - □ GNB (usually) needs less data
 - □ LR (usually) gets to better solutions in the limit

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Linear separability



- A dataset is linearly separable iff ∃ a separating hyperplane:
 - □ ∃ w, such that:
 - $\mathbf{w}_0 + \sum_i \mathbf{w}_i \mathbf{x}_i > 0$; if $\mathbf{x} = \{\mathbf{x}_1, \dots, \mathbf{x}_n\}$ is a positive example
 - $w_0 + \sum_i w_i x_i < 0$; if $\mathbf{x} = \{x_1, ..., x_n\}$ is a negative example

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Not linearly separable data



Some datasets are not linearly separable!

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Addressing non-linearly separable data – Option 1, non-linear features

- Choose non-linear features, e.g.,
 - □ Typical linear features: $w_0 + \sum_i w_i x_i$
 - □ Example of non-linear features:
 - Degree 2 polynomials, $\mathbf{w_0} + \sum_i \mathbf{w_i} \mathbf{x_i} + \sum_{ij} \mathbf{w_{ij}} \mathbf{x_i} \mathbf{x_j}$
- Classifier h_w(x) still linear in parameters w
 - □ Usually easy to learn (closed-form or convex/concave optimization)
 - □ Data is linearly separable in higher dimensional spaces
 - □ More discussion later this semester

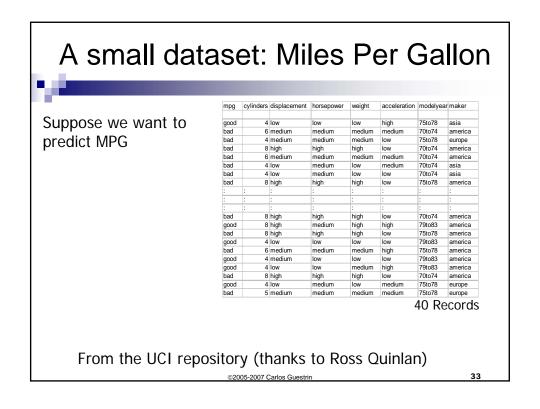
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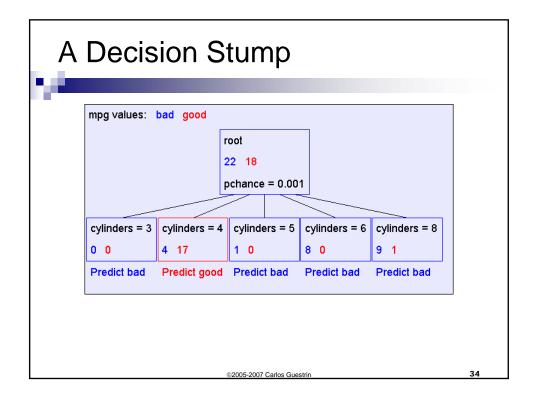
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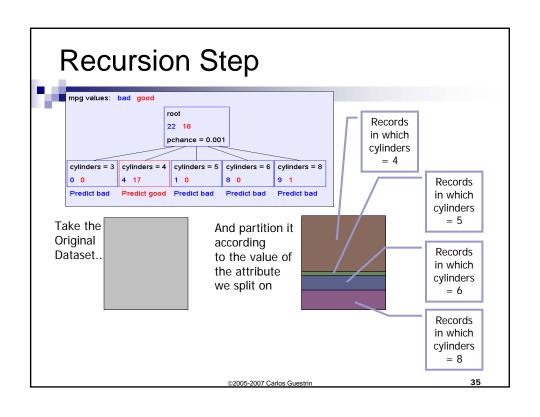
Addressing non-linearly separable data – Option 2, non-linear classifier

