# Classification and Regression Trees

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#### **Course outline**

#### 1. Motivating examples

- (a) Least squares regression: impact of air pollution on house prices
- (b) Poisson regression: defects in soldering circuit boards
- (c) Multiresponse data: interactions of variables in production of concrete
- (d) Longitudinal data: hourly wages of high-school dropouts
- (e) Censored data and differential treatment effects: breast cancer survival
- (f) Simple classification: Fisher's iris data
- (g) Classification with unequal costs: attitudes towards mammography
- (h) Unbalanced classes: characterizing dissatisfied credit card holders

#### 2. Classification tree algorithms

- (a) THAID (Messenger and Mandell, 1972), CART (Breiman et al., 1984), RPART (Therneau and Atkinson, 2013, 2012)
- (b) FACT (Loh and Vanichsetakul, 1988), QUEST (Loh and Shih, 1997), CRUISE (Kim and Loh, 2001, 2003), GUIDE (Loh, 2009)

- (c) C4.5 (Quinlan, 1993), CHAID (Kass, 1980), CTREE (Hothorn et al., 2006)
- (d) More examples: peptide binding; fish identification; car prediction
- (e) Missing values, selection bias, accuracy, speed, and tree complexity

#### 3. Regression tree algorithms

- (a) Piecewise constant least squares models: AID (Morgan and Sonquist, 1963), CART, RPART, GUIDE (Loh, 2002)
- (b) Piecewise linear least squares, quantile regression, subgroup identification of differential treatment effects, and longitudinal data effects: GUIDE (Loh and Zheng, 2013)
- (c) Others: M5 (Quinlan, 1992), MOB (Zeileis et al., 2008)
- (d) More examples: college tuition; primary biliary cirrhosis of the liver; progression of CD4 counts in AIDS
- (e) Missing values, selection bias, accuracy, speed, and tree complexity

#### 4. Conclusion

#### **Free software**

- C4.5—www.rulequest.com/Personal/c4.5r8.tar.gz; see also www.cs.uregina.ca/~dbd/cs831/notes/ml/dtrees/c4.5/tutorial.html
- CART, C4.5, M5, etc.—www.cs.waikato.ac.nz/~ml/weka/
- CRUISE, GUIDE, LOTUS, QUEST—www.stat.wisc.edu/~loh/
- RPART, CTREE, MOB, PARTY, RandomForest —cran.us.r-project.org/
- LATEX (text processing package)—http://www.ctan.org/ CRUISE, GUIDE, LOTUS, and QUEST produce LATEX tree diagrams

# Some review papers

- 1. Lemon et al. (2003), Classification and regression tree analysis in public health: methodological review and comparison with logistic regression, *Annals of Behavioral Medicine*
- 2. Loh (2008a), Classification and regression tree methods, *Encyclopedia of Statistics in Quality and Reliability*
- 3. Merkle and Shaffer (2011), Binary recursive partitioning: background, methods, and application to psychology, *British Journal of Mathematical and Statistical Psychology*
- 4. Loh (2011), Classification and regression trees, *Wiley Interdisciplinary Reviews: Data Mining and Knowledge Discovery*
- 5. Loh (2014), Fifty years of classification and regression trees (with discussion), *International Statistical Review*

# Linear regression: 1970 Boston housing data (Harrison and Rubinfeld, 1978; Belsley et al., 1980)

Var	Definition	Var	Definition
ID	census tract number	TOWN	township (92 values)
MEDV	median value in \$1000	AGE	% built before 1940
CRIM	per capita crime rate	DIS	distance to employment centers
ZN	% zoned for lots $>$ 25K sq.ft.	RAD	accessibility to radial highways
INDUS	% nonretail business	TAX	property tax rate per \$10000
CHAS	1 on Charles River, 0 else	PT	pupil/teacher ratio
NOX	nitrogen oxide conc. (p.p. $10^9$ )	В	$(\% black - 63)^2/10$
RM	average number of rooms	LSTAT	% lower-status population

Data: 506 observations (census tracts) in the greater Boston area

Objective: To examine the impact of air pollution on house price

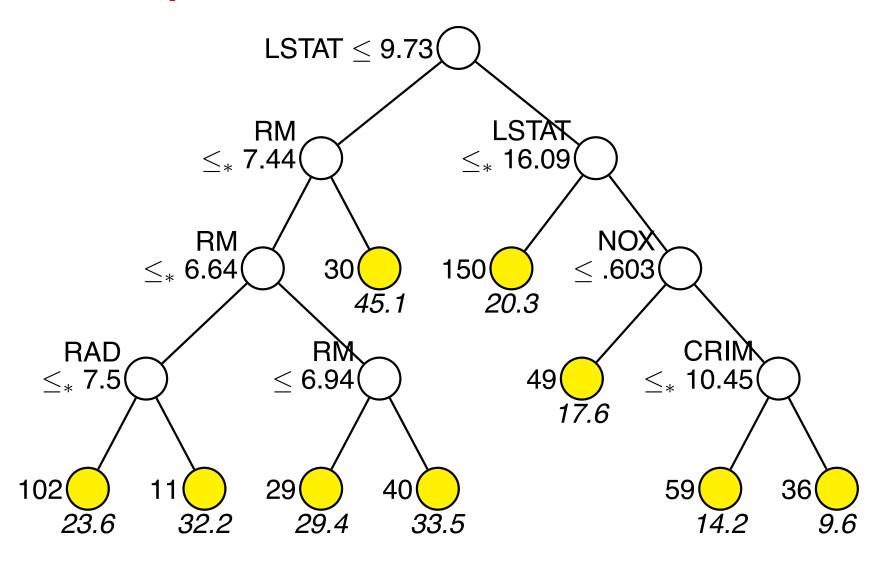
# Harrison & Rubinfeld model for log(MEDV)

X	eta	t	ho	X	$\beta$	t	ho
Constant	4.6	30.0		AGE	7.1E-5	0.1	-0.5
CRIM	-1.2E-2	-9.6	-0.5	log(DIS)	-2.0E-1	-6.0	0.4
ZN	9.2E-5	0.2	0.4	log(RAD)	9.0E-2	4.7	-0.4
INDUS	1.8E-4	0.1	-0.5	TAX	-4.2E-4	-3.5	-0.6
CHAS	9.2E-2	2.8	0.2	PT	-3.0E-2	-6.0	-0.5
$\mathtt{NOX}^2$	-6.4E-1	-5.7	-0.5	В	3.6E-4	3.6	0.4
${\tt RM}^2$	6.3E-3	4.8	0.6	log(LSTAT)	-3.7E-1	-15.2	-0.8

 $\beta$  = coefficient, t = t-statistic,  $\rho$  = corr(X, Y)

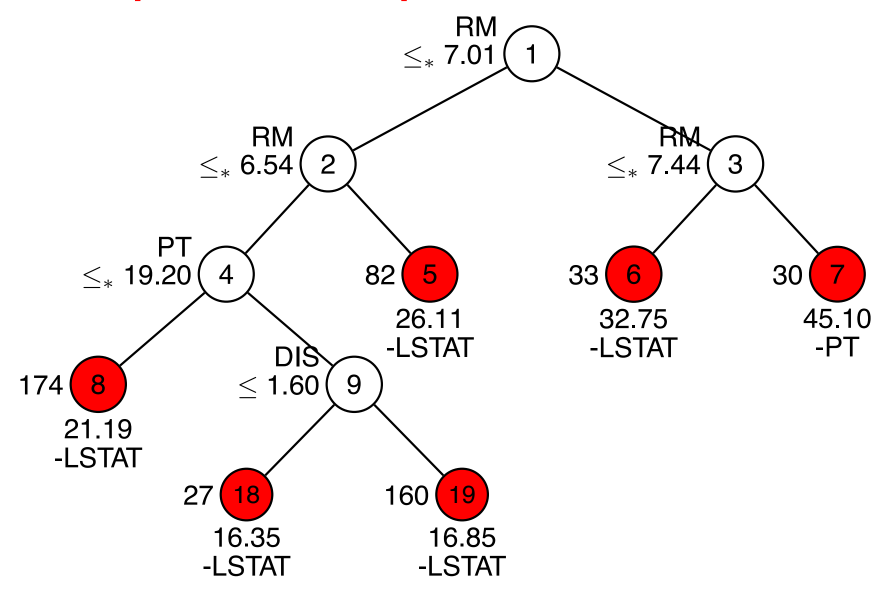
### What can we conclude from this model?

## **GUIDE** piecewise constant model for MEDV

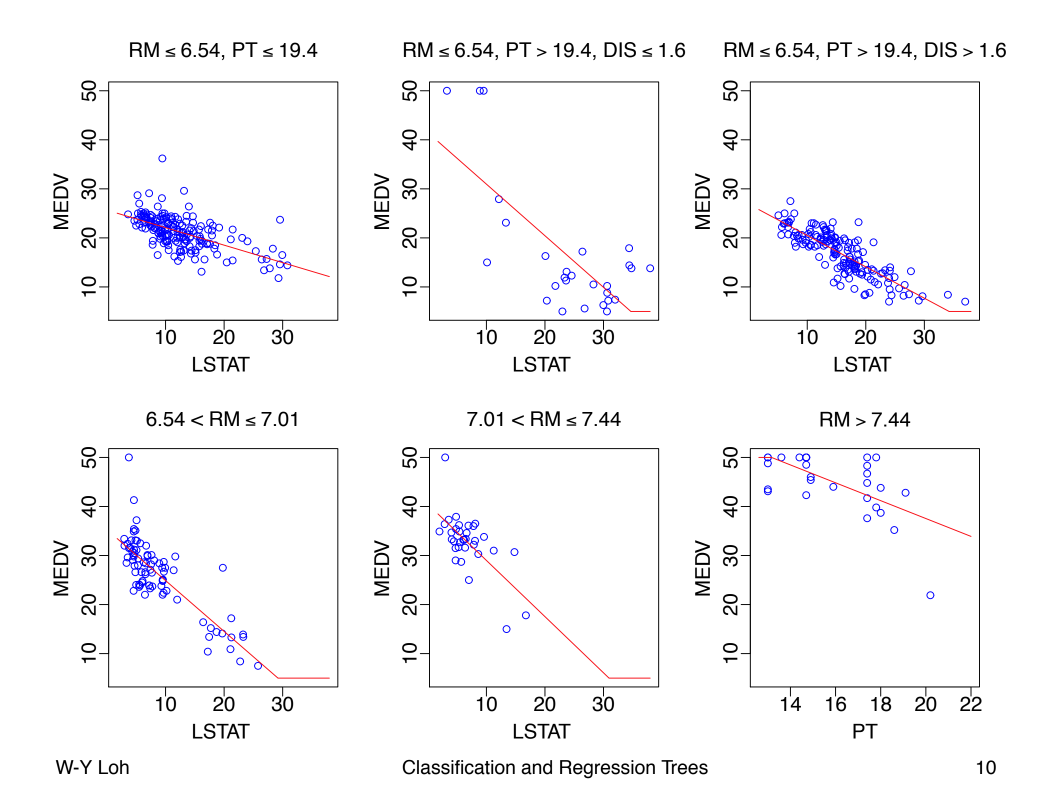


Sample means and sample sizes below and beside nodes. At each intermediate node, a case goes left if and only if the condition is true. Symbol " $\leq_*$ " means " $\leq$  or missing."

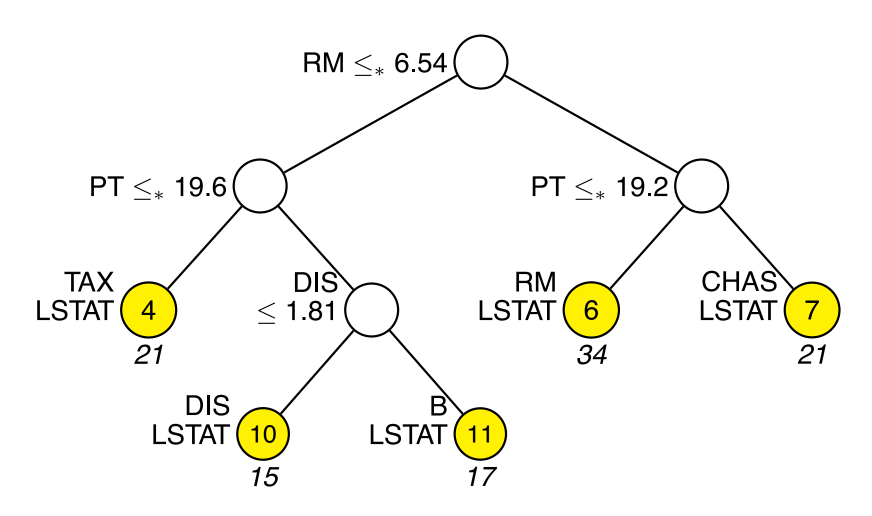
# **GUIDE** piecewise simple linear model for MEDV



Mean MEDV and signed linear predictor beneath each node

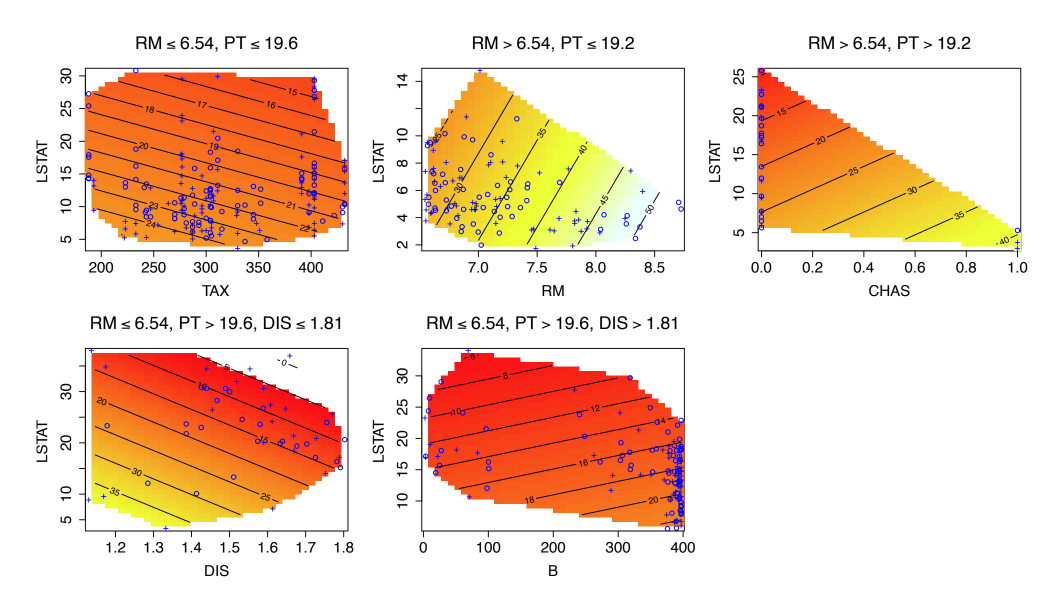


# **GUIDE** piecewise two-variable model for MEDV

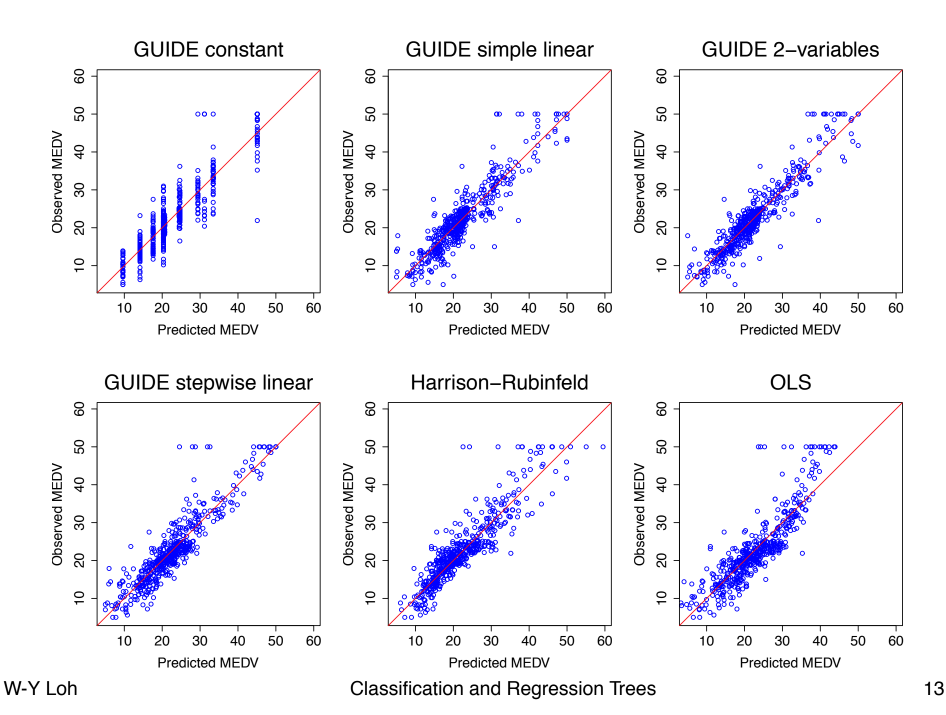


Mean MEDV beneath each node

### Data and fits in GUIDE two-variable model



# **Comparison of models**



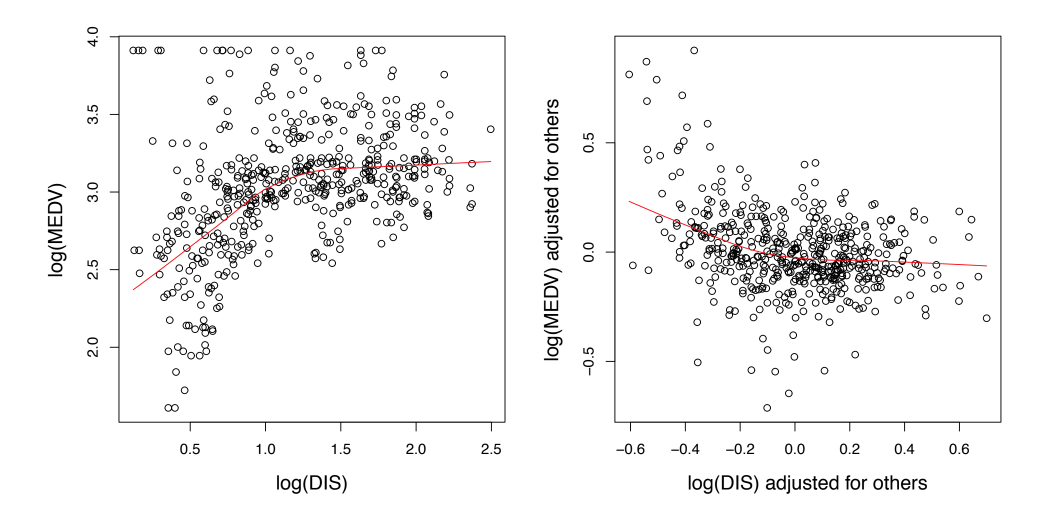
# Difficulties in interpreting regression coefficients: Harrison & Rubinfeld model for log(MEDV)

X	eta	t	ho	X	eta	t	ho
Constant	4.6	30.0		AGE	7.1E-5	0.1	-0.5
CRIM	-1.2E-2	-9.6	-0.5	log(DIS)	-2.0E-1	-6.0	0.4
ZN	9.2E-5	0.2	0.4	log(RAD)	9.0E-2	4.7	-0.4
INDUS	1.8E-4	0.1	-0.5	TAX	-4.2E-4	-3.5	-0.6
CHAS	9.2E-2	2.8	0.2	PT	-3.0E-2	-6.0	-0.5
$\mathtt{NOX}^2$	-6.4E-1	-5.7	-0.5	В	3.6E-4	3.6	0.4
${\tt RM}^2$	6.3E-3	4.8	0.6	log(LSTAT)	-3.7E-1	-15.2	-0.8

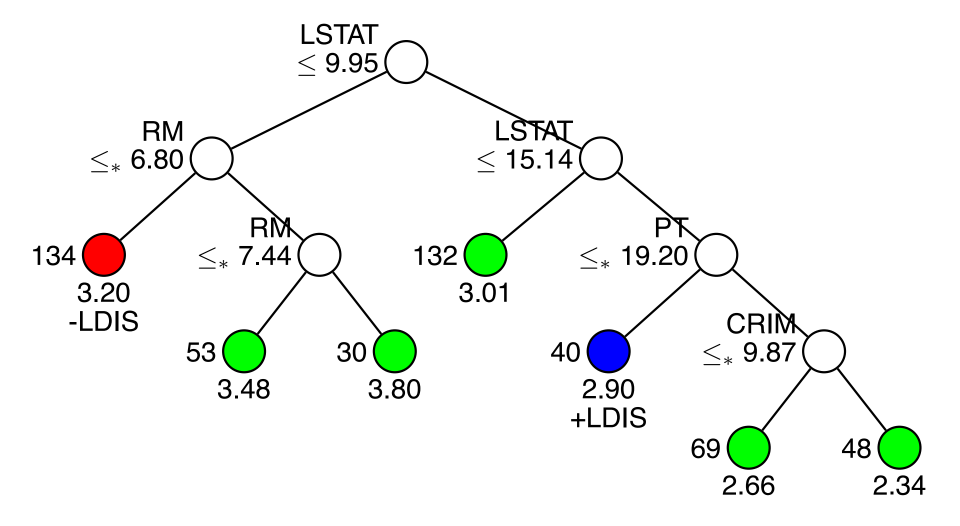
 $\beta$  = coefficient, t = t-statistic,  $\rho$  = corr(X, Y)

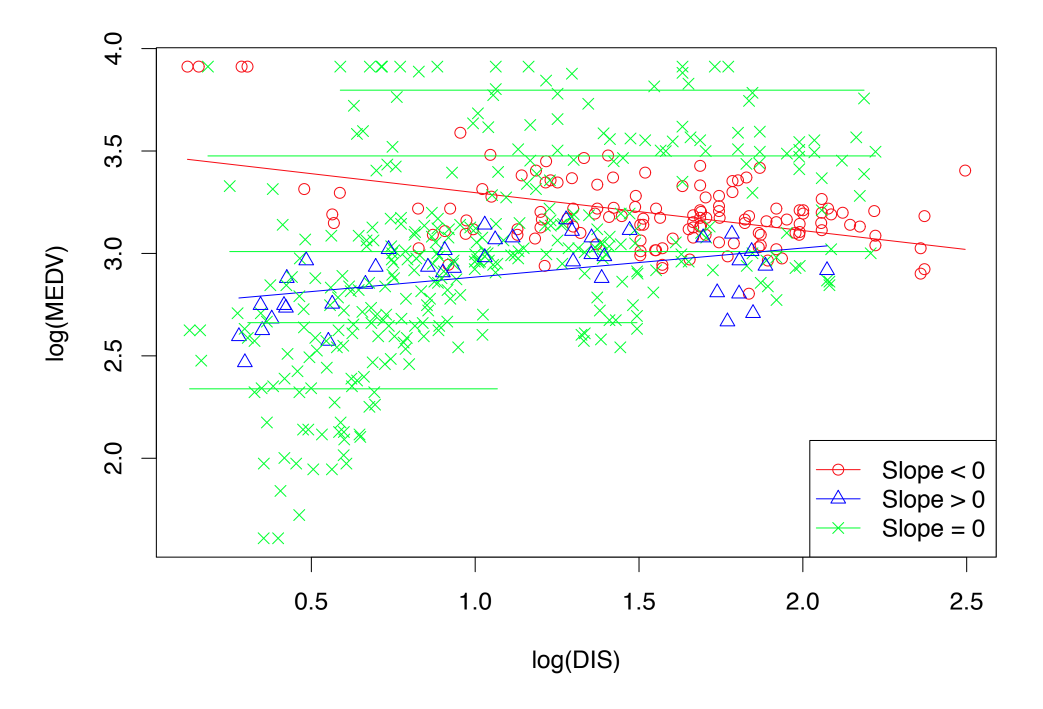
Why do  $\beta$  and  $\rho$  have opposite signs for  $\log(\text{DIS})$  and  $\log(\text{RAD})$ ?

# log(MEDV) vs. log(DIS)

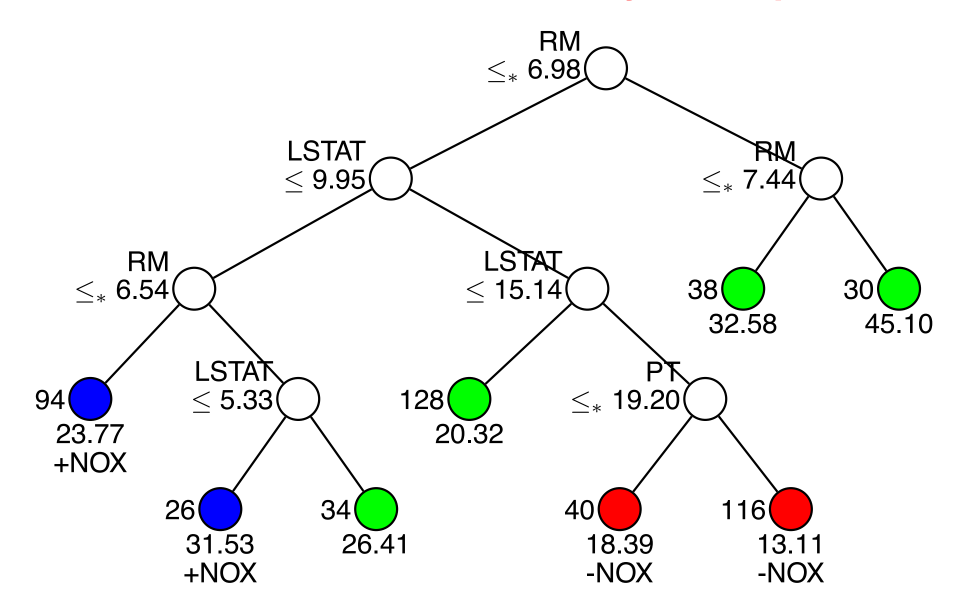


# Model for log(MEDV) with log(DIS) as linear predictor

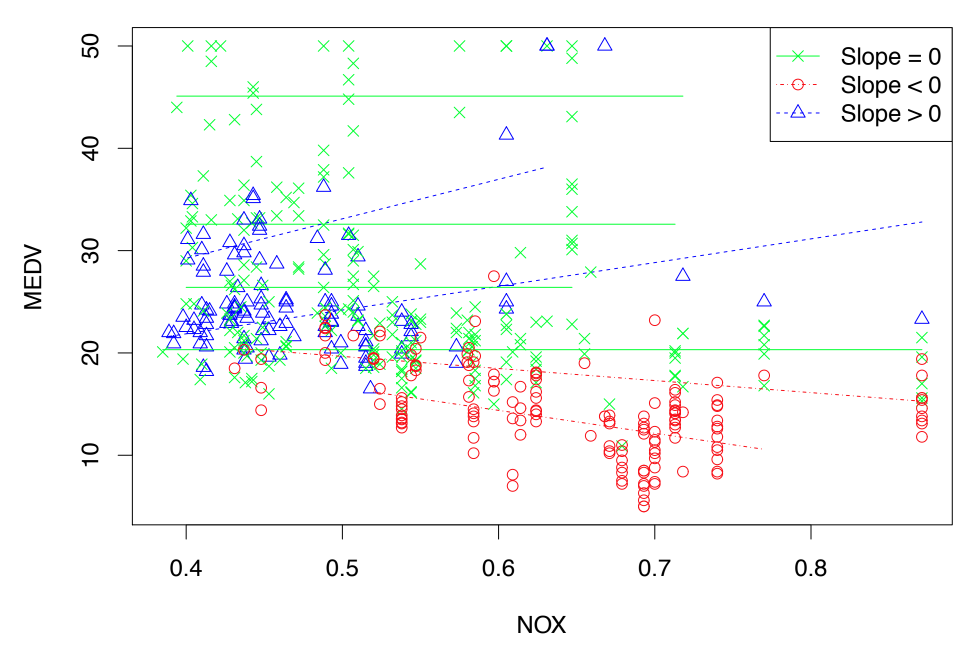




# Model for MEDV with NOX as only linear predictor



### **MEDV vs NOX**



# Poisson regression:

# Unreplicated $3 \times 2 \times 4 \times 10 \times 3$ soldering experiment

Opening: Amount of clearance around a mounting pad (small, medium, large)

**Solder:** Amount of solder (thin, thick)

Mask: Type and thickness of solder mask (A1.5, A3, B3, B6)

Pad: Shape and size of mounting pad (D4, D6, D7, L4, L6, L7, L8, L9, W4, W9)

**Panel:** Each board is divided into three panels (1, 2, 3)

Response: Number of solder skips (0–48)

Ref: Comizzoli et al. (1990), Chambers and Hastie (1992)

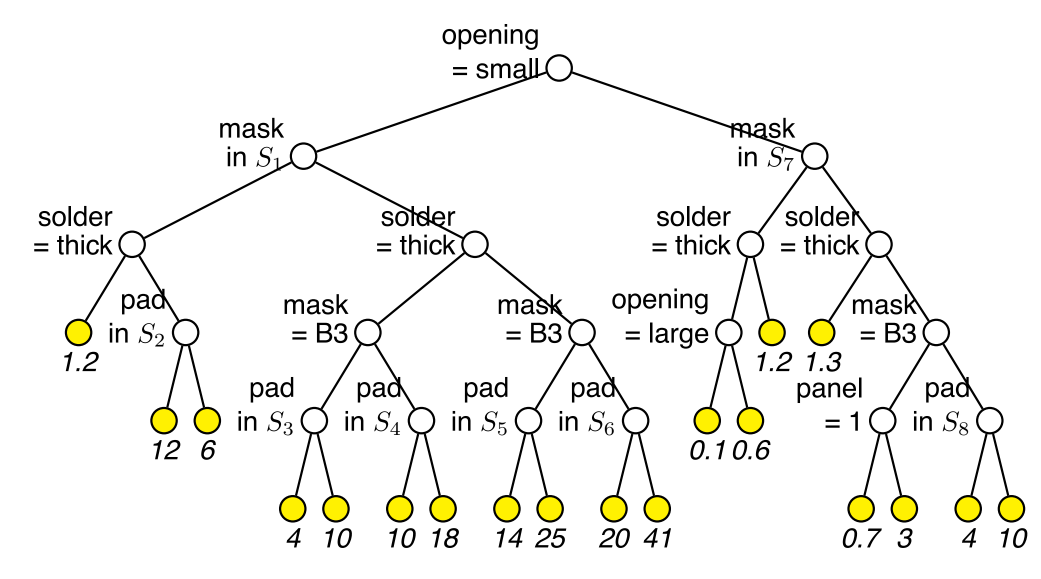
# Full 2nd-degree Poisson loglinear model

Term	df	Deviance	Р	Term	df	Deviance	Р
open	2	2524.6	0.000	open:pad	18	47.4	0.000
solder	1	937.0	0.000	open:panel	4	11.2	0.024
mask	3	1653.1	0.000	solder:pad	9	43.4	0.000
pad	9	542.5	0.000	solder:panel	2	6.0	0.050
panel	2	68.1	0.000	mask:pad	27	61.5	0.000
open:solder	2	28.0	0.000	mask:panel	6	21.2	0.002
open:mask	6	71.0	0.000	pad:panel	18	13.7	0.748
solder:mask	3	59.8	0.000				

# Chambers & Hastie (1992) model with three 2-factor interactions

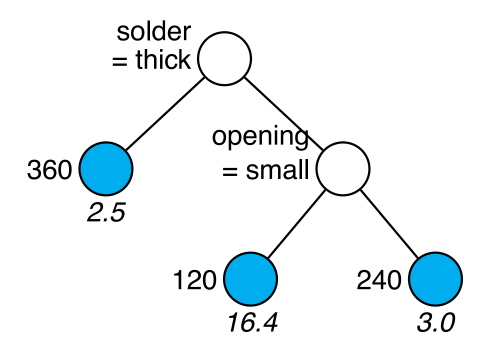
Regressor	Coef	t-stat	Regressor	Coef	t-stat
Constant	-2.668	-9.25			
maskA3	0.396	1.21	openmedium	0.921	2.95
maskB3	2.101	7.54	opensmall	2.919	11.63
maskB6	3.010	11.36	soldthin	2.495	11.44
padD6	-0.369	-5.17	maskA3:openmedium	0.816	2.44
padD7	-0.098	-1.49	maskB3:openmedium	-0.447	-1.44
padL4	0.262	4.32	maskB6:openmedium	-0.032	-0.11
padL6	-0.668	-8.53	maskA3:opensmall	-0.087	-0.32
padL7	-0.490	-6.62	maskB3:opensmall	-0.266	-1.12
padL8	-0.271	-3.91	maskB6:opensmall	-0.610	-2.74
padL9	-0.636	-8.20	maskA3:soldthin	-0.034	-0.16
padW4	-0.110	-1.66	maskB3:soldthin	-0.805	-4.42
padW9	-1.438	-13.80	maskB6:soldthin	-0.850	-4.85
panel2	0.334	7.93	openmedium:soldthin	-0.833	-4.80
panel3	0.254	5.95	opensmall:soldthin	-0.762	-5.13

# **GUIDE** piecewise-constant Poisson model



Estimated mean number of solder skips given under each leaf node

## **GUIDE** piecewise main effects Poisson model

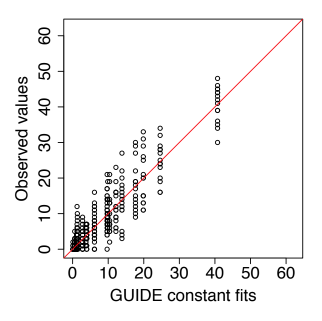


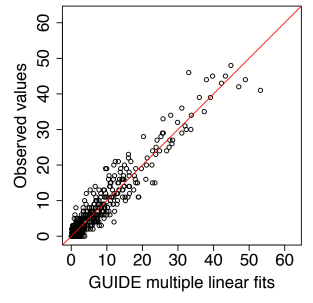
Number in italics below terminal node is sample mean of solder skips.

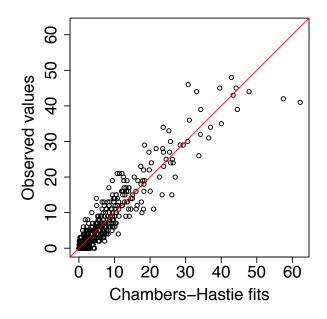
Number beside terminal node is sample size.

	solder = thick			solder	= thin		
			opening	= small	medium o	medium or large	
Regressor	Coef	t-stat	Coef	t-stat	Coef	t-stat	
Constant	-2.43	-10.68	2.08	21.5	-0.37	-1.9	
maskA3	0.47	2.37	0.31	3.3	0.81	4.5	
maskB3	1.83	11.01	1.05	12.8	1.01	5.8	
maskB6	2.52	15.71	1.50	19.3	2.27	14.6	
openmedium	0.86	5.57	aliased		0.10	1.4	
opensmall	2.46	18.18	aliased		aliased		
panel2	0.22	2.72	0.31	5.5	0.58	5.7	
panel3	0.07	0.81	0.19	3.2	0.69	6.9	
padD6	-0.32	-2.03	-0.25	-2.8	-0.80	-4.6	
padD7	0.12	0.85	-0.15	-1.7	-0.19	-1.3	
padL4	0.70	5.53	0.08	1.0	0.21	1.6	
padL6	-0.40	-2.46	-0.72	-6.8	-0.82	-4.7	
padL7	0.04	0.29	-0.65	-6.3	-0.76	-4.5	
padL8	0.15	1.05	-0.43	-4.5	-0.36	-2.4	
padL9	-0.59	-3.43	-0.64	-6.3	-0.67	-4.1	
padW4	-0.05	-0.37	-0.09	-1.0	-0.23	-1.6	
padW9	-1.32	-5.89	-1.38	-10.3	-1.75	-7.0	

### **Observed vs. fitted values**

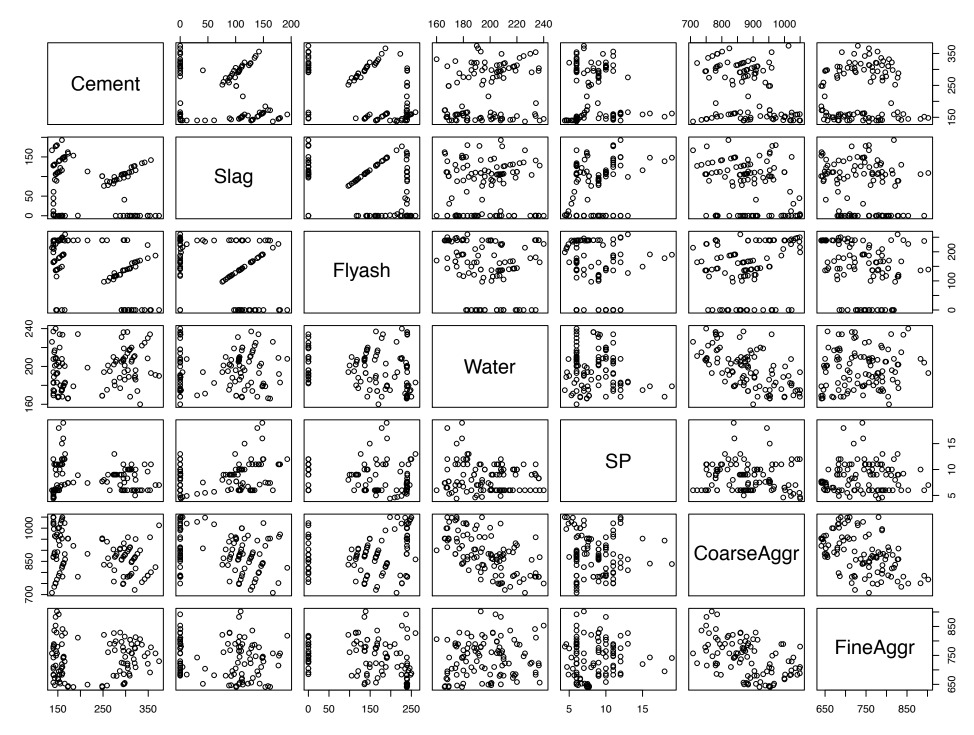






# Multiresponse data: viscosity and strength of concrete (Yeh, 2007)

- 103 observations on seven input variables (kg per cubic meter):
  - 1. Cement
  - 2. Slag
  - 3. Fly ash
  - 4. Water
  - 5. Superplasticizer
  - 6. Coarse aggregate
  - 7. Fine aggregate
- Three output variables:
  - 1. Slump (cm)
  - 2. Flow (cm)
  - 3. 28-day compressive strength (Mpa)



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Classification and Regression Trees

# **Separate linear models**

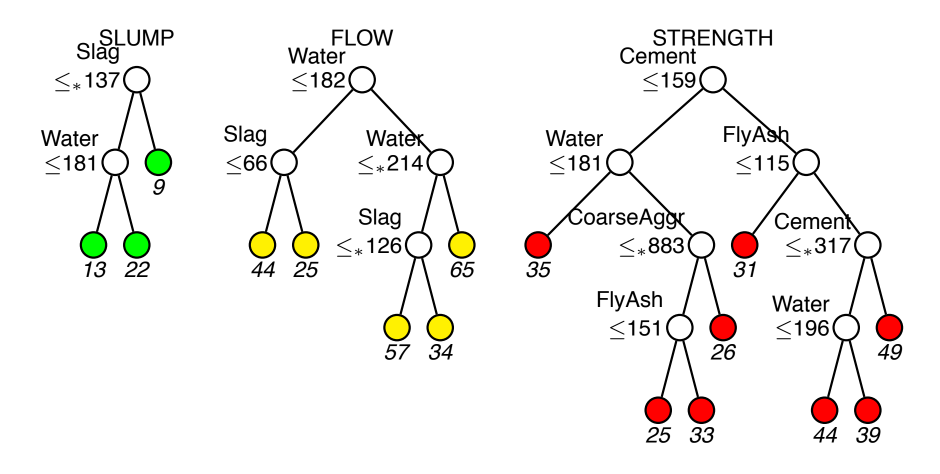
	Slui	mp	Flo	w	Strer	Strength	
	Estimate	P-value	Estimate	P-value	Estimate	P-value	
(Intercept)	-88.525	0.66	-252.875	0.472	139.782	0.052	
Cement	0.010	0.88	0.054	0.634	0.061	0.008	
Slag	-0.013	0.89	-0.006	0.971	-0.030	0.352	
Flyash	0.006	0.93	0.061	0.593	0.051	0.032	
Water	0.259	0.21	0.732	0.041	-0.23270	0.002	
SP	-0.184	0.63	0.298	0.654	0.103	0.445	
CoarseAggr	0.030	0.71	0.074	0.587	-0.056	0.045	
FineAggr	0.039	0.64	0.094	0.509	-0.039	0.178	

Is there really nothing significant for Slump?

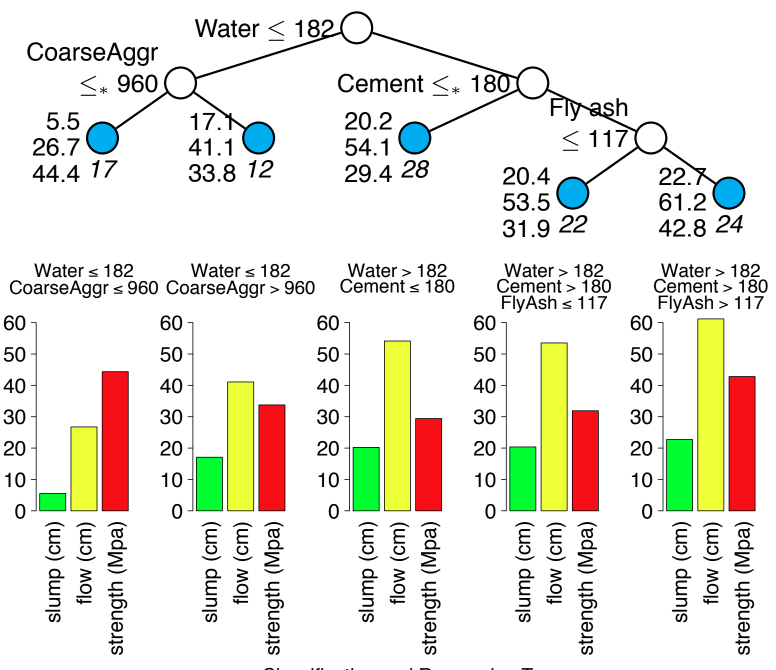
# Water and Slag are highly significant for Slump if no other variables are in the model!

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	-18.099	7.314	-2.475	0.01502 *
Water	0.199	0.036	5.455	3.56e-07 ***
Slag	-0.039	0.012	-3.227	0.00169 **
(Intercept)	11.370	9.683	1.174	0.243
Water	0.050	0.0486	1.025	0.308
Slag	-0.479	0.104	-4.604	1.23e-05 ***
Water:Slag	0.002	0.001	4.251	4.83e-05 ***

# One tree for each response variable



# One tree for all response variables



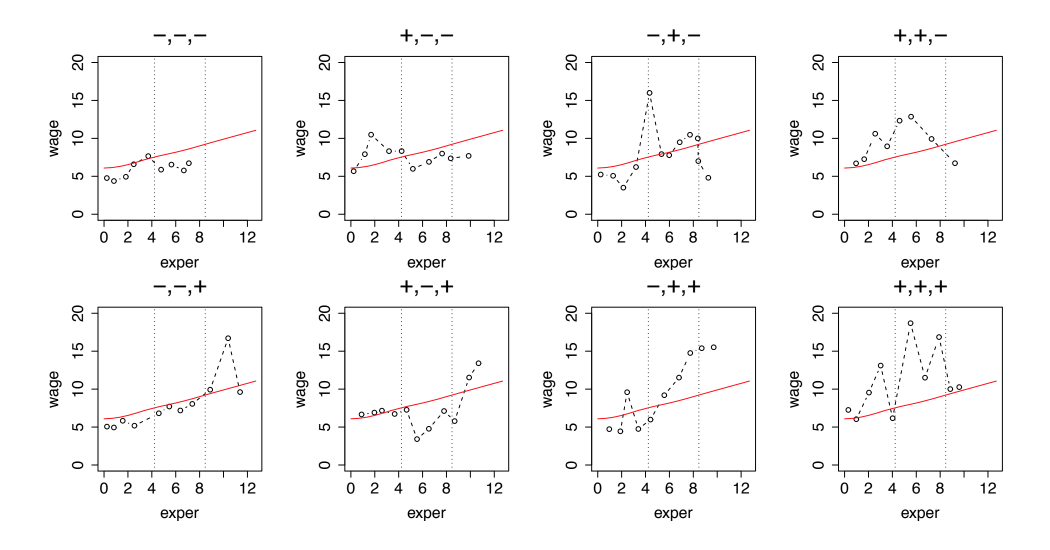
# Longitudinal data: Hourly wage of high-school dropouts

- 888 male high-school dropouts (246 Black, 204 Hispanic, 438 White) observed over time
- Response is hourly wage (in 1990 dollars)
- Predictor variables are:
  - 1. hgc: highest grade completed (6–12)
  - 2. exper: years in labor force (0.001–12.7 yrs)
  - 3. black: 1 if Black, 0 otherwise
  - 4. hisp: 1 if Hispanic, 0 otherwise
- Data from the National Longitudinal Survey of Youth
- References: Murnane et al. (1999), Singer and Willett (2003, Sec. 5.2.1)

# Design details and complications

- 1. At first wave of data collection, subjects varied in age from 14–17
- 2. Some subsequent waves separated by one year, others by two
- 3. Each wave's interviews conducted at different times in calendar year
- 4. Subjects observed at random times and random number of times: 77 have 1–2, 82 have 3–4, 166 have 5–6, 226 have 7–8, 240 have 9–10, and 97 have more than 10 observations
- 5. Subjects could describe more than one job at each interview
- 6. Subjects drop out of school and enter labor force at varying times
- 7. Subjects can change jobs at any time
- 8. Murnane et al. (1999) clocked time from each subject's first day of work

# Some individual trajectories



# Questions in analysis of longitudinal data

- 1. How does the outcome change over time?
- 2. Can we predict the differences in these changes?