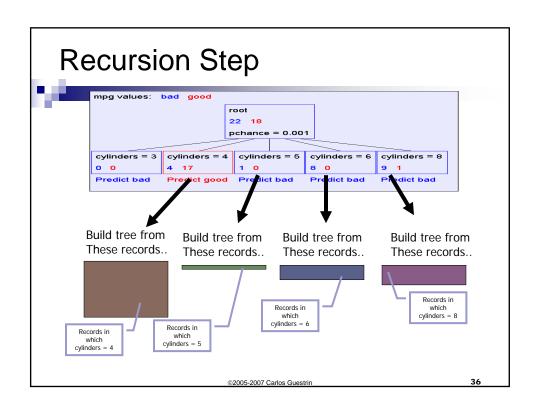
- Choose a classifier h_w(x) that is non-linear in parameters w, e.g.,
 - □ Decision trees, neural networks, nearest neighbor,...
- More general than linear classifiers
- But, can often be harder to learn (non-convex/concave optimization required)
- But, but, often very useful
- (BTW. Later this semester, we'll see that these options are not that different)

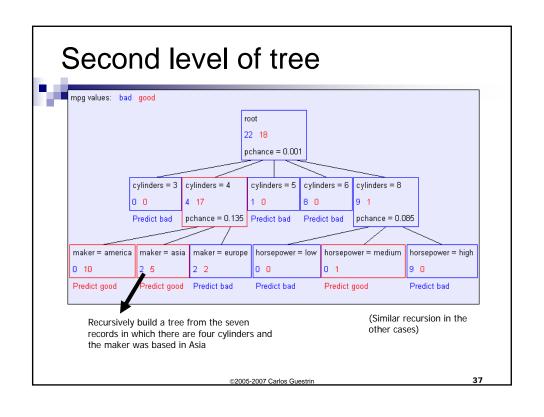
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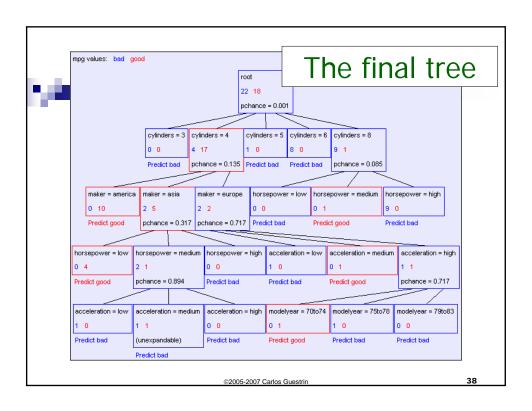


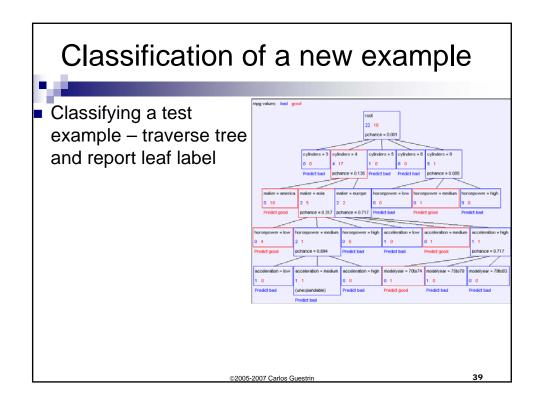














- Many trees can represent the same concept
- But, not all trees will have the same size!
 - \square e.g., $\varphi = A \wedge B \vee \neg A \wedge C \ \ \mbox{((A and B) or (not A and C))}$

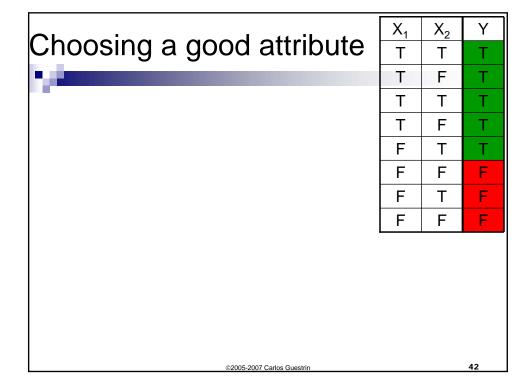
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Learning decision trees is hard!!!