# Two popular approaches

**Parametric:** Fit a *mixed model* (also called *individual growth model, random coefficient model, multilevel model*, and *hierarchical linear model*) and deduce the effect of predictor variables from the regression coefficients

**Nonparametric:** *Cluster* the subject trajectories, then *test* each predictor variable for its effect on the clusters

# Linear mixed model (Singer and Willett, 2003)

$$\log(\text{wage}) = \beta_0 + \beta_1 \text{hgc} + \beta_2 \text{exper} + \beta_3 \text{black} + \beta_4 \text{hisp}$$
  $+ \beta_5 \text{exper} \times \text{black} + \beta_6 \text{exper} \times \text{hisp}$   $+ b_0 + b_1 \text{exper} + \epsilon$ 

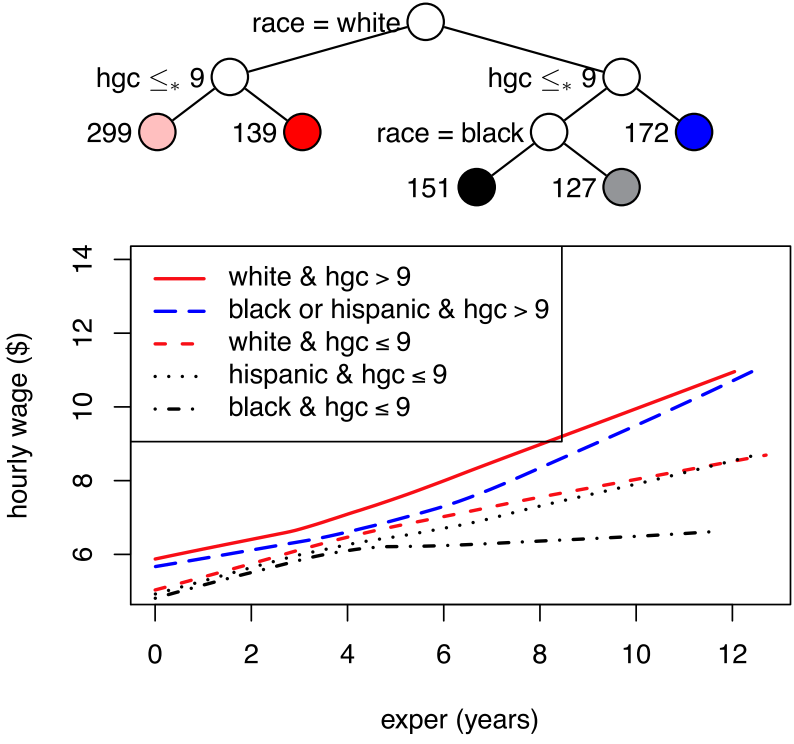
#### Assumptions/limitations:

- 1. Random (subject) intercepts and slopes  $b_0 \sim N(0, \sigma_0^2)$  and  $b_1 \sim N(0, \sigma_1^2)$ ;  $\epsilon \sim N(0, \sigma^2)$ ; all independent
- 2. Log transformation of wage to address skewness, linearize individual wage trajectories, and overcome range restriction
- 3. Predictions of wage requires exponentiation of fitted values of log(wage)
  - least-squares fit on log-dollar scale not best for dollar scale

## Coefficients of fixed effect terms

	Value	Std.Error	DF	t-value	p-value
(Intercept)	1.382	0.059	5511	23.43	0.000
hgc	0.038	0.006	884	5.94	0.000
exper	0.047	0.003	5511	14.57	0.000
black	0.006	0.025	884	0.25	0.804
hisp	-0.028	0.027	884	-1.03	0.302
exper×black	-0.015	0.006	5511	-2.65	0.008
exper×hisp	0.009	0.006	5511	1.51	0.131

<sup>&</sup>quot;Analyses not shown here suggest that we cannot distinguish statistically between the trajectories of Hispanic and White dropouts." (Singer and Willett, 2003, p. 149)

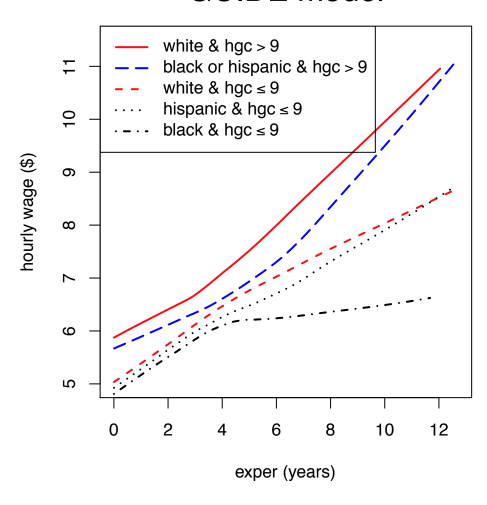


## LME vs. GUIDE fits

#### Linear mixed model

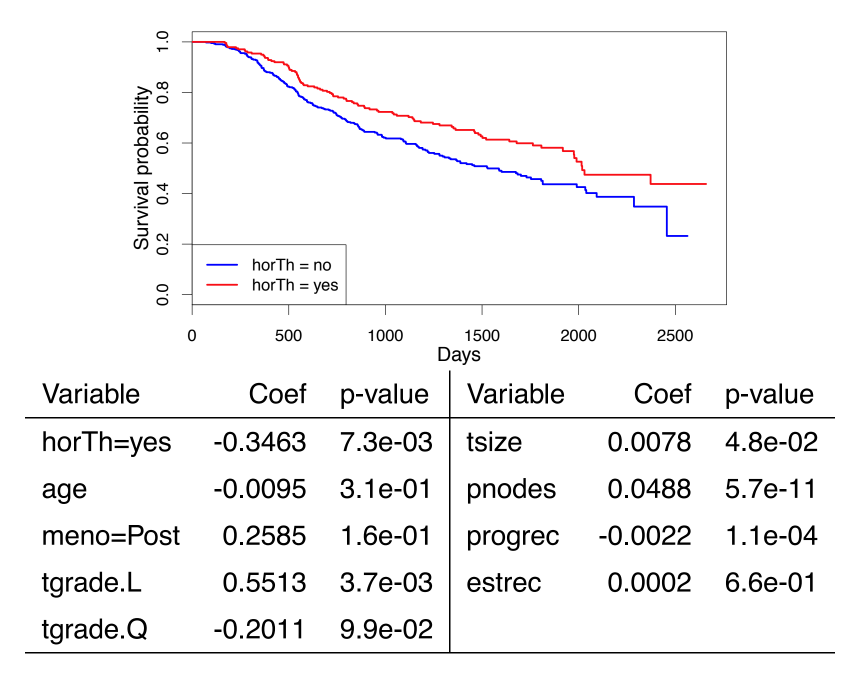
#### 12th grade non-black 12th grade black 6th grade non-black 6th grade black 10 0 wage (\$) $\infty$ 9 2 10 12 0 2 6 8 exper (years)

#### **GUIDE** model



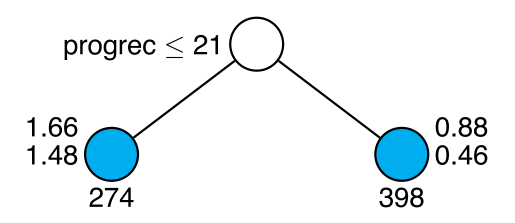
## Censored response data: breast cancer

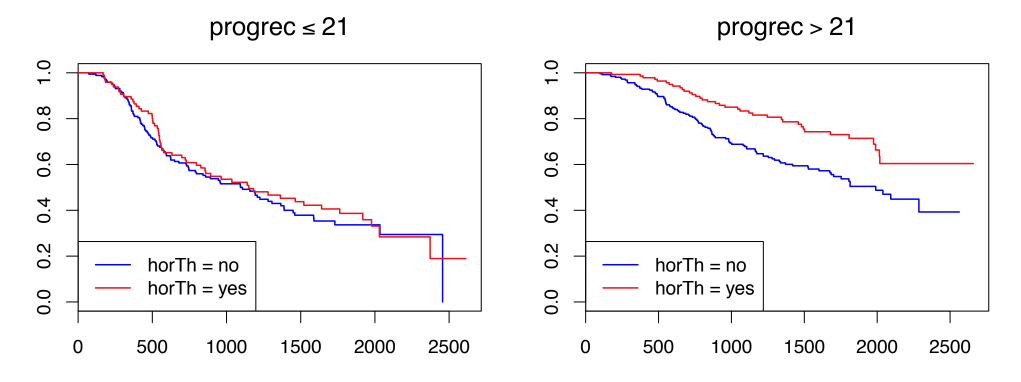
- Randomized clinical trial of 672 subjects with primary node positive breast cancer (Schumacher et al., 1994; data from **ipred** R package; 14 subjects with censored times less than smallest uncensored time excluded)
- Response is recurrence-free survival time (8–2659 days, 299 uncensored, 387 censored)
- Eight predictor variables with no missing values:
  - 1. **horTh** (hormone therapy, yes/no)
  - 2. **age** (21–80 years)
  - 3. **tsize**(tumor size, 3–120 mm)
  - 4. **pnodes**(number of positive lymph nodes, 1–51)
  - 5. **progrec** (progesterone receptor status, 0–2380 fmol)
  - 6. **estrec** (estrogen receptor status, 0–1144 fmol)
  - 7. **menostat** (menopausal status, pre/post)
  - 8. **tgrade** (tumor grade, 1, 2, 3)



#### Is there a subgroup where hormone therapy is ineffective?

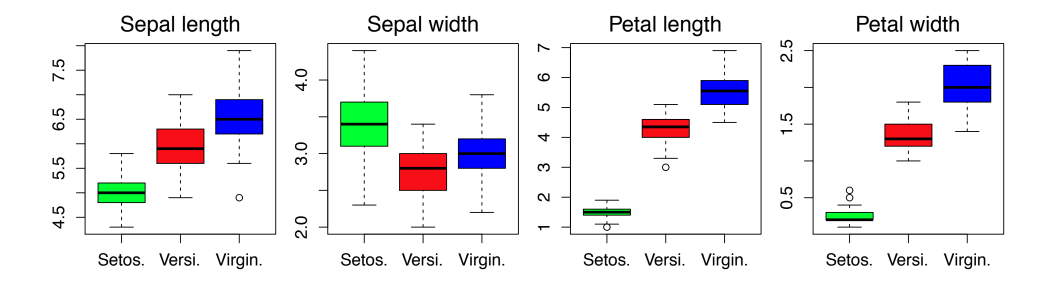
## **GUIDE** tree for subgroup identification



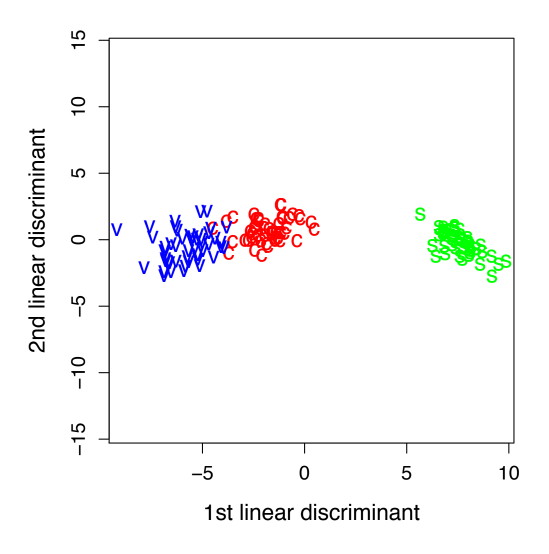


## Classification: Fisher's iris data

- 3 classes (Setosa, Versicolour, Virginica)
- 50 observations per class
- 4 predictor variables (petal length and width, sepal length and width)

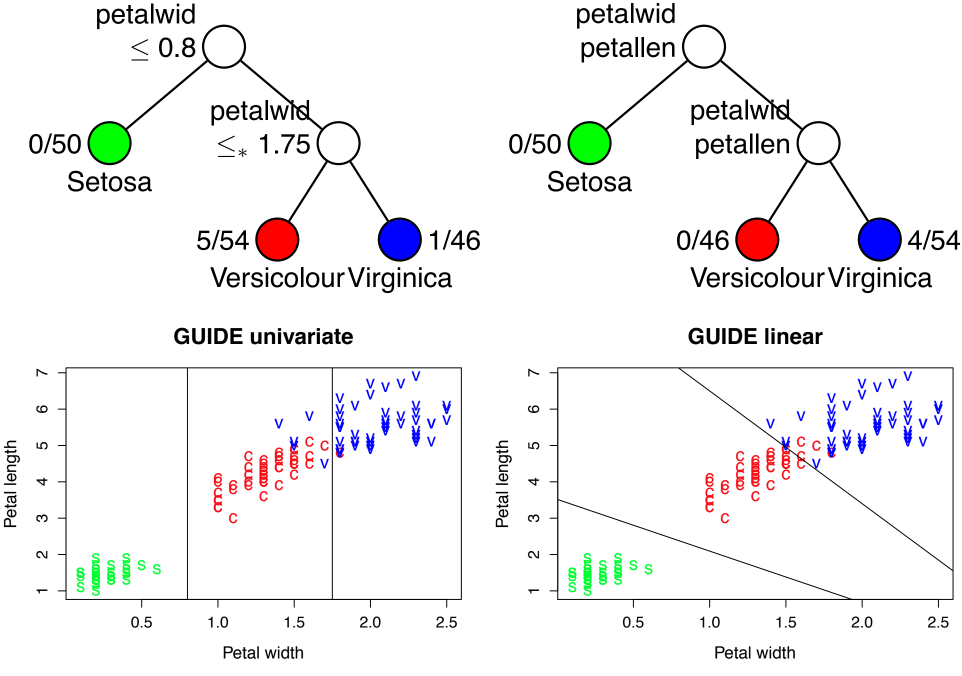


## Plot of iris data in first 2 discriminant coords



s = Setosa, c = Versicolour, v = Virginica

## Classification trees for iris data



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## Highly unbalanced classes: credit card data

- Goal: A major credit card company wants to find out why 14.8% of its card holders are dissatisfied
- Data: 22,242 card holder records with information on 24 predictor variables
- Missing values: 1,752 records contain one or more missing values; 0.34% missing values overall
- Response variable: whether a card holder is satisfied with the card
- Problem: Low percent of dissatisfied card holders makes most methods classify everyone as "satisfied"—a useless result
- Two solutions: Use equal priors or make cost of misclassifying dissatisfied
   = 5.5 × that of satisfied (more emphasis on identifying dissatisfied card holders)

## Predictor variables for credit card data

numadv30 How many times did you get cash advances in last 30 days?

spend30 How much money did you spend on purchases in last 30 days? (\$)

numpur30 How many times did you make purchases in last 30 days?

over30 Have you gone over limit in last 30 days? (1=yes 0 = no)

otherbal How much balance do you carry on other bank cards?

(0=0K, 1=0-2.5k, 2=2.5K-5K, ..., 8 = 17.5k-20k, 9 = 20k+)

othercred How much credit do you have on other bank cards?

(0=0K, 1=0-2.5k, 2=2.5K-5K, ..., 8 = 17.5k-20k, 9 = 20k+)

apply How many times did you apply for credit card in last year?

joint Do you have a joint account? (1 = yes 0 = no)

employ Are you currently employed? (1 = yes 0 = no)

cardyrs How many years have you had any credit card?

dailybal The average daily balance, unit in \$

currentbal The current balance, unit in \$

credlim The current credit limit, unit in \$100

mpastdue How many months the customer is past due

apr The annual percent rate, unit in %

worthy Historical index, credit worthiness, range [0,400]

months How many months has the customer had the card?

init Initial credit limit when account was opened, unit in \$100

adv1 Cash advance indicator for month -1, 1 = yes, 0 = no

adv2 Cash advance indicator for month -2, 1 = yes, 0 = no

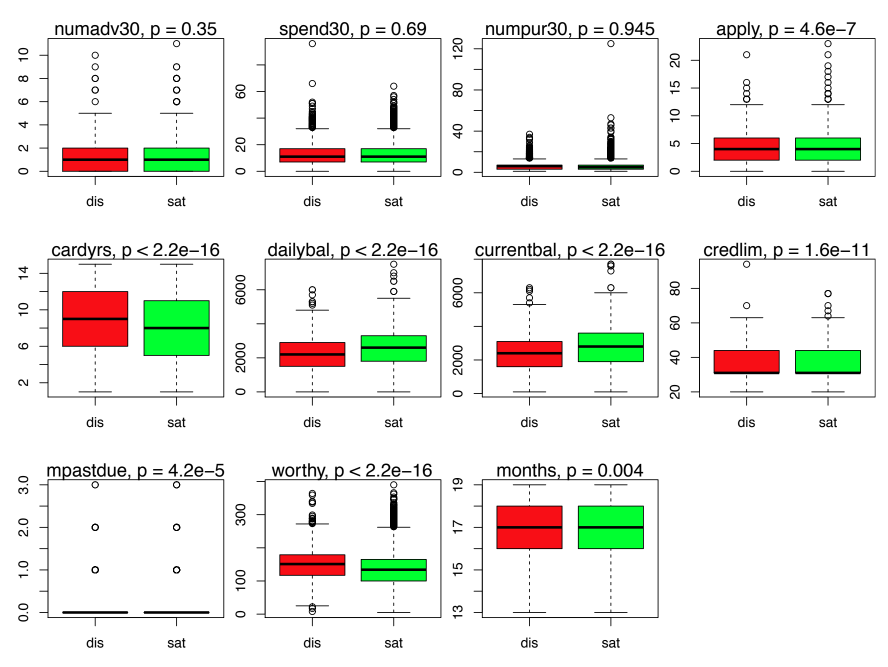
adv3 Cash advance indicator for month -3, 1 = yes, 0 = no

adv4 Cash advance indicator for month -4, 1 = yes, 0 = no

adv5 Cash advance indicator for month -5, 1 = yes, 0 = no

adv6 Cash advance indicator for month -6, 1 = yes, 0 = no

## t-tests on ordered predictors



W-Y Loh

Classification and Regression Trees

# **Chi-squared tests of categorical predictors**

	over30 ( <i>p</i>	= 0.13)	joint (p	0 = 0.47	employ ( $p = 0.002$ )		
Satisfied	No	Yes	No	Yes	No	Yes	
Yes	17951	836	3875	15079	2394	16560	
No	3132	125	691	2597	351	2937	

	otherbal ( $p = 1.5 \times 10^{-13}$ )									
Satisfied	1	2	3	4	5	6	7	8	9	
Yes	9281	4711	1610	1308	497	471	199	194	533	
No	1370	947	356	242	98	92	19	34	109	

	othercred ( $p < 2.2 \times 10^{-16}$ )									
Satisfied	1	2	3	4	5	6	7	8	9	
Yes	3304	6107	2393	2469	1056	1075	505	522	1435	
No	312	915	491	501	227	256	120	110	343	

# Chi-squared tests of categorical predictors (cont'd.)

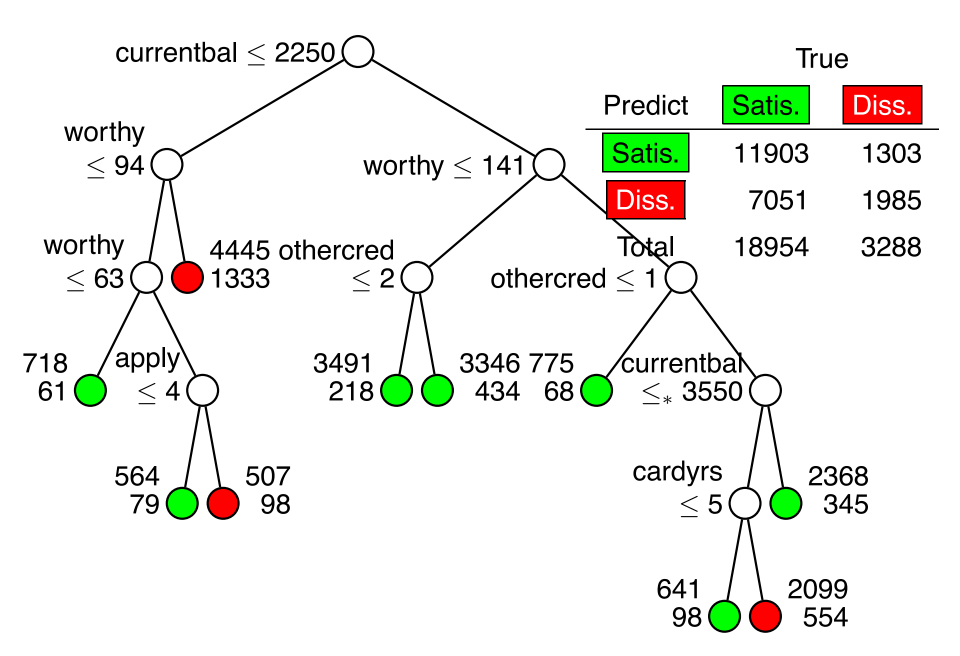
	apr ( $p = 0.002431$ )								
Satisfied	4	5	6	7	8	9	10	11	15
Yes	164	5	273	36	459	4	145	17386	482
No	24	6	42	11	59	1	27	3044	74

	init ( $p < 2.2 \times 10^{-16}$ )							
Satisfied	20	24	31	44				
Yes	3375	13	8062	7367				
No	773	8	1375	1114				

# Logistic regression model for Pr(Dissatisfied)

Variable	Estimate	p-value	Variable	Estimate	p-value
(Intercept)	-1.802e+00	7.12e-07	credlim	4.218e-02	8.20e-05
numadv30	-1.442e-02	0.517144	mpastdue	4.479e-01	3.42e-06
spend30	2.661e-03	0.399596	apr	1.556e-02	0.375681
numpur30	3.477e-03	0.594214	worthy	5.604e-03	<2e-16
over30	6.561e-02	0.529030	months	-4.112e-02	0.003214
otherbal	-7.053e-02	2.22e-05	init	-5.195e-02	2.19e-06
othercred	1.351e-01	< <b>2e-16</b>	adv1	-9.934e-02	0.374672
apply	3.229e-02	8.97e-05	adv2	-8.055e-03	0.938932
joint	-8.693e-02	0.081735	adv3	-3.709e-02	0.752908
employ	2.313e-01	0.000356	adv4	-2.381e-02	0.827685
cardyrs	3.080e-02	4.05e-09	adv5	1.072e-01	0.310609
dailybal	-5.665e-05	0.161080	adv6	-2.010e-02	0.841265
currentbal	-2.623e-04	1.83e-12			

# **GUIDE** tree with equal priors (or 5.5 to 1 costs)



## Properties of an ideal classifier

High predictive accuracy: classify unseen cases with low error

**Intuitive, comprehensible structure:** give insight into the roles and relative importance of the predictor variables

Correct, unbiased inference: draw inferences without bias

Fast training time: construct models quickly

### **Definition**

A classifier or classification rule is a function  $d(\mathbf{x})$  defined on  $\mathcal{X}$  such that for every  $\mathbf{x}$ ,  $d(\mathbf{x})$  is equal to one of the numbers  $1, 2, \dots, J$ .

A classifier is a partition of the sample space  $\mathcal{X}$  such that

$$A_j = \{\mathbf{x} : d(\mathbf{x}) = j\}$$
  
 $\mathcal{X} = \cup_j A_j$ 

## **Notations**

Y: response variable

J: number of classes

 $C = \{1, \dots, J\}$ : set of classes

N: training sample size

K: number of predictor variables

 $\mathbf{X} = (X_1, \dots, X_K)$ : vector of predictor variables

 $\mathcal{X}$ : Space of predictor variables

# AID (Morgan and Sonquist, 1963)

AID is the first published regression tree algorithm. It works as follows.

- 1. Recursively partition the data with splits of the form " $X \le c$ " (ordinal X) and " $X \in S$ " (categorical X).
- 2. At each stage, choose the split that minimizes a measure of node impurity, e.g., sum of squared deviations from mean:  $\sum (y_i \bar{y})^2$ .
- 3. Stop splitting if reduction in impurity is below preset value.

# **THAID** (Messenger and Mandell, 1972)

THAID is the first published classification tree algorithm (categorical Y)

- At each node, count the number of observations in the most frequent Y
  category (modal category)
- Choose the split that maximizes the sum of observations in the modal categories of the subnodes
- Follow the rest of the AID algorithm

## CART (Breiman et al., 1984)

- 1. Choose the split that maximizes the decrease in node impurity (Gini index for classification, sum of squared errors for regression)
- 2. For classification, let C(i|j) be cost of misclassifying a class j as class i. Assign terminal node t to class  $j^*$  if it minimizes the misclassification cost

$$\sum_{j} C(j^*|j)p(j|t) = \min_{i} \sum_{j} C(i|j)p(j|t)$$

For regression, use the sample Y mean in t as predicted value

- 3. Prune tree using test sample or cross-validation
- 4. Use surrogates splits to deal with missing values

## Estimates of misclassification error

Resubstitution estimate. Use the training data:

$$R(d) = N^{-1} \sum_{n=1}^{N} I(d(\mathbf{x}_n) \neq j_n)$$

**Test sample estimate.** Divide  $\mathcal{L}$  into  $\mathcal{L}_1$  and  $\mathcal{L}_2$ . Let  $N_2 = \#\mathcal{L}_2$ . Construct d from  $\mathcal{L}_1$ . Then

$$R^{ts}(d) = N_2^{-1} \sum_{\mathcal{L}_2} I(d(\mathbf{x}_n) \neq j_n)$$

#### V-fold cross-validation estimate.

- 1. Divide  $\mathcal{L}$  into subsets  $\mathcal{L}_1, \dots, \mathcal{L}_V$ . Let  $d^{(v)}$  be constructed from  $\mathcal{L} \mathcal{L}_v$ .
- 2. Define

$$R^{ts}(d^{(v)}) = N_v^{-1} \sum_{\mathcal{L}_v} I(d^{(v)}(\mathbf{x}_n) \neq j_n)$$

3. The *V*-fold cross-validation estimate is

$$R^{cv}(d) = V^{-1} \sum_{v=1}^{V} R^{ts}(d^{(v)})$$

## More notation

- t denotes a node
- J is the number of classes in training sample
- $J_t$  is the number of classes in t
- N(t) is the number of training samples in t
- $N_j$  is the number of class j training samples
- $N_i(t)$  is the number of class j training samples in t
- T denotes a tree
- $ilde{T}$  is the set of terminal nodes of T
- $|\tilde{T}|$  is number of terminal nodes of T
- $T_t$  is a subtree of T with root node t
- $\{t\}$  is a subtree of  $T_t$  containing only the root node t

# **Node impurity measures**

Let p(j|t) be the proportion of class j learning samples in node t. Define the **node impurity measure** 

$$i(t) = \phi(p(\cdot|t)) \ge 0$$

where  $\phi$  is a symmetric function with maximum value  $\phi(J^{-1},J^{-1},\dots,J^{-1})$  and

$$\phi(1,0,\ldots,0) = \phi(0,1,\ldots,0) = \ldots = \phi(0,0,\ldots,0,1) = 0$$

**Entropy:**  $i(t) = -\sum_{j=1}^{J} p(j|t) \log p(j|t)$ 

Gini index:  $i(t) = 1 - \sum_{j} p^2(j|t)$ 

- We use g(t) to denote the Gini index
- If J=2, then g(t)=2p(1|t)p(2|t), i.e., two times binomial variance

## Split set selection

1. Define the goodness of a split s as

$$\Delta i(s,t) = i(t) - p_L i(t_L) - p_R i(t_R)$$

where  $t_L$  and  $t_R$  are the left and right subnodes of t and  $p_L$  and  $p_R$  are the probabilities of being in those subnodes.

2. Define a set S of binary splits of the form  $X \in A$ , where,

$$A=(-\infty,c], \qquad ext{if } X ext{ is ordinal}$$
  $A\subset \mathcal{X}, \qquad ext{if } X ext{ is categorical}$ 

- (a) If X is ordinal with k unique values, there are (k-1) splits
- (b) If X is categorical with k unique values, there are  $(2^{k-1}-1)$  splits
- 3. Find  $s^* \in \mathcal{S}$  such that  $\Delta i(s^*, t) = \max_{s \in \mathcal{S}} \Delta i(s, t)$ .

# Shortcut for categorical splits with 2 classes

**Theorem 1** Let X be a categorical variable taking values in  $\{b_1, \ldots, b_L\}$ .

Suppose  $i(t) = \phi(p(1|t))$ , where  $\phi$  is strictly concave.

Define  $(b_{l(i)}; i = 1, ..., L)$  such that

$$p(1|X = b_{l(1)}) \le p(1|X = b_{l(2)}) \le \dots \le p(1|X = b_{l(L)})$$

Then the split on X that maximizes the decrease in impurity is one of the splits:

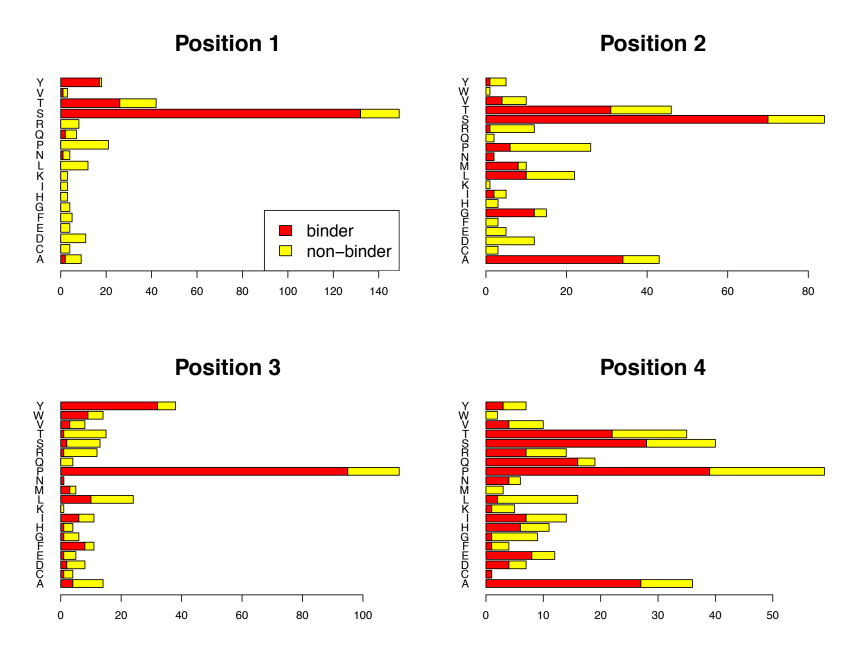
$$X \in \{b_{l(1)}, \dots, b_{l(h)}\}, h = 1, \dots, L-1$$

This reduces the search from  $(2^{L-1}-1)$  subsets to (L-1) subsets Proof: See Breiman et al. (1984, Sec. 9.4)

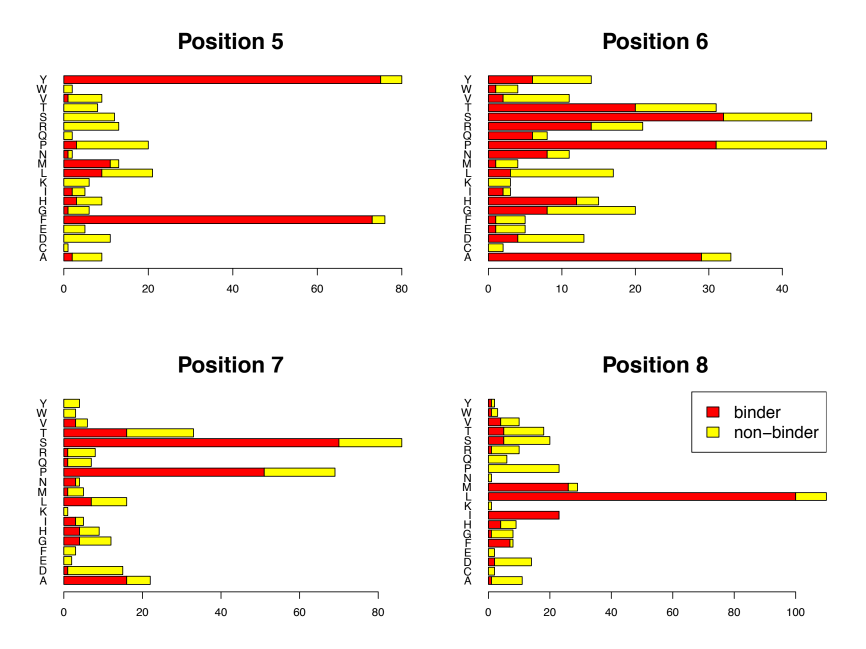
# Categorical predictors: peptide-binding data

- 310 amino acid sequences of peptides
- 181 bind to a class of MHC molecule, 129 do not
- Each amino acid sequence has length 8
- Each position in a sequence is one of 18–20 amino acids
- Problem: What amino acids in which positions are predictive of binding?
- Milik et al. (1998) convert amino acid info into 104 numerical "property variables" and use neural networks
- Segal et al. (2001) use CART
   http://repositories.cdlib.org/cbmb/peptide\_binding

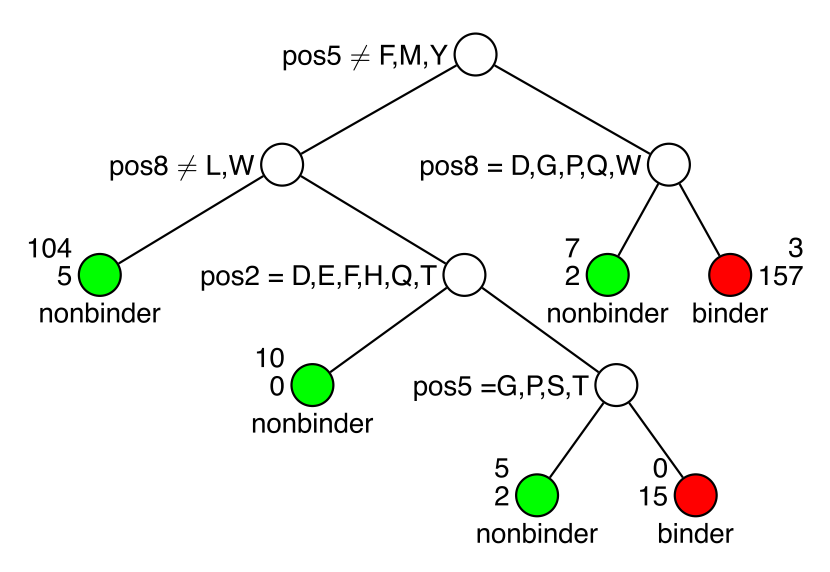
## Distributions of peptide-binding data



## Distributions of peptide-binding data (cont'd.)



# RPART (Therneau and Atkinson, 2012) tree for peptide data



Counts of nonbinder and binder, resp., are beside terminal nodes

# Levels of Pos5 ordered by P(Y = 0)

Class						Cla	SS		
Level	0	1	Total	Prop.	Level	0	1	Total	Prop.
F	3	73	76	0.039	V	8	1	9	0.889
Υ	5	75	80	0.063	С	1	0	1	1
M	2	11	13	0.154	D	11	0	11	1
Ν	1	1	2	0.5	Е	5	0	5	1
L	12	9	21	0.571	K	6	0	6	1
1	3	2	5	0.6	Q	2	0	2	1
Н	6	3	9	0.667	R	13	0	13	1
Α	7	2	9	0.778	S	12	0	12	1
G	5	1	6	0.833	Т	8	0	8	1
Р	17	3	20	0.85	W	2	0	2	1

## Resubstitution estimate of misclassification cost

- Let  $\pi(j)$  be the prior probability of class j
- Let  $N_i(t)$  be the number of class j observations in node t
- Let  $N_j$  be the number of class j observations in the training sample
- Let  $p(j,t) = \pi(j)N_j(t)/N_j$  be the estimated probability of being in class j and in node t
- Define  $p(t) = \sum_{j} p(j,t)$  and p(j|t) = p(j,t)/p(t)
- ullet The resubstitution estimate of expected misclassification cost of node t is

$$r(t) = \min_{i} \sum_{j} C(i|j)p(j|t)$$

ullet The resubstitution estimate of expected misclassification cost of a tree T is

$$R(T) = \sum_{t \in \tilde{T}} r(t)p(t)$$

# Why not use R(t) as impurity function?

- Optimal split is not unique: possible for  $R(t)-R(t_L)-R(t_R)=0$  for some or all splits
- Shortcut algorithm for categorical split is not applicable because R(t) is not a strictly concave function of  $\{p(j|t)\}$

# **CART** pruning

- 1. Given  $\alpha$  and tree T, define the cost-complexity  $R_{\alpha}(T) = R(T) + \alpha |\tilde{T}|$
- 2. For each  $\alpha$ , there is a tree T that minimizes the cost-complexity
- 3. Let t be any node and  $T_t$  be the branch of T with root node t. Then

$$R_{\alpha}(\{t\}) = R(t) + \alpha$$
  
 $R_{\alpha}(T_t) = R(T_t) + \alpha |\tilde{T}_t|$ 

- 4.  $R_{\alpha}(T_t) = R_{\alpha}(\{t\})$  when  $\alpha = u(t) = [R(t) R(T_t)]/[|\tilde{T}_t| 1]$
- 5. Prune branches at nodes  $t_1$  for which  $u(t_1) = \min\{u(t) : t \in T \tilde{T}\}$
- 6. Define  $\alpha_1 = u(t_1)$  and iterate to obtain a nested sequence of trees

Sequence of minimal cost-complexity trees is a subsequence of the subtrees constructed by finding the minimum cost subtree for a given number of terminal nodes.

### Subtree selection by V-fold cross-validation

- 1. Let  $\alpha_1 < \alpha_2 < \dots$  be the  $\alpha$ -values associated with the pruned sequence of subtrees  $T_1 \succ T_2 \succ \dots$  Define  $\alpha_k' = \sqrt{\alpha_k \alpha_{k+1}}$
- 2. Divide  $\mathcal{L}$  into V subsets  $\mathcal{L}_1, \ldots, \mathcal{L}_V$
- 3. Let  $T^{(v)}(\alpha_k')$  be the minimal cost-complexity tree grown from  $\mathcal{L} \mathcal{L}_v$ ,  $v = 1, \dots, V$
- 4. Let  $R'(T^{(v)}(\alpha_k'))$  be the estimate of the misclassification cost of  $T^{(v)}(\alpha_k')$  based on the test sample  $\mathcal{L}_v$
- 5. The V-fold CV estimate for subtree  $T_k$  is

$$R^{cv}(T_k) = V^{-1} \sum_{v=1}^{V} R'(T^{(v)}(\alpha_k'))$$

6. Select the subtree with the smallest CV cost

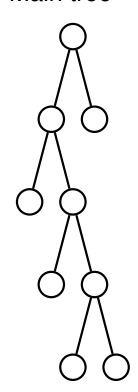
#### V-fold cross-validation

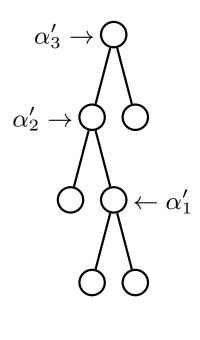
Main tree

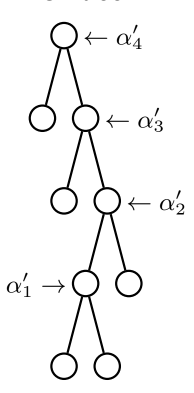
CV tree 1

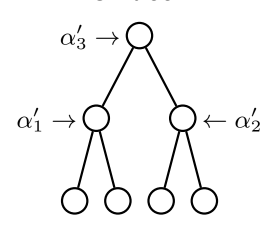
CV tree 2

CV tree V







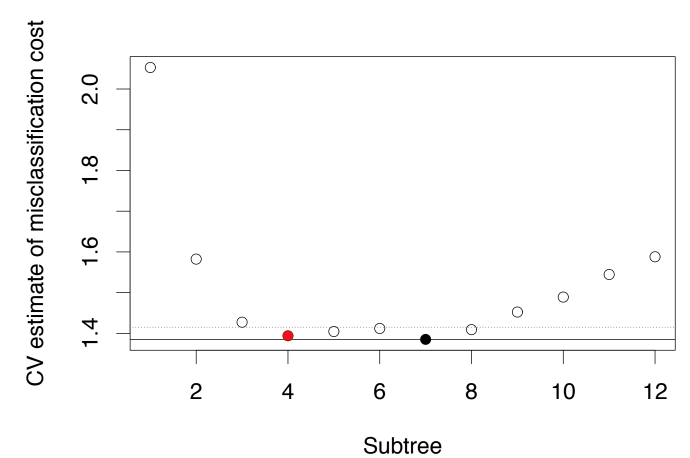


- Main tree is grown using all the data
- ullet Each CV tree is grown using (V-1) subsets

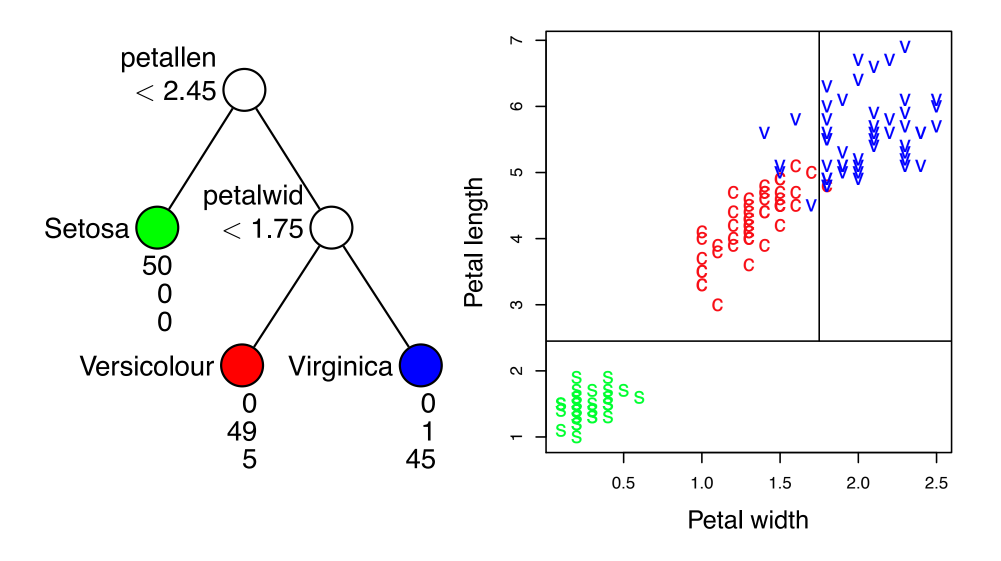
#### k-SE rule

- 1. Let  $\hat{R}(T_j)$  be the CV estimate of misclassification cost of  $T_j$ , let  $T^*$  be the tree with min. value of  $\hat{R}(T_j)$ , and let SE be the standard error of  $\hat{R}(T^*)$
- 2. The k-SE tree  $T^{**}$  is the smallest subtree such that

$$\hat{R}(T^{**}) \le \min_{j} \hat{R}(T_{j}) + k \times SE$$



#### **RPART** tree for iris data



### Unequal misclassification costs via Gini

The Gini index can be generalized to:

$$i(t) = \sum_{i,j} C(i|j)p(i|t)p(j|t)$$

This reduces for J=2 to

$$i(t) = [C(2|1) + C(1|2)]p(1|t)p(2|t)$$

which gives the same split criterion as for unit costs

Disadvantage: Index symmetrizes the cost matrix

#### Unequal misclassification costs via altered priors

- Let  $\pi(j)$  be the prior probability of class  $j \in \mathcal{C}$
- Let Q(i|j) be the proportion of class j cases in  $\mathcal{L}$  classified as class i by T
- Resubstitution estimate of T is  $R(T) = \sum_{i,j \in \mathcal{C}} C(i|j)Q(i|j)\pi(j)$
- The value of R(T) is the same if  $\{\pi'(j)\}$  and  $\{C'(i|j)\}$  satisfy

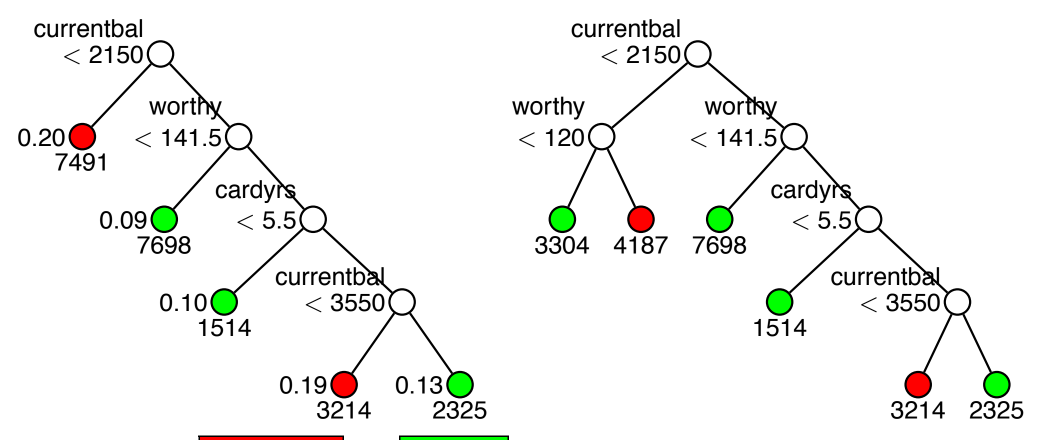
$$C'(i|j)\pi'(j) = C(i|j)\pi(j), i, j \in \mathcal{C}$$

- Thus unequal C(i|j) can be accommodated by altering  $\pi(j)$  to  $\pi'(j)$
- If C(i|j) = C(j),  $i \neq j$  for each j, define C'(i|j) = 1,  $i \neq j$  and

$$\pi'(j) = \frac{C(j)\pi(j)}{\sum_{i} C(i)\pi(i)}$$

- Otherwise, use  $C(j) = \sum_{i} C(i|j)$  in the above formula for  $\pi'(j)$
- Disadvantage: Only uses the values of  $\sum_{i} C(i|j)$

# RPART trees for credit card data: equal priors (left), 5.5:1 costs (right)

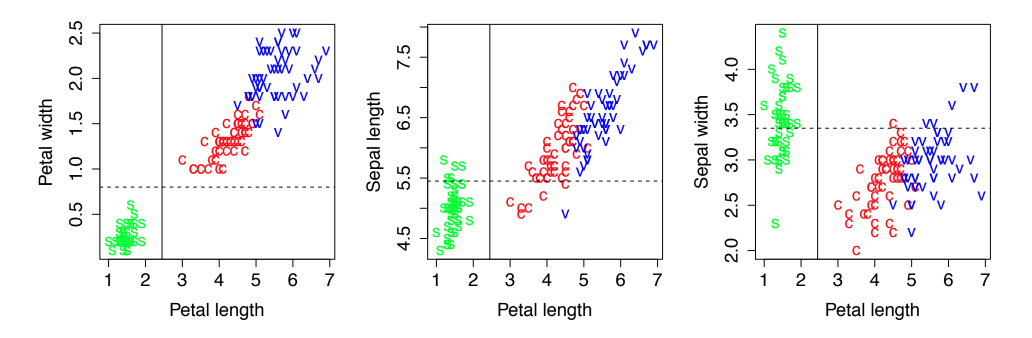


Dissatisfied and satisfied nodes in red and green colors
P(Dissatisfied) beside node in left tree; Sample sizes beneath nodes

### Missing values: CART surrogate splits

Suppose  $X \in S$  is selected to split node t

- 1. For each  $X_i \neq X$ , find the split  $X_i \in S_i$  that best predicts  $X \in S$  in terms of maximizing the number,  $M_i$ , of observations going to the corresponding subnodes
- 2. Order the  $X_i$  in terms of  $M_i$  to form a preferential set of surrogate splits



s = Setosa, c = Versicolour, v = Virginica

### **CART** surrogate splits: the details

- 1. Recall that  $p(j,t) = \pi(j)N_j(t)/N_j$  and  $p(t) = \sum_j p(j,t)$
- 2. Let  $s^*$  be the best split of t into  $t_L$  and  $t_R$
- 3. For each k, let  $S_k$  be the set of all splits on  $x_k$
- 4. Let  $s \in \mathcal{S}_k$  with subnodes  $t_L'$  and  $t_R'$
- 5. Let  $N_j(LL)$  be the number of class j cases in  $t_L \cap t_L'$
- 6. Define  $p(t_L \cap t_L') = \sum_j \pi(j) N_j(LL) / N_j$
- Let  $p_{LL}(s^*, s)$  be an estimate of  $P(\text{both } s^* \text{ and } s \text{ send a case left})$ :

$$p_{LL}(s^*, s) = p(t_L \cap t_L')/p(t)$$

- Similarly, define  $p_{RR}(s^*,s)=p(t_R\cap t_R')/p(t)$
- Estimate  $P(s \text{ predicts } s^*)$  by  $p(s^*, s) = p_{LL}(s^*, s) + p_{RR}(s^*, s)$
- $\tilde{s}_k$  is called a surrogate split on  $x_k$  for  $s^*$  if

$$p(s^*, \tilde{s}_k) = \max\{p(s^*, s) : s \in \mathcal{S}_k\}$$

### Measure of association for surrogate splits

- Let  $p_L$  and  $p_R$  be the probabilities that  $s^*$  sends a case to  $t_L$  and  $t_R$ , resp.
- The naive predictor sends every case to  $t_L$  if  $p_L \ge p_R$  and to  $t_R$  otherwise
- Error probability of the naive predictor is  $min(p_L, p_R)$
- Define the measure of association between  $s^*$  and s as the relative reduction in error:

$$\lambda(s^*, s) = \frac{\min(p_L, p_R) - [1 - p(s^*, s)]}{\min(p_L, p_R)}$$

- Rank the surrogate splits according to their  $\lambda(s^*, \tilde{s}_k)$  values
- If  $\lambda(s^*, \tilde{s}_k) \leq 0$ ,  $\tilde{s}_k$  is not used as a surrogate split

### **Uses of surrogate splits in CART**

- Enable tree construction when there are missing values in the learning sample
- 2. Enable classification of new cases with missing values
- 3. Rank variables by their order of importance (not available in RPART)
- 4. Detect masking of variables

# CART classification tree construction when there are missing values in the learning sample

Univariate splits: Find the best split  $s_k^*$  on each  $x_k$  using only cases non-missing in  $x_k$ . Select the split  $s^*$  that maximizes  $\Delta i(s,t)=i(t)-p_Li(t_L)-p_Ri(t_R)$ . Note: i(t) is constant for all splits but  $p_L$ ,  $p_R$ ,  $i(t_L)$ , and  $i(t_R)$  are computed from the non-missing values only. This induces a selection bias (Therneau and Atkinson, 2013, pp.18–19).