- Learning the simplest (smallest) decision tree is an NP-complete problem [Hyafil & Rivest '76]
- Resort to a greedy heuristic:
 - □ Start from empty decision tree
 - ☐ Split on **next best attribute (feature)**
 - □ Recurse

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Measuring uncertainty



- Good split if we are more certain about classification after split
 - □ Deterministic good (all true or all false)
 - □ Uniform distribution bad

Y=A) = 1/2 $P(Y=B) = 1/4$ $P(Y=C) = 1/8$ $P(Y=D) = 1$	/8
---	----

P(Y=A) = 1/4 $P(Y=B) = 1/4$ $P(Y=C) = 1/4$ $P(Y=D) = 1/4$

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Entropy

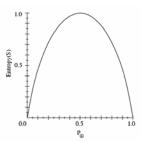


Entropy H(X) of a random variable Y

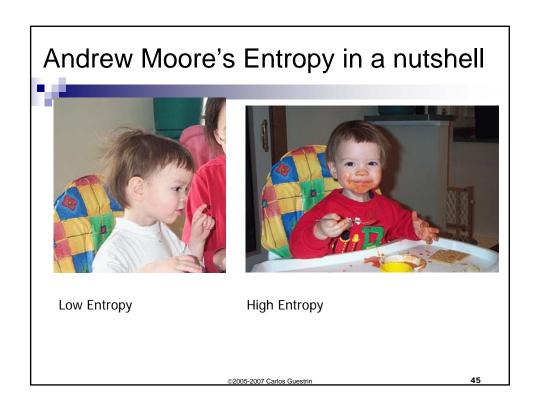
$$H(Y) = -\sum_{i=1}^{k} P(Y = y_i) \log_2 P(Y = y_i)$$

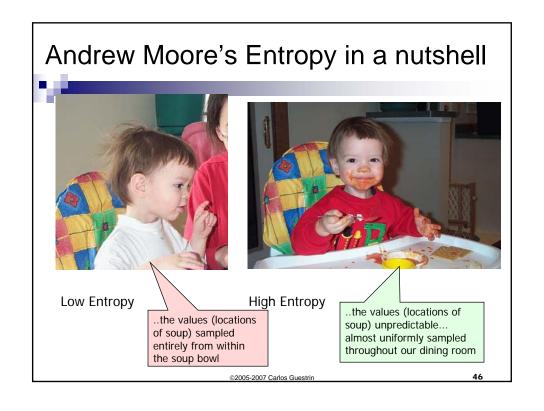
More uncertainty, more entropy!

Information Theory interpretation: H(Y) is the expected number of bits needed to encode a randomly drawn value of Y (under most efficient code)



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Information gain



- Advantage of attribute decrease in uncertainty
 - □ Entropy of Y before you split
 - □ Entropy after split
 - Weight by probability of following each branch, i.e., normalized number of records

$$H(Y \mid X) = -\sum_{j=1}^{v} P(X = x_j) \sum_{i=1}^{k} P(Y = y_i \mid X = x_j) \log_2 P(Y = y_i \mid X = x_j)$$

■ Information gain is difference $IC(X) = H(Y) - H(Y \mid X)$

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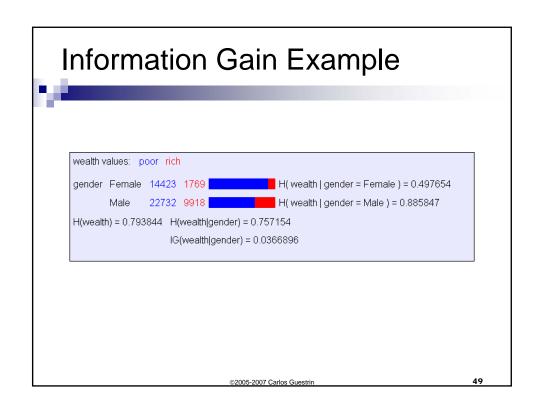
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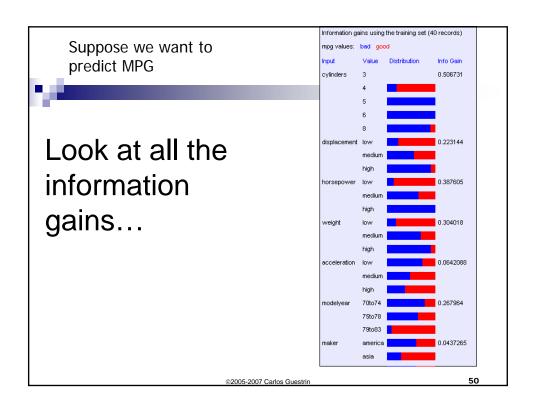
Learning decision trees