**Linear combination splits:** Find the best split  $s^*$  using only cases non-missing in all variables

Passing a case with missing values through the split: Let  $\tilde{s}_m$  be the surrogate split based on each variable  $x_m$  that is nonmissing for the case. Let  $\tilde{s}_{m^*}$  be the surrogate split among them with the highest measure of association with  $s^*$ . The split  $\tilde{s}_{m^*}$  is used on the case in place of  $s^*$ .

# CART classification of a new case with missing values

- Let  $s^*$  be the split at a node. Suppose the new case is missing some variable(s) that are required by  $s^*$ .
- Among all nonmissing variables in the case, find the one whose surrogate split  $\tilde{s}_k$  (say) has the highest measure of association with  $s^*$ .
- Send the case down using  $\tilde{s}_k$ . If no  $\tilde{s}_k$ , send the case to the larger node.

#### **Notes on RPART:**

- 1. If a split variable has no missing training values, it has no surrogate splits. In that case, new cases with missing values are sent to the larger node.
- 2. If a split is on a categorical variable X and a new case has an X value not in the training sample, RPART will return an error.

### Importance ranking of predictor variables in CART

• The importance of variable  $x_k$  is measured by

$$M(x_k) = \sum_{t \in T} \Delta i(\tilde{s}_k, t)$$

CART reports the standardized values

$$100M(x_k)/\max_m M(x_m)$$

The more obvious alternative measure

$$\sum_{t \in T} \Delta i(s_k^*, t)$$

is not used because it was found to be inferior

#### **Problems with CART classification**

- Biased toward variables with more splits: A k-valued ordered variable has (k-1) splits; a k-valued categorical variable has  $(2^{k-1}-1)$  splits.
- Biased toward predictors with more missing values: Split method uses only
  proportions of nonmissing cases—it ignores the number of missing values.
  A variable taking a unique value for exactly one case in each class and
  missing on all other cases yields the largest decrease in impurity. Bias
  exists for surrogate splits too.
- Computation: Impractical when there are three or more classes and categorical variables with many values. Note: Because CART and RPART encode each categorical variable split with a 32-bit binary integer, they do not properly deal with categorical variables having more than 32 values.
- Prediction accuracy: Often no better than linear discriminant analysis.

# FACT (Loh and Vanichsetakul, 1988) Classification trees with two or more splits/node

An approximate, quick, and fairly accurate solution with J splits per node:

- 1. Replace missing values by means and modes at each node
- 2. Convert each categorical variable to a dummy vector and then transform to largest discriminant variable (crimcoord)
- 3. For linear splits, use recursive linear discriminant analysis (LDA)
- 4. For univariate splits:
  - (a) Use one-way ANOVA to choose split variable or crimcoord
  - (b) Use LDA on selected variable or crimcoord to split node
  - (c) If split is on crimcoord, re-express it as an univariate split  $X \in S$
- 5. Use weighted sums of ANOVA F-statistics as importance scores

### **FACT** method for categorical variable splits

- 1. Suppose X takes values in the set  $\{a_1, \ldots, a_c\}$
- 2. Define dummy vector  $D = (d_1, \ldots, d_{c-1})$  with  $d_i = I(X = a_i)$
- 3. Project the D-data onto the largest discriminant coordinate (crimcoord)  $U = \sum_i b_i I(X = a_i)$
- 4. Search for a split of the form ' $U \leq c$ '
- 5. Re-express the split as ' $X \in A$ ' with  $A = \{a_i : b_i \leq c\}$

# QUEST (Loh and Shih, 1997) First algorithm with unbiased variable selection

- 1. If J > 2, use 2-means clustering of class means to form 2 superclasses
- 2. For univariate splits:
  - (a) Find p-value of 1-way ANOVA for each ordinal variable
  - (b) Find p-value of  $\chi^2$  test of independence for each categorical variable
  - (c) Select variable with smallest p-value to split node
  - (d) Transform each categorical variable to a crimcoord
  - (e) Use QDA on selected variable or crimcoord to find split
- 3. For linear combination splits, use FACT method (LDA on ordinal and crimcoord variables)
- 4. Use mean/mode imputation for missing values at each node
- 5. Use CART method to prune the tree

## CRUISE (Kim and Loh, 2001, 2003) First unbiased algorithm with multiple splits

- 1. Find p-value of  $\chi^2$  test of Y vs. each variable, with ordinal variables discretized (replaces F test of QUEST)
- 2. Find p-value of  $\chi^2$  test of Y vs. each **pair** of variables (adds ability to detect local interactions)
- 3. Select the variable(s) with smallest p-value; if latter is from an interaction test, select the variable with smaller marginal p-value
- 4. If selected variable is categorical, transform it to a crimcoord
- 5. Use Box-Cox transformations and LDA to split on selected variable
- 6. For linear combination splits, use LDA on all variables
- 7. Use different surrogate split methods for missing values
- 8. Optionally fit bivariate LDA models in nodes
- 9. Use CART method to prune the tree

### CRUISE 'alternate variable' missing value method

#### 1. For univariate splits:

- (a) Compute  $\chi^2$  tests using non-missing cases in the respective variables
- (b) For tree construction, impute missing values with class mean/mode
- (c) For predicting new cases, use the next best split at the node to predict the class and then impute with its mean/mode

#### 2. For linear combination splits:

- (a) For tree construction, impute with class mean/mode
- (b) For predicting new cases:
  - Use best univariate split to predict class; then impute with estimated class mean/mode
  - ii. If variable in best univariate split is also missing, impute with grand mean/mode

#### Prob. of surrogate/alternate variable selection

	CART				CRUISE					
	Percent missing $X_1$					Percent missing $X_1$				1
	1	2	3	4	25	1	2	3	4	25
$X_1$	.18	.12	.09	.05	.00	.19	.20	.18	.20	.18
$X_2$	.25	.25	.26	.24	.30	.18	.22	.18	.19	.19
$X_3$	.21	.23	.26	.27	.25	.22	.19	.20	.21	.19
$X_4$	.20	.23	.20	.23	.23	.22	.19	.22	.22	.21
$X_5$	.17	.17	.19	.21	.22	.20	.20	.22	.18	.23

- $Y \sim \text{Bernoulli}(1/2), X_0 \sim N(0.3Y, 1), \text{ and } X_1, \dots, X_5 \text{ indep. } N(0, 1)$
- Variable  $X_1$  has missing values but others do not
- Estimates based on 1000 iterations and n = 200 in each iteration
- Simulation standard errors about 0.015

# GUIDE classification (Loh, 2009) Improving on FACT, QUEST, and CRUISE

- 1. Use marginal and interaction  $\chi^2$  contingency table tests (as in CRUISE)
- 2. Use two-deep search to choose variable if split is based on interaction test (more powerful than CRUISE)
- 3. Allow linear splits on pairs of variables (new; useful for collinearity)
- 4. Use Bonferroni to control frequencies of interaction and linear splits (corrects CRUISE's propensity to split on interactions)
- 5. Allow kernel and nearest-neighbor node models (new; reduces tree size and yields predicted probabilities, *à la* logistic regression)
- 6. Treat missing values as a separate category in split selection (new; replaces imputation and surrogate splits)
- 7. Use CART method to prune the tree

### **GUIDE** marginal tests for ordinal X

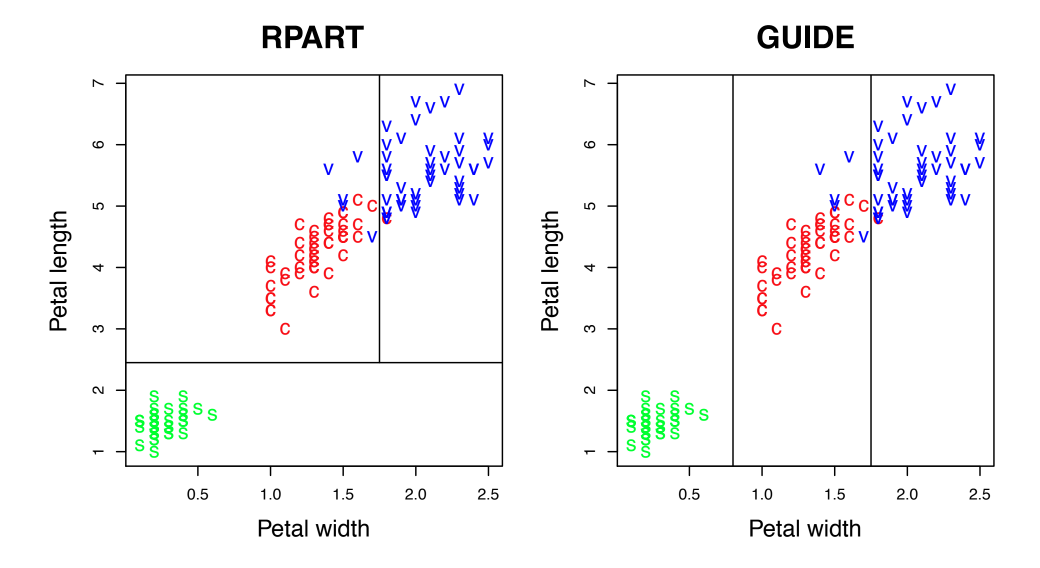
- 1. Compute the sample mean  $\bar{x}$  and SD s of X in t.
- 2. Define k=3 if  $N(t)<20J_t$ ; else k=4. Define  $b=2s\sqrt{3}/k$ .
- 3. Divide the range of X into k intervals with boundaries  $\bar{x} s\sqrt{3} + bj$ ; j = 1, 2, ..., k-1. Add one "interval" for missing values, if any.
- 4. Form a table with class values as rows and intervals as columns.
- 5. Let  $\nu$  be df of the table. Compute the chi-squared statistic  $\chi^2_{\nu}$  for testing independence.
- 6. Convert  $\chi^2_{\nu}$  to a 1-df chi-squared (Wilson and Hilferty, 1931)

$$W_M(X) = \max\left(0, \left[\frac{7}{9} + \sqrt{\nu} \left\{ \left(\frac{\chi_{\nu}^2}{\nu}\right)^{1/3} - 1 + \frac{2}{9\nu} \right\} \right]^3 \right).$$

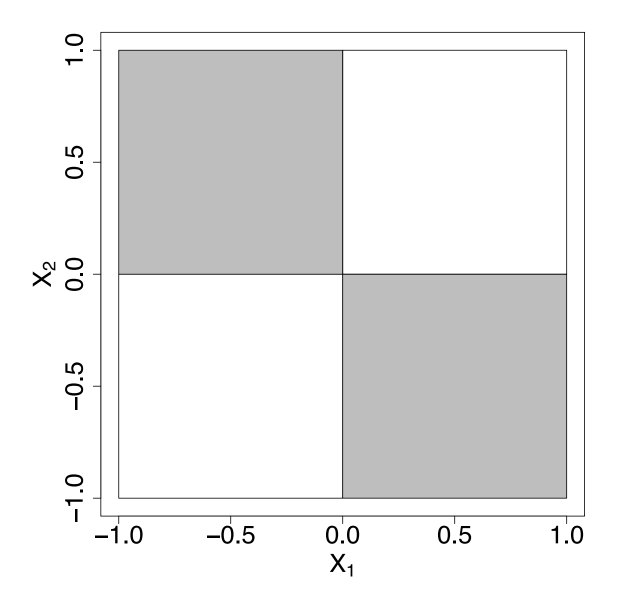
Note: For categorical X, use its values to form the columns of the table

### **Chi-squared tests**

	Petal length ( $\chi_6^2$ = 223.9)					Petal width ( $\chi_6^2 = 226.0$ )			
	≤2.2	(2.2, 3.7]	(3.7, 5.2]	>5.2	≤0.5	(0.5, 1.1]	(1.1, 1.8]	>1.8	
Setosa	50	0	0	0	49	1	0	0	
Versicol	0	7	43	0	0	10	40	0	
Virginica	0	0	18	32	0	0	16	34	
	5	Sepal length	$(\chi_6^2 = 109.2$	)	Sepal width ( $\chi_6^2$ = 64.6)				
	≤5.1	(5.1, 5.8]	(5.8, 6.5]	>6.5	≤2.6	(2.6, 3.0]	(3.0, 3.4]	>3.4	
Setosa	36	14	0	0	1	7	21	21	
Versicol	4	20	18	8	16	26	8	0	
Virginica	1	5	22	22	7	26	16	3	



### Two-class problem with interaction



## GUIDE split variable selection: interaction tests for $X_1, X_2$

1. Divide the  $(X_1, X_2)$ -space into sets

$$B_{k,m} = \{(x_1, x_2) : x_1 \in A_{1k}, x_2 \in A_{2m}\}, \quad k, m = 1, 2, \dots$$

where  $A_{1k}$  and  $A_{2m}$  are the respective intervals or categories

- 2. Form a contingency table with class labels as rows and  $\{B_{k,m}\}$  as columns
- 3. Compute chi-squared statistic and use Wilson-Hilferty approximation to convert it to a 1-df chi-squared value  $W_I(X_1,X_2)$

# Women's knowledge, attitude, and behavior toward mammography (Hosmer and Lemeshow, 2000)

- Data on 412 women and 3 classes:
  - 234 had no mammography experience;
  - 104 had a mammogram within the last year;
  - 74 had one more than a year ago
- 5 predictor variables: 2 binary; 2 ordered categorical; 1 ordinal

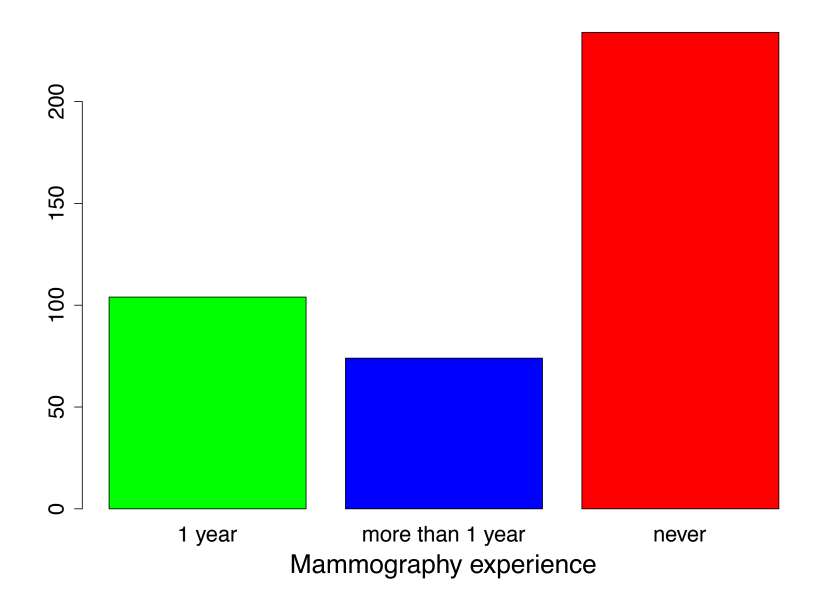
#### **Unequal misclassification costs**

	True class				
Predicted	1 (≤ 1 yr)	2 (> 1 yr)	3 (never)		
1 (≤ 1 yr)	0	1	2		
2 (> 1 yr)	1	0	1		
3 (never)	2	1	0		

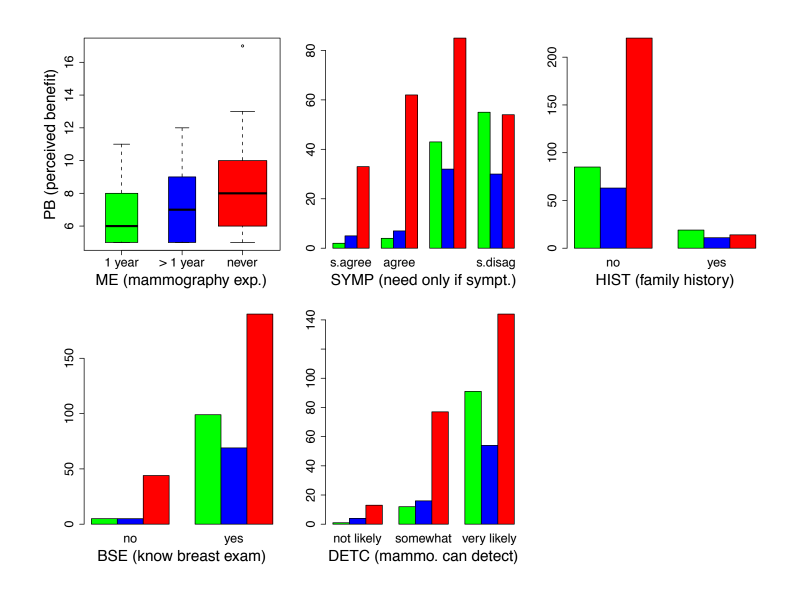
### Mammography variables

Name	Description	Values
ME	Mammography experience	within one year (1), over one year ago (2), never (3)
SYMP	You do not need a mam- mogram unless you develop symptoms	Strongly agree (1), agree (2), disagree (3), strongly disagree (4)
PB	Perceived benefit of mam- mography	5, 6,, 20 (low values imply greater perceived benefit)
HIST	Mother or sister with history of breast cancer	no (0), yes (1)
BSE	Anyone taught you how to examine your own breasts?	no (0), yes (1)
DETC	How likely is it that a mammo- gram can find a new case of breast cancer?	Not likely (1), somewhat likely (2), very likely (3)

#### **Distribution of classes**



#### Distributions of predictor variables



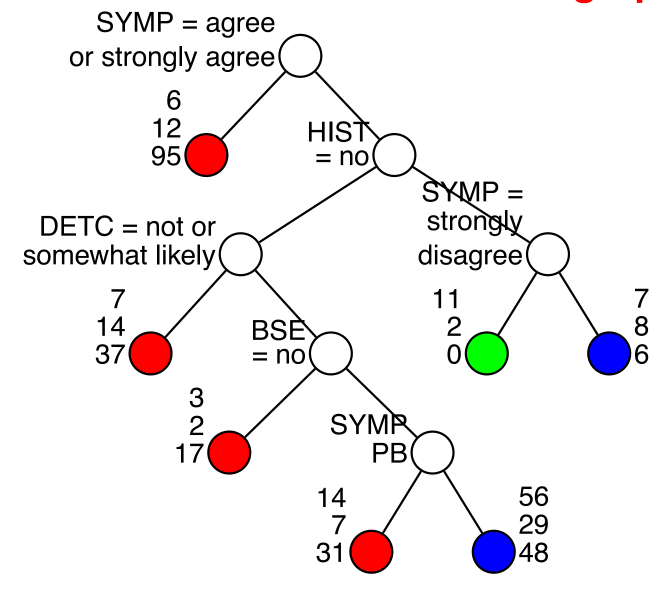
# Multinomial logistic regression model with "ME = never" as baseline category

Logit(ME = within 1 year)				Logit(ME = more than 1 year)			
Variable	Coef	SE	P-value	Variable	Coef	SE	P-value
Constant	-2.62	0.93	0.005	Constant	-1.82	0.86	0.033
SYMPD*	2.10	0.46	< 0.001	SYMPD*	1.13	0.36	0.002
PB	-0.25	0.07	0.001	PB	-0.15	0.07	0.034
HIST	1.31	0.43	0.003	HIST	1.06	0.45	0.019
BSE	1.24	0.53	0.019	BSE	0.96	0.51	0.056
DETCD**	0.89	0.36	0.019	DETCD**	0.11	0.32	0.720

<sup>\*</sup> SYMPD = 1 if SYMP = "disagree" or "strongly disagree", SYMPD = 0 otherwise

<sup>\*\*</sup> DETCD = 1 if DETC = "very likely", DETCD = 0 otherwise

#### **GUIDE** classification tree for mammography data



Within 1 year in green, more than one year in blue, never in red

### **Chi-squared tests**

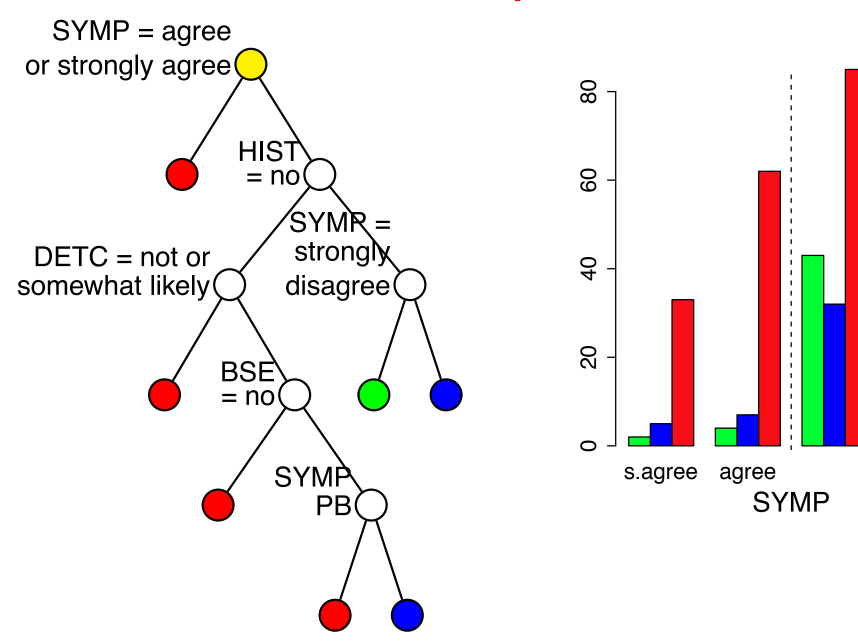
	SYMP ( $\chi_6^2$ = 57.2; $\chi_1^2 \approx$ 47)						
	strongly			strongly			
ME	agree	agree	disagree	disagree			
Never	33	62	85	54			
1 year	2	4	43	55			
> 1 yr	5	7	32	30			

	PB ( $\chi_6^2$ = 31.3; $\chi_1^2 \approx$ 19)						
ME	≤ <b>5.7</b>	(5.7, 7.6]	(7.6, 9.4]	> 9.4			
Never	33	68	65	68			
1 year	31	43	22	8			
> 1 yr	19	25	18	12			