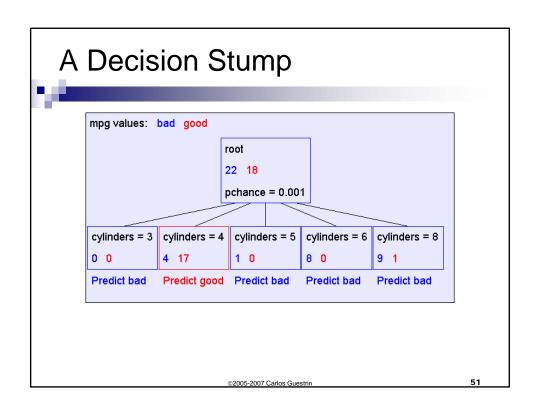


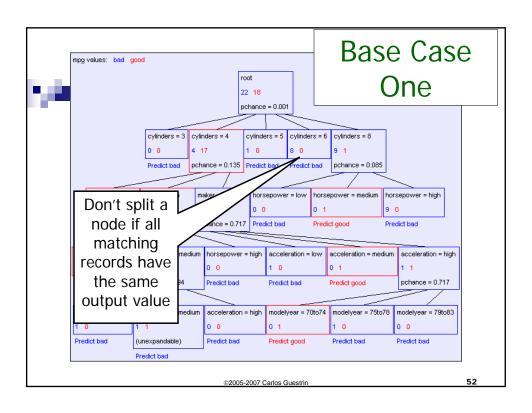
- Start from empty decision tree
- Split on next best attribute (feature)
 - $\hfill\square$ Use, for example, information gain to select attribute
 - \square Split on $\max_{i} IG(X_i) = \arg\max_{i} H(Y) H(Y \mid X_i)$
- Recurse

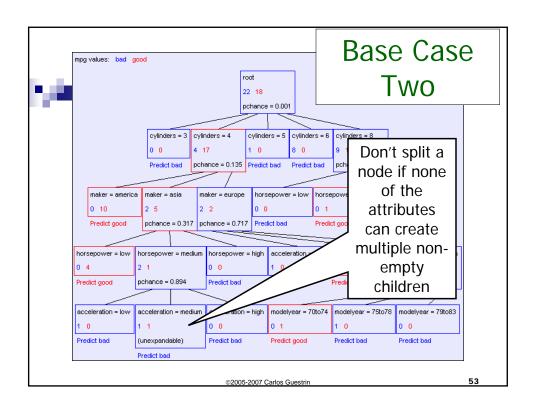
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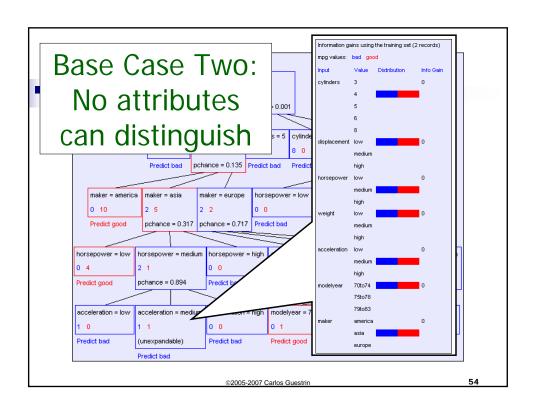












Base Cases



- Base Case One: If all records in current data subset have the same output then don't recurse
- Base Case Two: If all records have exactly the same set of input attributes then don't recurse