	DETC ( $\chi_4^2$ = 24.1; $\chi_1^2 \approx$ 16			
	not	somewhat	very	
ME	likely	likely	likely	
Never	13	77	144	
1 year	1	12	91	
> 1 yr	4	16	54	

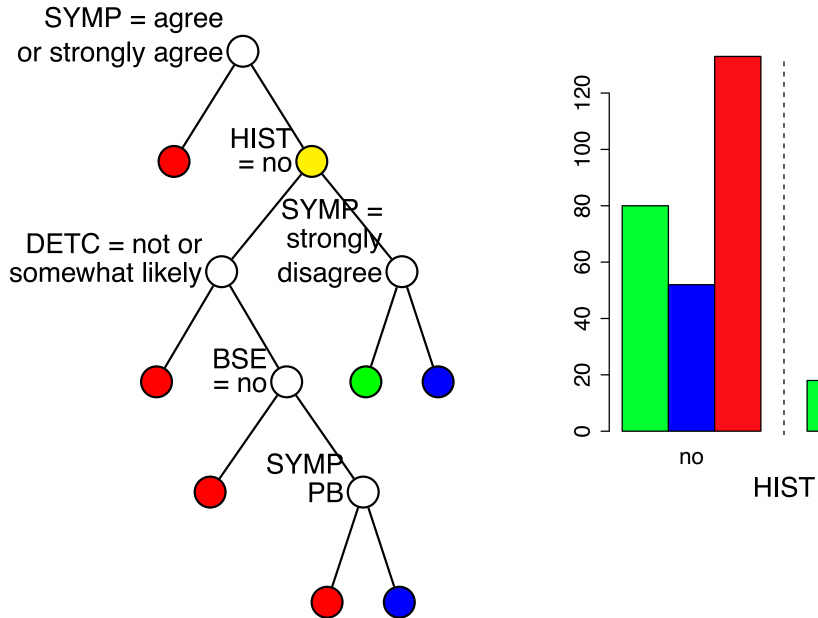
	BSE ( $\chi_2^2 = 1$	5.6, $\chi_1^2 pprox$ 13)	HIST	$(\chi_2^2$ = 13.1, $\chi_1^2 \approx$ 10)
ME	no	yes	no	yes
Never	44	190	220	14
1 year	5	99	85	19
> 1 yr	5	69	63	11

#### 1st split



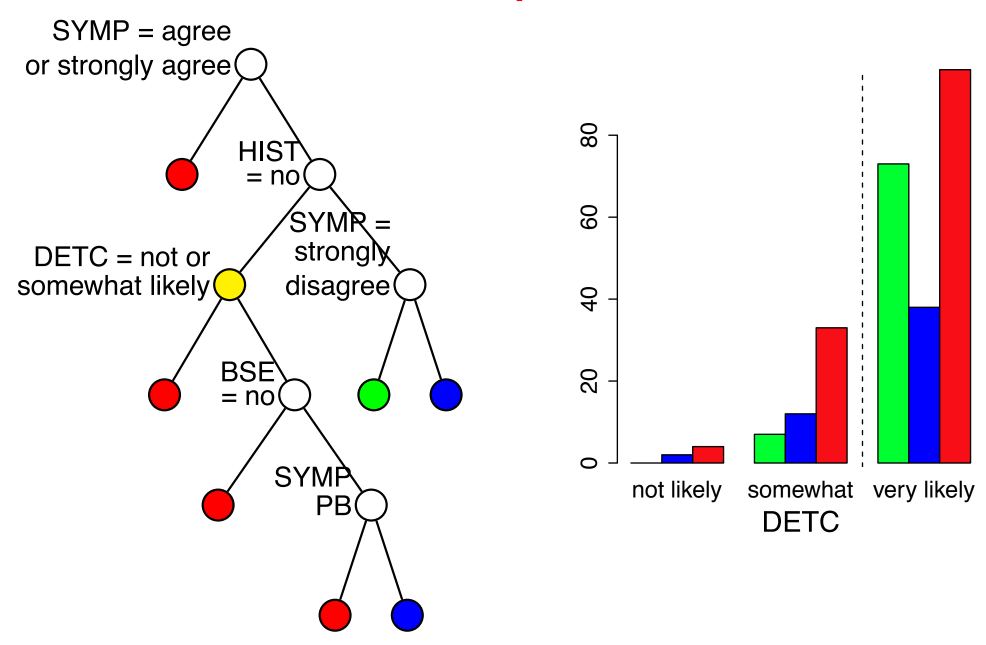
s.disagree

#### 2nd split

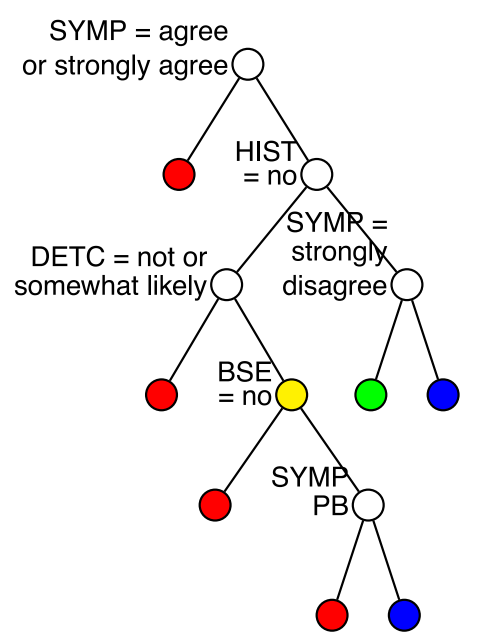


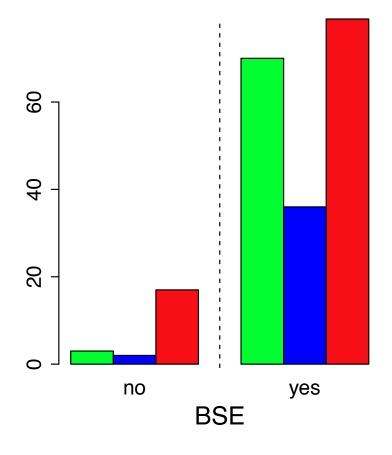
yes

#### 3rd split

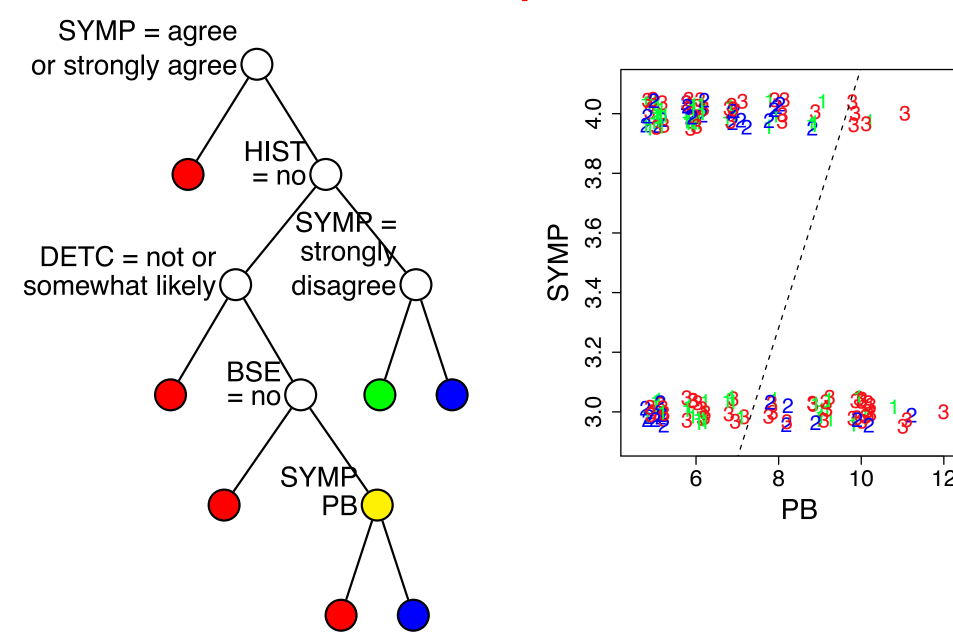


#### 4th split





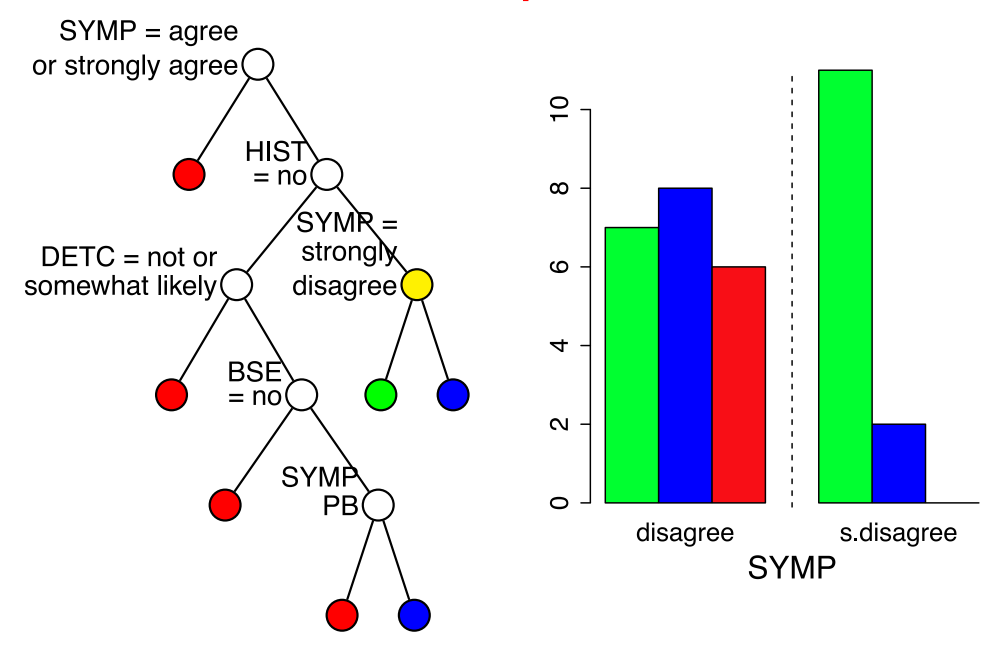
#### 5th split



12

10

#### 6th split



#### **SYMP-BSE** interaction test

	SYMP									
	stro	ngly	strongly							
	agree		agree		disagree		disagree			
	BSE		BSE		BSE		BSE			
ME	no	yes	no	yes	no	yes	no	yes		
0	6	27	15	47	15	70	8	46		
1	1	1	0	4	0	43	4	51		
2	1	4	0	7	2	30	2	28		

$$\chi_{14}^2$$
 = 72,  $\chi_1^2$  = 45, p =  $9 \times 10^{-10}$ 

### **GUIDE** split variable selection

- 1. Let K be the number of non-constant predictor variables in node t.
- 2. Let  $\chi^2_{\nu,\alpha}$  be the upper- $\alpha$  quantile of the chi-squared distribution with  $\nu$  df and define

$$\alpha = \frac{0.05}{K}, \quad \beta = \frac{0.1}{K(K-1)}$$

- 3. Find  $W_M(X_i)$  for each  $X_i$ .
- 4. (a) If  $\max_i W_M(X_i) > \chi^2_{1,\alpha}$ , select the variable with the largest  $W_M(X_i)$ .
  - (b) Otherwise, find  $W_I(X_i, X_i)$  for each pair of predictor variables.
    - i. If  $\max_{i\neq j} W_I(X_i, X_j) > \chi^2_{1,\beta}$ , select pair with largest  $W_I(X_i, X_j)$ .
    - ii. Otherwise, select variable with largest  $W_M(X_i)$ .

### Split set selection for ordinal X

Search all splits of the form  $X \leq c$  to minimize misclassification cost

### Split set selection for categorical X

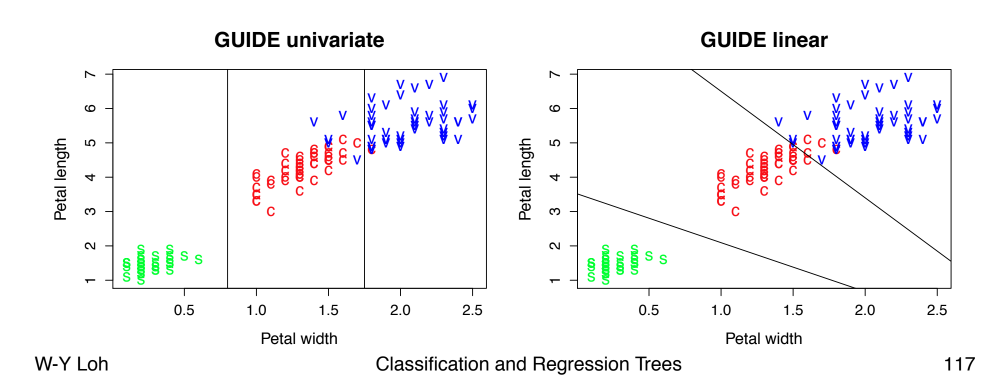
Suppose X takes distinct values  $\{a_1, a_2, \ldots, a_n\}$  in node t

- 1. If J=2 or  $n \leq 11$ , search all subsets S to find  $t_L=\{X \in S\}$
- 2. If  $J \le 11$  and n > 20, let class  $j_i$  minimize misclassification cost in  $t \cap \{X = a_i\}$ 
  - (a) Define  $X' = \sum_i j_i I(X = a_i)$  and search for the split based on X' that minimizes the decrease in impurity
  - (b) Express the split as  $t_L = \{X \in S\}$
- 3. Otherwise, use linear discriminant analysis on the dummy vectors

#### **Bivariate linear discriminant split**

Let  $X_1$  and  $X_2$  be two s variables in node t

- 1. For the jth class and each  $X_i$ , find node class mean  $\bar{x}_{i,j}$  and SD  $s_{i,j}$
- 2. Find trimmed set  $S_j$  of class j samples in t such that  $|X_i \bar{x}_{i,j}| \leq 2.5 s_{i,j}$
- 3. Find the larger linear discrim. coord. Z from observations in  $S_1 \cup \ldots \cup S_J$
- 4. Project the data in t onto the Z-axis
- 5. Compute the Wilson-Hilferty 1-df chi-squared  $W_L(X_1, X_2)$  from the Z values



### Summary of GUIDE classification split selection

Let K be the number of non-constant predictor variables and let  $K_1$  (< K) be the number that are ordinal. Define

$$\alpha = 0.05/K$$
,  $\beta = 0.1/\{K(K-1)\}$ ,  $\gamma = 0.1/\{K_1(K_1-1)\}$ 

- 1. Split on the  $X_i$  with the largest marginal  $\chi^2$ , if it is significant at level  $\alpha$
- 2. Otherwise,
  - (a) If  $(X_i, X_j)$  has largest interaction  $\chi^2$  and is significant at level  $\beta$ , use a two-deep search to find the best split on  $X_i$  or  $X_j$
  - (b) Otherwise, compute linear discriminant  $\chi^2$  for all ordinal pairs
    - i. Use most significant linear split if it is significant at level  $\gamma$
    - ii. Otherwise, choose the  $X_i$  with largest marginal  $\chi^2$

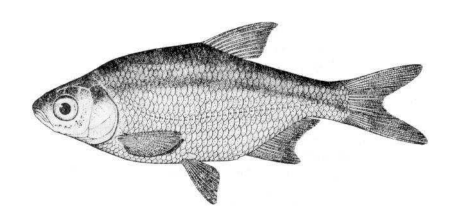
#### Fish classification

- 159 fish caught from the same lake near Tampere, Finland
- The fish are from 7 species: (1) 35 Bream, (2) 11 Parkki, (3) 56 Perch,
  (4) 17 Pike, (5) 20 Roach, (6) 14 Smelt, (7) 6 Whitefish

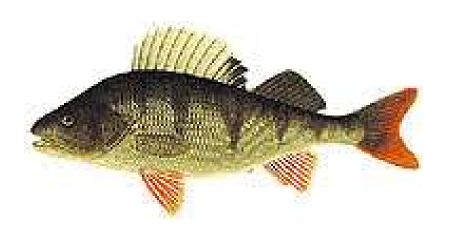
Predictor	Definition
Weight	Weight of the fish (in grams); one missing value
Length1	Length from the nose to the beginning of the tail (in cm)
Length2	Length from the nose to the notch of the tail (in cm)
Length3	Length from the nose to the end of the tail (in cm)
Height	Maximal height as % of Length3
Width	Maximal width as % of Length3
Sex	female, male, unknown

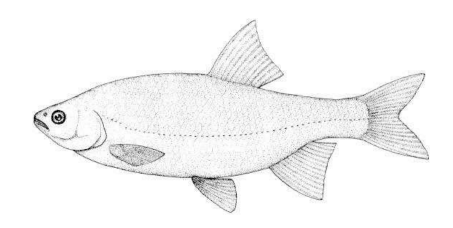
#### **Bream (left) and Parkki (right)**





## Perch (left) and Whitefish (right)



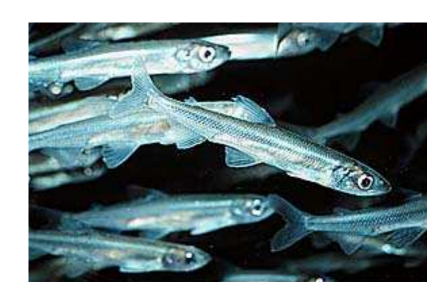


#### **Pike**

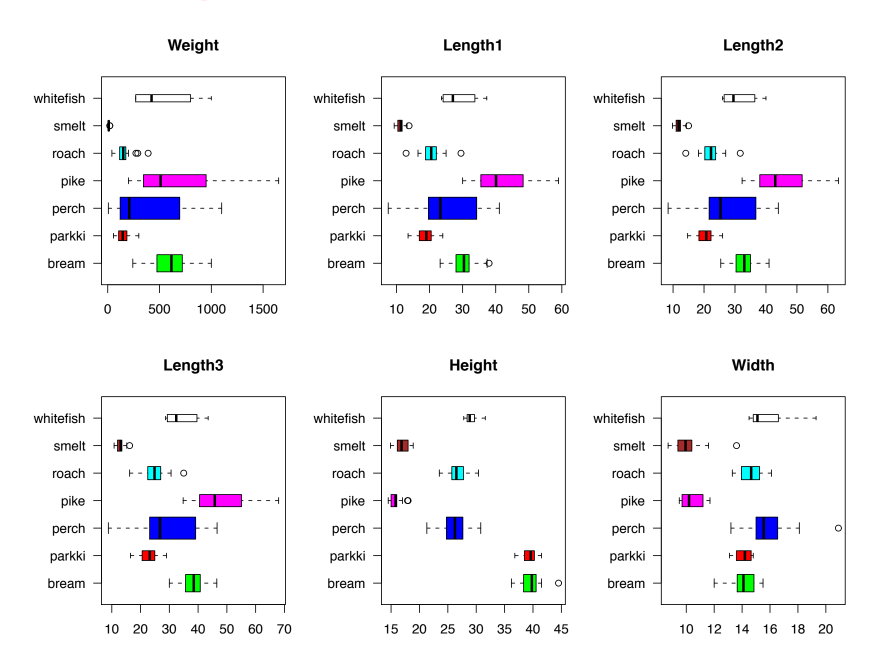


## Roach (left) and Smelt (right)





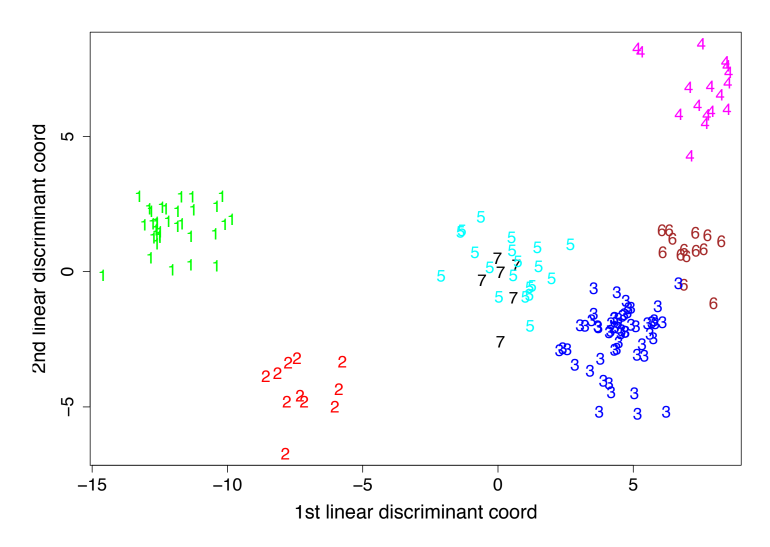
#### **Boxplots of continuous variables**



## Sex by species

	Species							
Sex	Bream	Parkki	Perch	Pike	Roach	Smelt	White	Total
female	3	4	25	5	8	9	1	55
male	6	3	2	1	0	5	0	17
unknown	26	4	29	11	12	0	5	87
Total	35	11	56	17	20	14	6	159

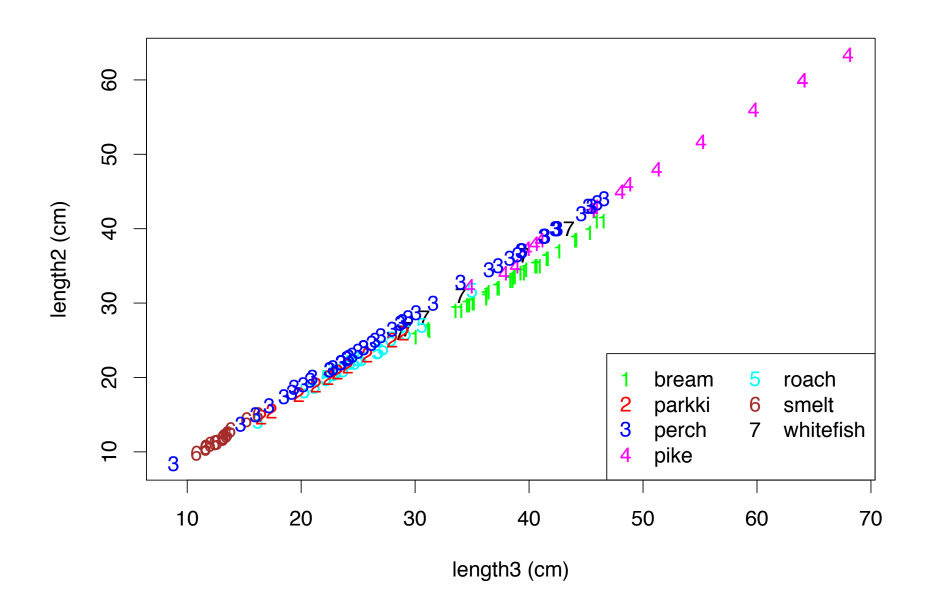
### Linear discriminant analysis



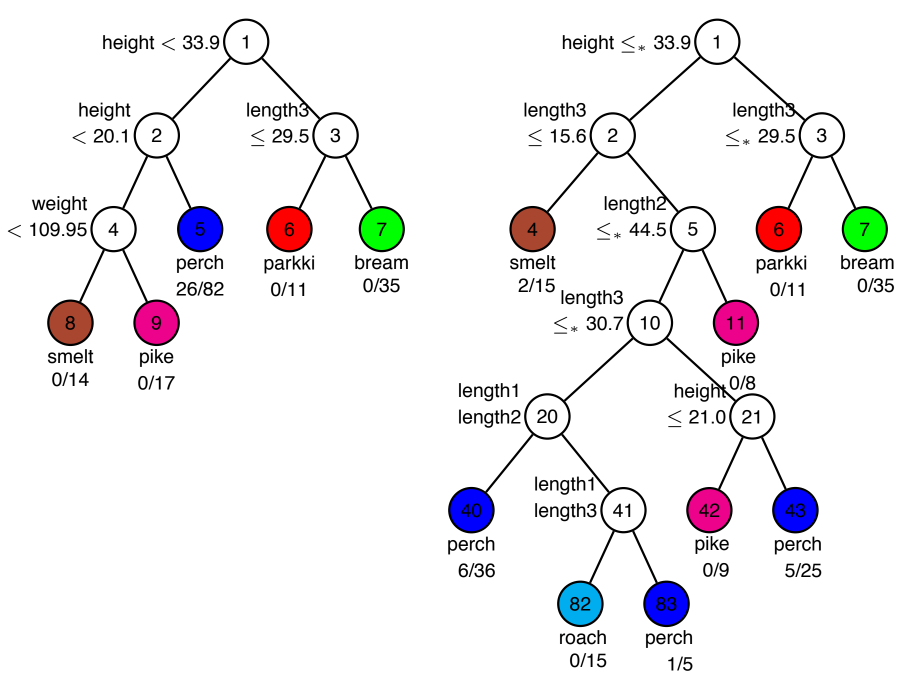
1 = Bream, 2 = Parkki, 3 = Perch, 4 = Pike, 5 = Roach, 6 = Smelt, 7 = Whitefish