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Base Cases: An idea



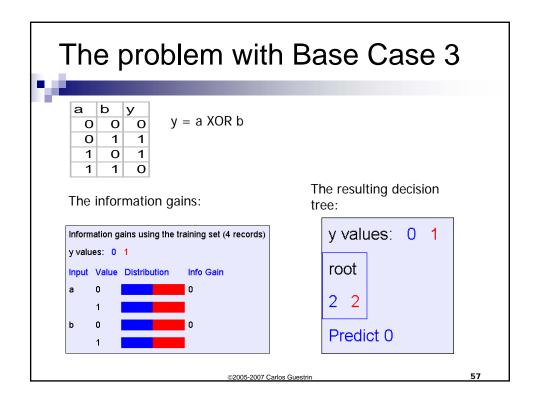
- Base Case One: If all records in current data subset have the same output then don't recurse
- Base Case Two: If all records have exactly the same set of input attributes then don't recurse

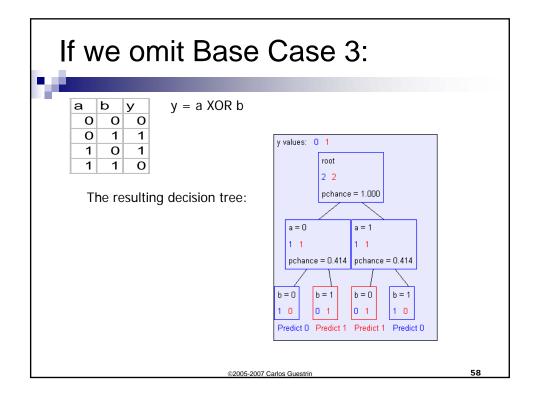
Proposed Base Case 3:

If all attributes have zero information gain then don't recurse

• Is this a good idea?

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Basic Decision Tree Building Summarized



BuildTree(DataSet, Output)

- If all output values are the same in DataSet, return a leaf node that says "predict this unique output"
- If all input values are the same, return a leaf node that says "predict the majority output"
- Else find attribute X with highest Info Gain
- Suppose X has n_X distinct values (i.e. X has arity n_X).
 - \Box Create and return a non-leaf node with n_X children.
 - ☐ The *i*th child should be built by calling BuildTree(*DS_i*, *Output*)

Where DS_i built consists of all those records in DataSet for which X = ith distinct value of X.

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Real-Valued inputs



What should we do if some of the inputs are real-valued?

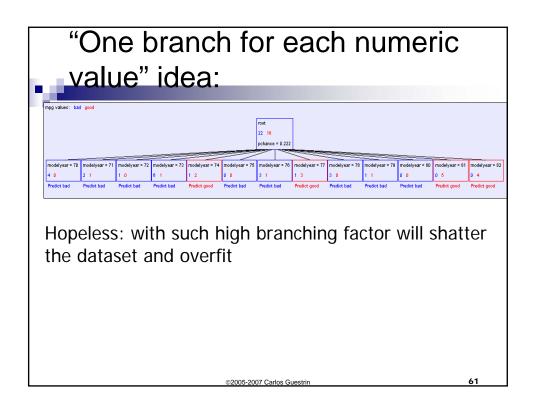
mpg	cylinders	displacemen	norsepower	weignt	acceleration	modelyear	maker
good	4	97	75	2265	18.2	77	asia
bad	6	199	90	2648	15	70	america
bad	4	121	110	2600	12.8	77	europe
bad	8	350	175	4100	13	73	america
bad	6	198	95	3102	16.5	74	america
bad	4	108	94	2379	16.5	73	asia
bad	4	113	95	2228	14	71	asia
bad	8	302	139	3570	12.8	78	america
:			:	:	:	:	
:	:	:	:	:	:	:	:
:			:	:	:	:	
good	4	120	79	2625	18.6	82	america
bad	8	455	225	4425	10	70	america
good	4	107	86	2464	15.5	76	europe
bad	5	131	103	2830	15.9	78	europe