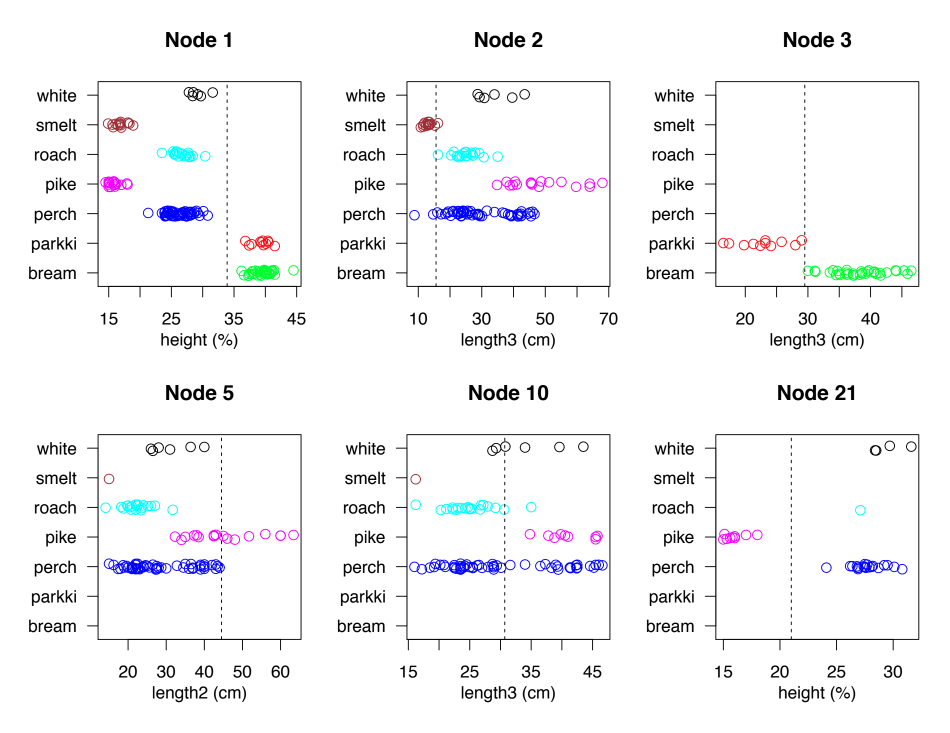
With Sex: 0/71 errors. Without Sex: 1/158 errors

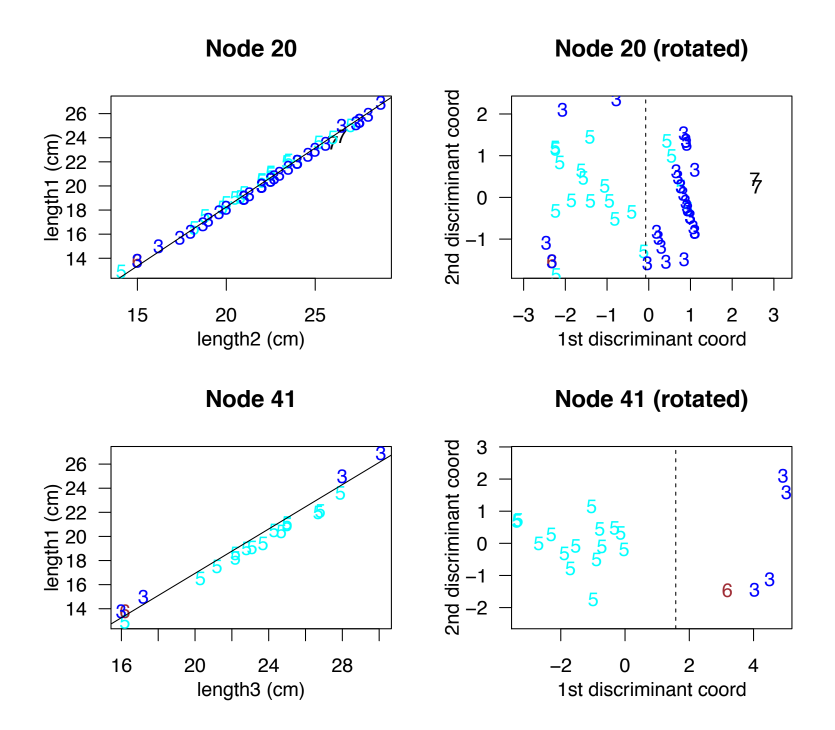
### Plot of Length2 vs. Length3



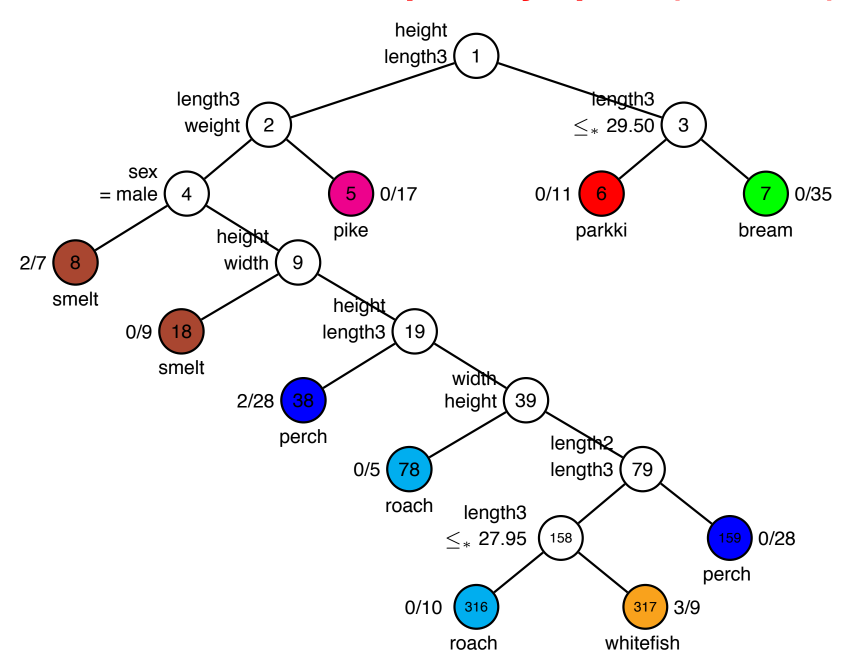
#### RPART (26 errors) and GUIDE (14 errors)

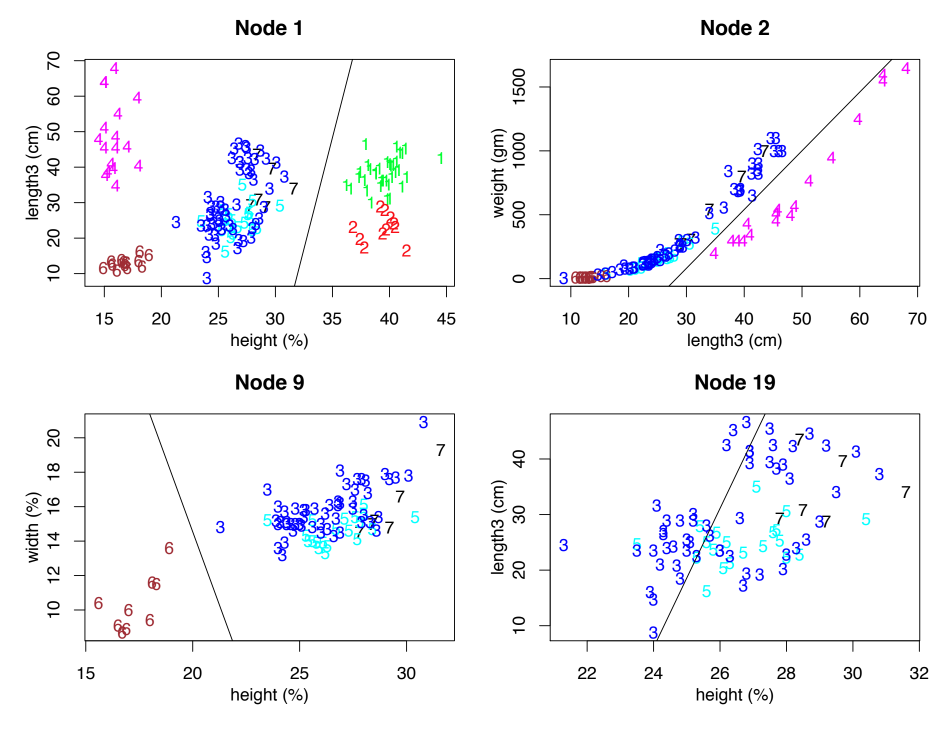


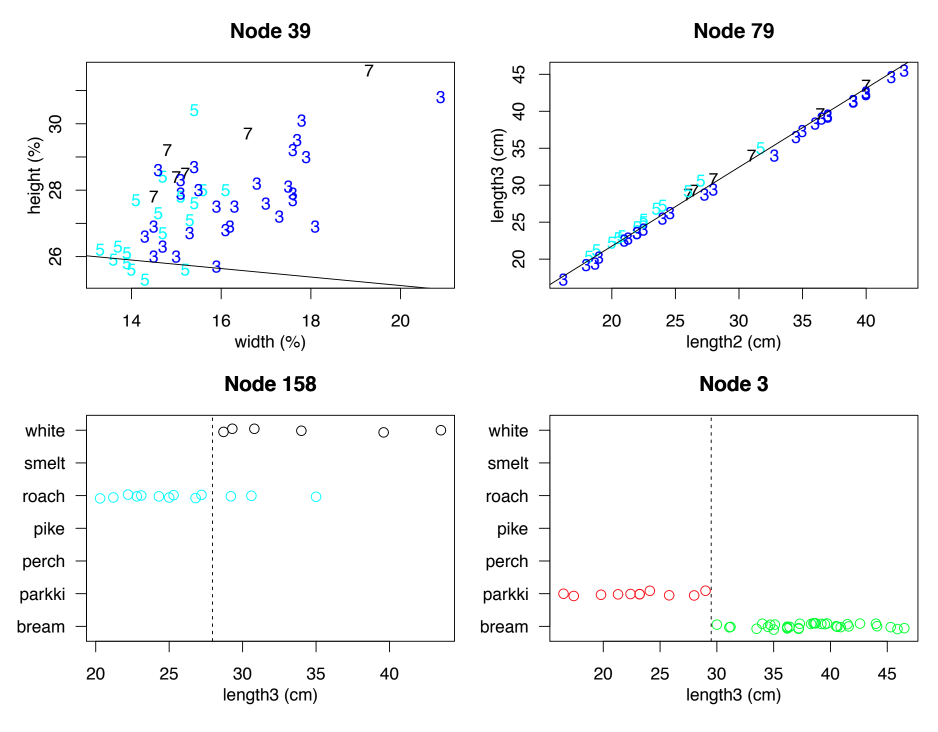




### Fish data with linear priority splits (7 errors)







#### Importance ranking of variables

Importance score of  $X_i$  is

$$\mathsf{IMP}(i) = \sum_{t} n(t) W_M(t, i)$$

- ullet  $W_M(t,i)$  is the Wilson-Hilferty marginal chi-squared value of  $X_i$  at t
- n(t) is the training sample size at node t
- sum is over all intermediate nodes t

If  $X_i$  is constant at t, set  $W_M(t,i) = 1$ 

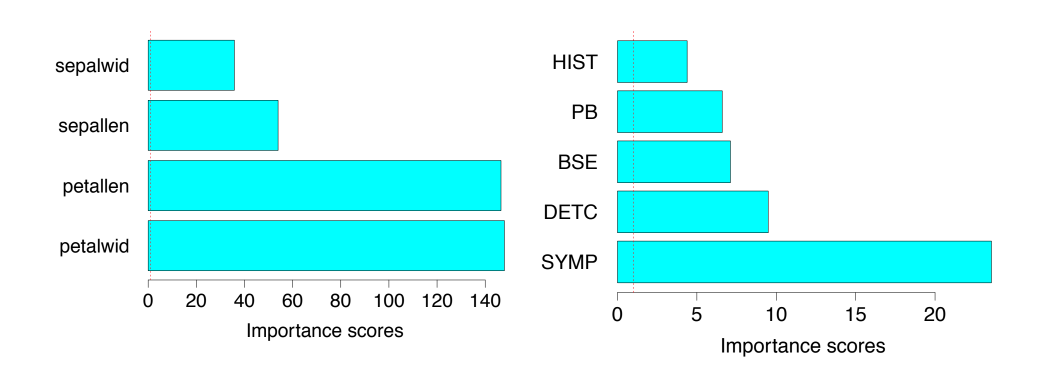
### **Null distribution of importance scores**

- If  $X_i$  is independent of Y, then
  - IMP(i) is a linear combination of independent chi-squared variables
  - Use Satterthwaite (1946) method to approximate distribution of IMP(i)
- Cut-off score for separating important from unimportant variables is the upper- $\alpha$  quantile of the corresponding chi-squared distribution, where

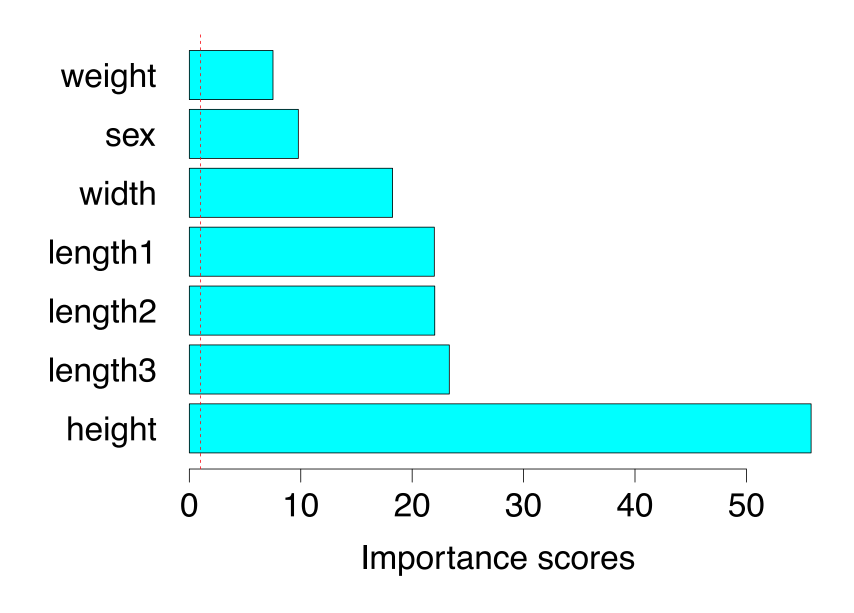
$$\alpha = k_0/K$$

and  $k_0$  is a user-specified expected number of unimportant variables erroneously identified as important (default value of  $k_0$  is 2 for classification and 1 for regression)

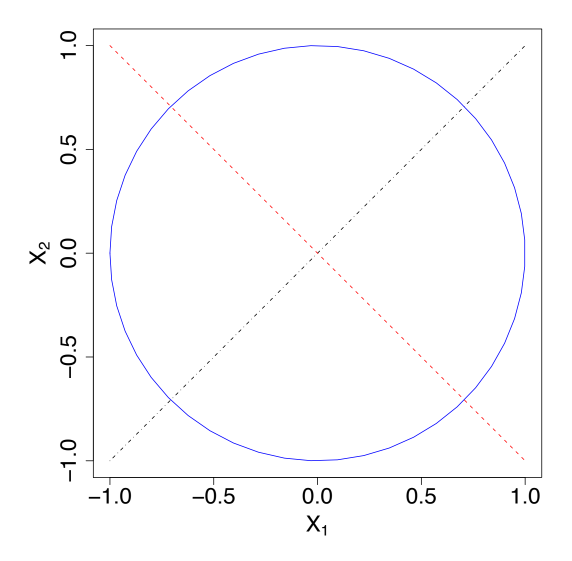
# Importance scores for iris (left) and mammography (right) data



#### Importance scores for fish data



#### A hard three-class problem with 8 predictors



Class 1 dist. on circle, class 2 on one diagonal, class 3 on other diagonal

### Kernel density estimation

- 1. Let s and r be the SD and inter-quartile range of  $x_1, x_2, \ldots, x_n$
- 2. The kernel density estimate is

$$\hat{f}(x) = (nh)^{-1} \sum_{i=1}^{n} \phi\{(x - x_i)/h\}$$

where  $\phi$  is the standard normal density function and h is the bandwidth

$$h = \begin{cases} 2.5 \min(s, 0.7413r) n^{-1/5}, & \text{if } r > 0 \\ 2.5 s n^{-1/5}, & \text{otherwise} \end{cases}$$

#### Kernel node models

#### Let Y denote the class variable

- 1. If the split is due to a marginal chi-squared, let X be the selected variable and fit a kernel density estimate to X for each class in t
- 2. If the split is due to an interaction chi-squared, let  $X_1$  and  $X_2$  be the selected variables. Fit a bivariate density estimate to  $(X_1, X_2)$  for each class in t:
  - (a) If  $X_1$  and  $X_2$  are categorical, use their sample class joint density
  - (b) If  $X_1$  is categorical and  $X_2$  is ordinal, for each combination of  $(X_1, Y)$  values in t, let  $h(Y, X_1)$  be the bandwidth and  $\bar{h}(Y)$  their average. For each value of  $X_1$  and Y, find a kernel density estimate for  $X_2$  using  $\bar{h}(Y)$  as bandwidth.
  - (c) If  $X_1$ ,  $X_2$  are ordinal, fit a bivariate Gaussian kernel density to each class with correlation equal to the class sample correlation

The predicted class is the one with the largest estimated density

#### **Nearest-neighbor node models**

Given n, define  $k = \max(3, \lceil \log n \rceil)$ 

- 1. If the split is due to a marginal chi-squared, let X be the selected variable
  - (a) If X is categorical,  $\hat{Y}$  is the highest probability class among the observations in t with the same X value as the one to be classified
  - (b) If X is ordinal, use k-NN classifier based on X with n = N(t)
- 2. If the split is due to an interaction chi-squared, let  $X_1$  and  $X_2$  be selected
  - (a) If both are categorical,  $\hat{Y}$  is the highest probability class among the cases in t with the same  $(X_1, X_2)$  values as the one to be classified
  - (b) If  $X_1$  is categorical and  $X_2$  is ordinal,  $\hat{Y}$  is given by the k-NN classifier based on  $X_2$  applied to the set S of observations in t that have the same  $X_1$  value as the one to be classified, with n being the size of S
  - (c) If both variables are ordinal, use the bivariate k-NN classifier based on  $(X_1, X_2)$  with the Mahalanobis distance and n = N(t)

#### **GUIDE** treatment of missing values

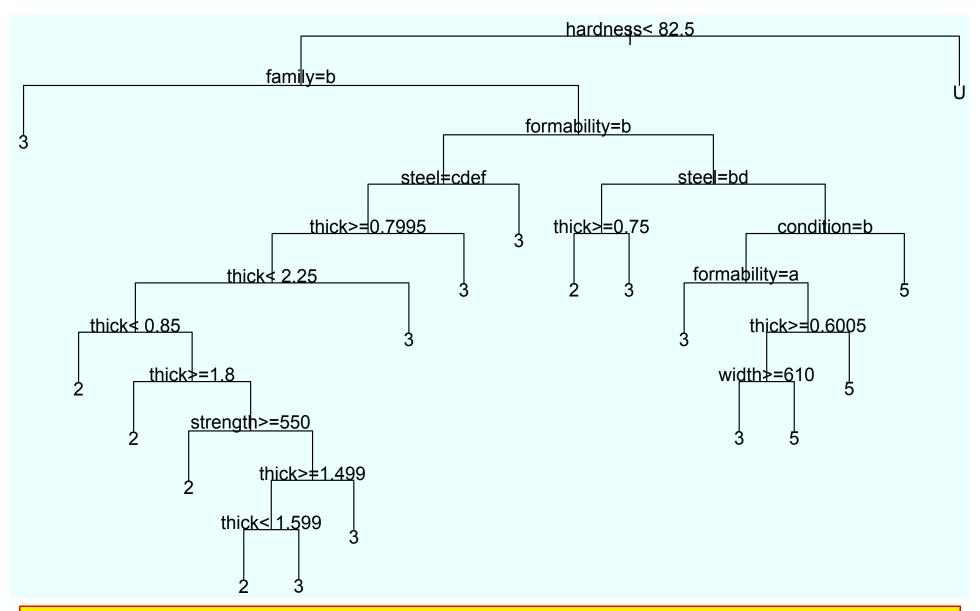
- 1. Cases with missing Y-values are not used for tree construction
- 2. For categorical X, missing values are assigned a separate "missing" category
- 3. For ordinal X:
  - (a) Cases with missing values are assigned to a "missing" interval for selection of split variables
  - (b) A split on missingness is always considered for split point selection
  - (c) Two splits are evaluated for each split on a non-missing value: one for each way of sending the missing values

### **Annealing data: lots of missing values**

Variable	С	M	Variable	С	М	Variable	С	M
class	5		surfacequality	4	217	exptl	1	796
family	2	687	enamelability	2	785	ferro	1	772
steel	7	70	bc	1	797	bright	3	793
carbon	0		bf	1	680	lustre	1	753
hardness	0		bt	1	736	shape	2	
temperrolling	1	675	bwme	2	609	width	0	
condition	2	271	bl	1	662	len	0	
formability	4	283	chrom	1	775	oil	2	740
strength	0		phos	1	791	bore	3	
nonageing	1	703	cbond	1	730	packing	2	789
surfacefinish	1	790	thick	0				

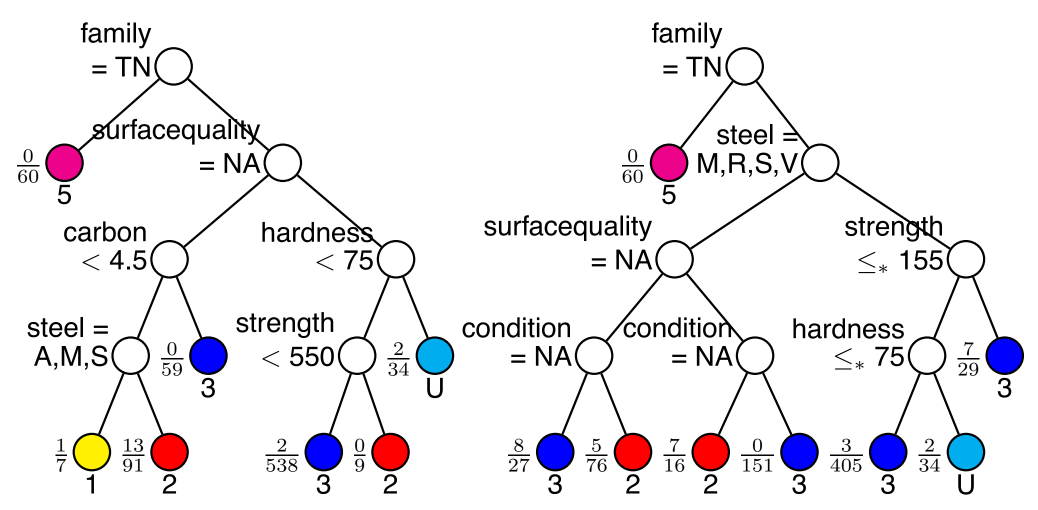
- 798 observations; 6 ordinal and 25 categorical variables
- Cols. C and M give numbers of categories and missing values (o = ordinal) W-Y Loh