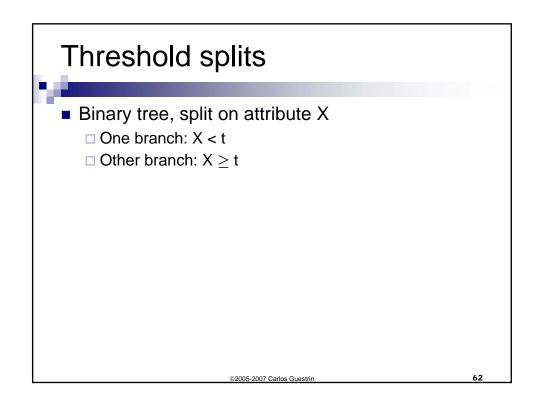
Infinite number of possible split values!!!

Finite dataset, only finite number of relevant splits!

Idea One: Branch on each possible real value

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Choosing threshold split



- Binary tree, split on attribute X
 - □ One branch: X < t
 - □ Other branch: X > t
- Search through possible values of t
 - □ Seems hard!!!
- But only finite number of *t*'s are important
 - □ Sort data according to X into $\{x_1,...,x_m\}$
 - \Box Consider split points of the form $x_i + (x_{i+1} x_i)/2$

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A better idea: thresholded splits

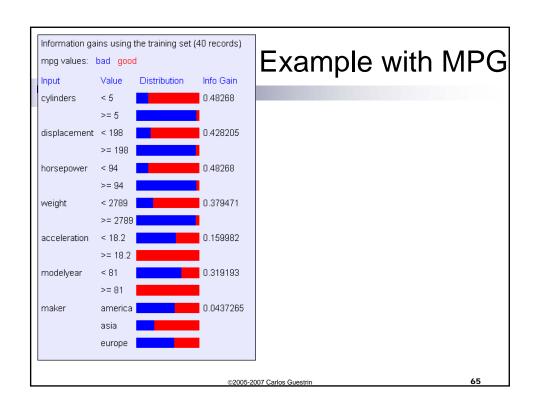


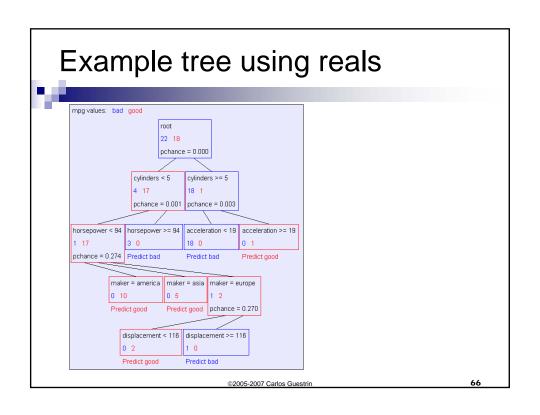
- Suppose X is real valued
- Define *IG*(*Y*|*X:t*) as *H*(*Y*) *H*(*Y*|*X:t*)
- Define *H*(*Y*|*X:t*) =

$$H(Y|X < t) P(X < t) + H(Y|X >= t) P(X >= t)$$

- *IG*(*Y*|*X*:*t*) is the information gain for predicting Y if all you know is whether X is greater than or less than *t*
- Then define $IG^*(Y|X) = max_t IG(Y|X:t)$
- For each real-valued attribute, use $IG^*(Y|X)$ for assessing its suitability as a split

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What you need to know about decision trees

- Decision trees are one of the most popular data mining tools
 - □ Easy to understand
 - ☐ Easy to implement
 - □ Easy to use
 - □ Computationally cheap (to solve heuristically)
- Information gain to select attributes (ID3, C4.5,...)
- Presented for classification, can be used for regression and density estimation too
- It's possible to get in trouble with overfitting (more next lecture)

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Acknowledgements



- Some of the material in the presentation is courtesy of Tom Mitchell, and of Andrew Moore, from his excellent collection of ML tutorials:
 - □ http://www.cs.cmu.edu/~awm/tutorials

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