### **RPART** tree for annealing data



RPART does not split on missing values and on 1-level categorical variables

# RPART (left, with missing as separate category) and GUIDE (right) trees for annealing data



#Misclassified/sample size beside each node

#### C4.5 (Quinlan, 1993)

- Univariate splits only
- Binary splits on ordered predictors via exhaustive search; splits at data values
- Multiway splits on categorical predictors
  - one subnode for each categorical value (with option to merge categories)
- Pruning based on statistical heuristics; no cross-validation
- Missing values handled by case weights
- Priors and misclassification costs cannot be specified
- Cross-validation error estimate available

# C4.5: Gain ratio split criterion

Define the "info" at node t as the entropy

$$\mathsf{info}(t) = -\sum_{j} p(j|t) \log_2 \{p(j|t)\}$$

• Suppose t is split into subnodes  $t_1, \ldots, t_n$  by predictor X. Define

$$\begin{split} &\inf \mathsf{o}_X(t) &= \sum_i \mathsf{info}(t_i) \frac{N(t_i)}{N(t)} \\ &\operatorname{gain}(X) &= \mathsf{info}(t) - \mathsf{info}_X(t) \\ &\operatorname{split} \mathsf{info}(X) &= -\sum_i \frac{N(t_i)}{N(t)} \log_2 \frac{N(t_i)}{N(t)} \\ &\operatorname{gain} \mathsf{ratio}(X) &= \frac{\mathsf{gain}(X)}{\mathsf{split} \mathsf{info}(X)} \end{split}$$

Split that yields the highest gain ratio is selected

# C4.5: Case weights for missing values

- Initialize the weight for each case to be 1 at the root node
- Suppose t is split by X into subnodes  $t_1, \ldots, t_n$
- Let  $W(t_i)$  be the sum of the weights of cases with known X that land in  $t_i$  and let  $W(t) = \sum_i W(t_i)$
- If a case in learning sample with weight w is missing X, send it down each subnode with weight in  $t_i$  equal to

$$w_i = \frac{W(t_i)}{W(t)} w$$

 Do the same for each test case. If a test case ends up in more than 1 terminal node, assign it the class with largest total weight

# Generalization when there are missing values

• Let  $p_w(j|t) = \frac{\text{sum of class } j \text{ weights in } t}{\text{total weight in } t}$  and define:

$$\begin{split} &\inf (t) &= -\sum_{j} p_w(j|t) \log_2 \{p_w(j|t)\} \\ &\inf (x_i) = -\sum_{j} \inf (t_i) \frac{W(t_i)}{W(t)} \end{split}$$

 Let f be the fraction of learning cases in t that are nonmissing X and define

$$\begin{array}{lcl} \operatorname{gain}(X) & = & f \times \{\operatorname{info}(t) - \operatorname{info}_X(t)\} \\ \operatorname{split} \operatorname{info}(X) & = & -\sum_i \frac{W(t_i)}{W(t)} \log_2 \frac{W(t_i)}{W(t)} - (1-f) \log_2 (1-f) \\ \\ \operatorname{gain} \operatorname{ratio}(X) & = & \frac{\operatorname{gain}(X)}{\operatorname{split} \operatorname{info}(X)} \end{array}$$

# C4.5: Pruning

- Suppose  $N_E(t)$  learning cases are misclassified in node t
- C4.5 estimates the true misclassification probability with the upper 75% confidence bound p where

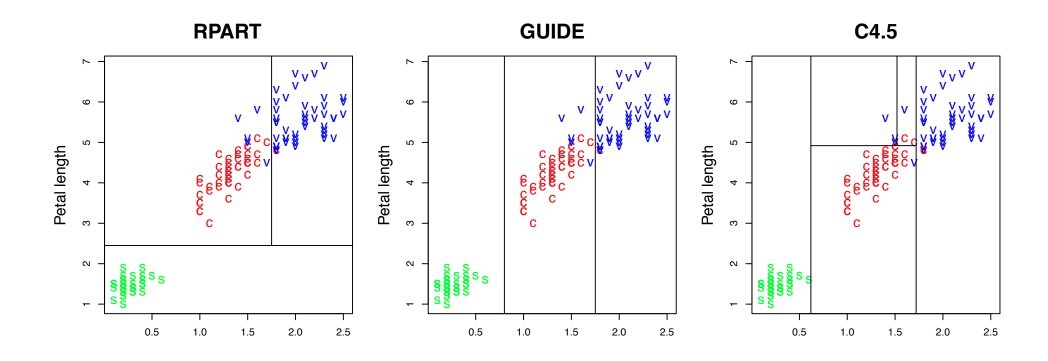
$$\sum_{i=0}^{N_E(t)} \frac{N(t)!}{i! (N(t)-i)!} p^i (1-p)^{N(t)-i} = 0.25$$

• Let  $\nu_1 = 2(N(t) - N_E(t) + 1)$ ,  $\nu_2 = 2N_E(t)$  and  $F_{\nu_1,\nu_2;0.75}$  be the 75% percentile of the  $F_{\nu_1,\nu_2}$  dist. Then (Owen 1962, p. 273)

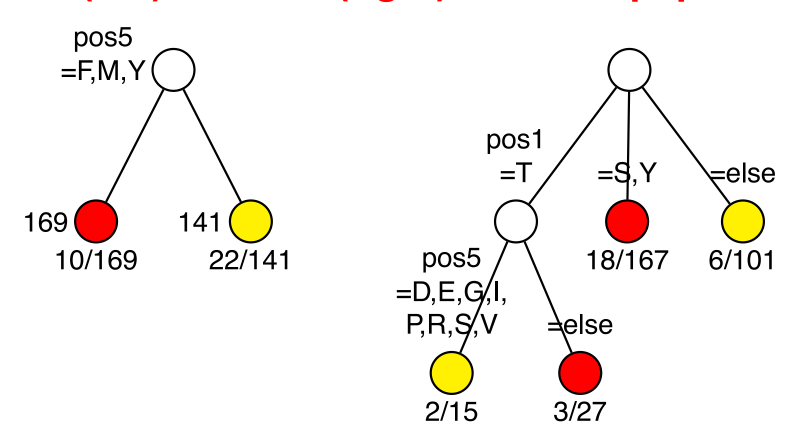
$$p = 1 - \frac{N_E(t)}{N_E(t) + (N(t) - N_E(t) + 1)F_{\nu_1, \nu_2; 0.75}}$$

- The misclassification cost at t is estimated by N(t)p
- A branch is pruned if its estimated cost is larger than its root node

# RPART, GUIDE and C4.5 trees for iris data



## RPART (left) and J48 (right) trees for peptide data

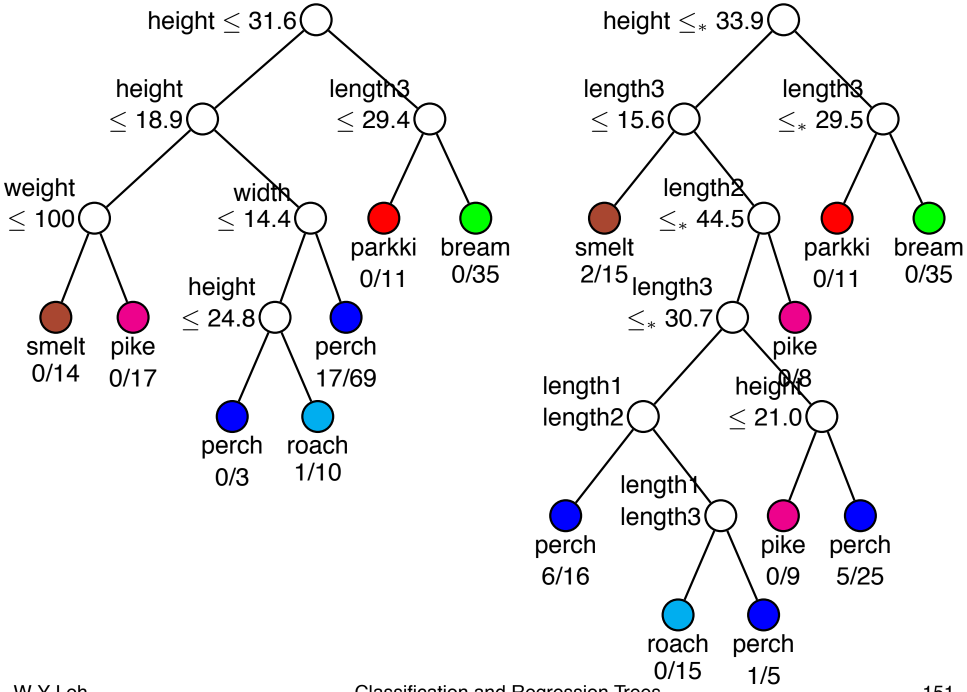


Red denotes binder, yellow denotes non-binder

Numbers beneath nodes are misclassified/sample size

RPART and J48 misclassify 32 and 29 cases, respectively

# J48 (left) and GUIDE (right) trees for fish data



### **CHAID** (Kass, 1980)

- Extends AID to categorical and ordered dependent variables
- Uses a direct stopping rule; no pruning
- Uses significance tests to select split variables and split points
- Uses Bonferroni method to control for multiple testing
- Can split each node into more than two subnodes

# **CHAID** predictor types

**Monotonic:** Ordered or ordinal categorical

Free: Nominal categorical

Floating: Ordinal categorical with exception of a single category that either does not belong to the rest or whose position on the ordinal scale is unknown, e.g., "missing" category

Note: A variable is treated as floating only if it has some missing values in the learning sample. Otherwise it is treated as either monotonic or free. Therefore if a learning sample has no missing values, the tree may not be able to classify future cases that have missing values.

### **CHAID** algorithm

Let  $\alpha_1 > \alpha_2$  and  $\alpha_3$  be three given significance levels.

**Prepare predictors.** Discretize values of each ordinal X into 10 interval groups. For categorical X, the groups are the categories.

#### **Merge categories.** Do for each predictor variable:

- 1. For classification, take each pair of categories and compute the p-value of the chi-squared test of independence between categories and class
- 2. For regression, take each pair of categories and compute the p-value of the two-sample two-sided t-test, using the categories as groups
- 3. Find least significant pair of categories. If  $p > \alpha_1$ , merge the two categories and repeat this step.
- 4. For each compound category containing three or more of the original categories, find the most significant binary split. If  $p < \alpha_2$ , split the compound category and return to Step 3.

**Select split.** Compute Bonferroni-adjusted p-value for each predictor. If smallest  $p < \alpha_3$ , split the node; otherwise stop.

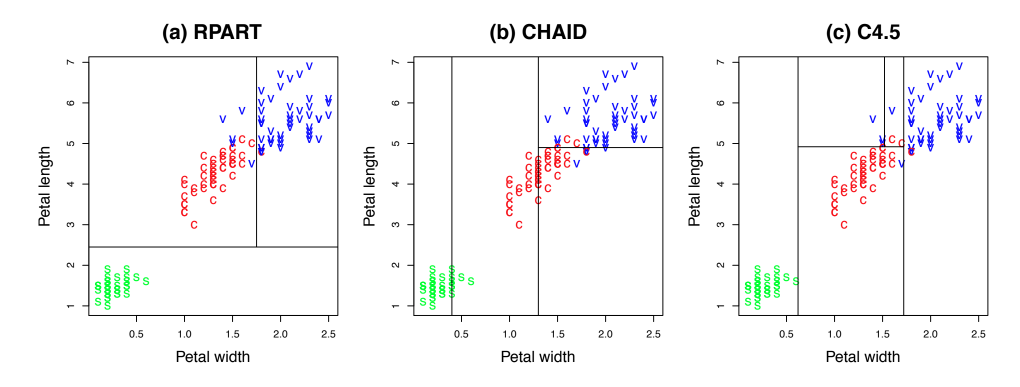
# **CHAID Bonferroni multipliers**

Suppose a predictor with c original categories is merged into r categories. The Bonferroni adjustments to the p-values are:

Monotonic: 
$$B = \begin{pmatrix} c-1 \\ r-1 \end{pmatrix}$$

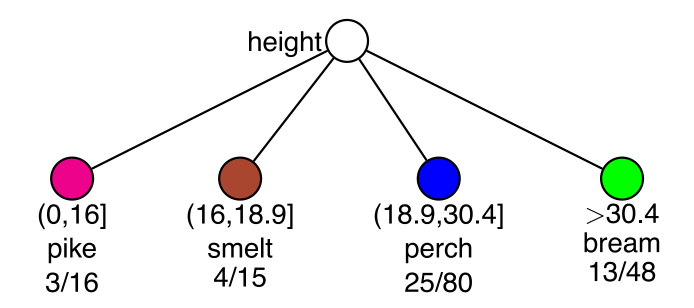
Free: 
$$B = \sum_{i=0}^{r-1} (-1)^i \frac{(r-i)^c}{i!(r-i)!}$$

Floating: 
$$B = {c-2 \choose r-2} + r{c-2 \choose r-1}$$



s = Setosa, c = Versicolour, v = Virginica

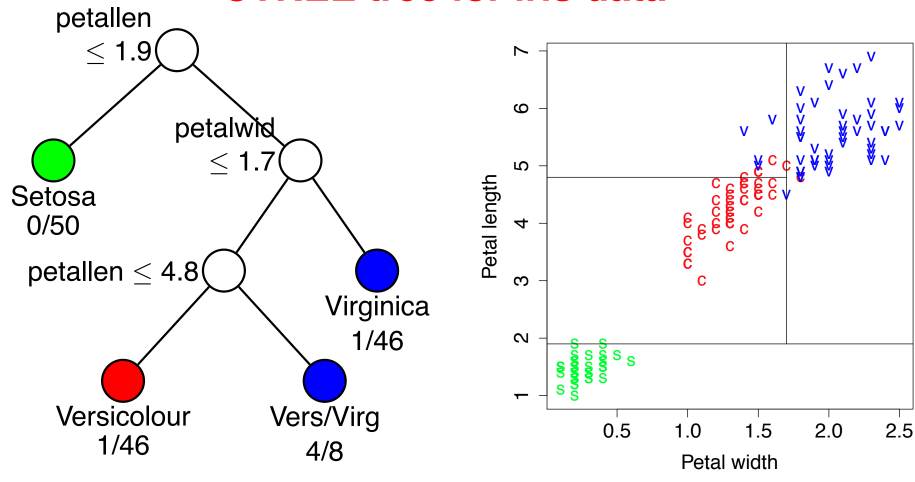
### CHAID tree for fish data (45 misclassified)



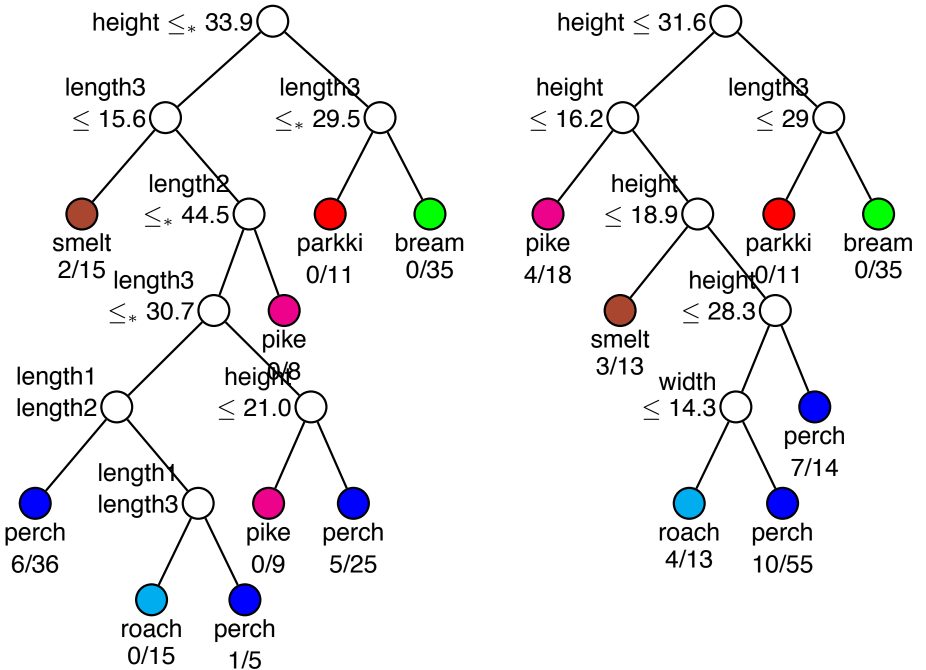
## CTREE (Hothorn et al., 2006)

- 1. Use conditional permutation tests to select variables
  - Requires computation of p-values, either by exact calculation, Monte Carlo simulation, or asymptotic approximation
- 2. Use stopping rules controlled by Bonferroni adjustments; no pruning
- 3. Surrogate splits are used to deal with missing values
- 4. Permutation tests (with subnode label as response variable) are used to find the surrogate variables

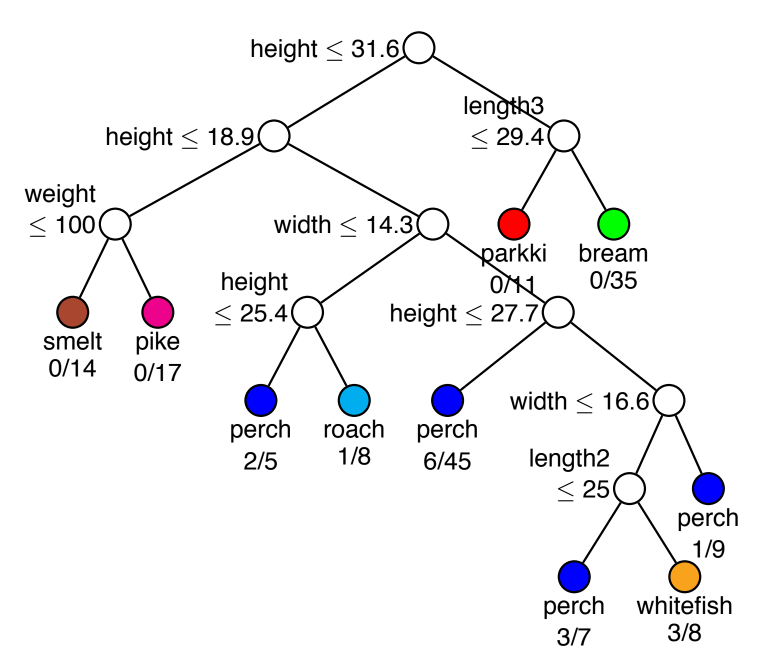
#### **CTREE** tree for iris data



# GUIDE (14 errors) and CTREE (28 errors) for fish data



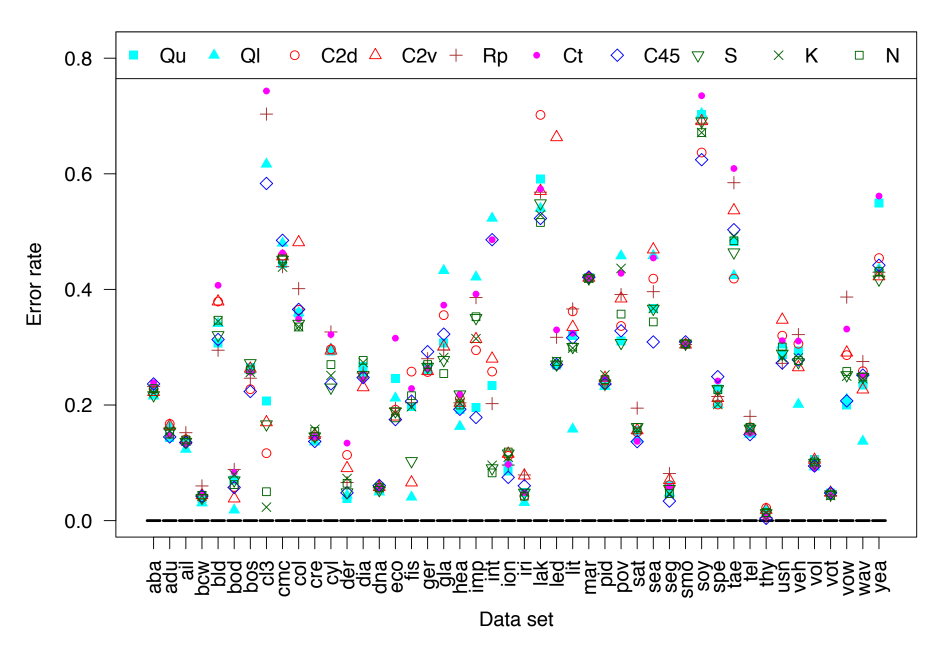
# J48 (16 errors) for fish data



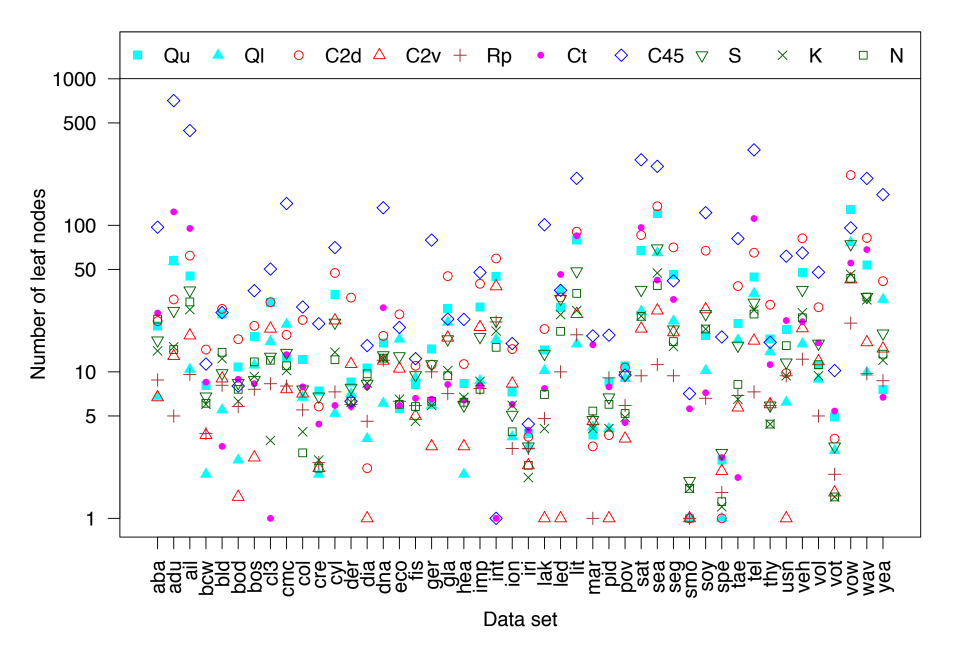
# Comparisons on 46 datasets (Loh, 2009)

C45	C4.5
C2d	CRUISE with interaction detection and simple node models
C2v	CRUISE with interaction detection and linear discriminant node models
Qu	QUEST with univariate splits
QI	QUEST with linear splits
Rp	RPART
Ct	CTree
S	GUIDE with simple node models
K	GUIDE with kernel node models
N	GUIDE with nearest-neighbor node models

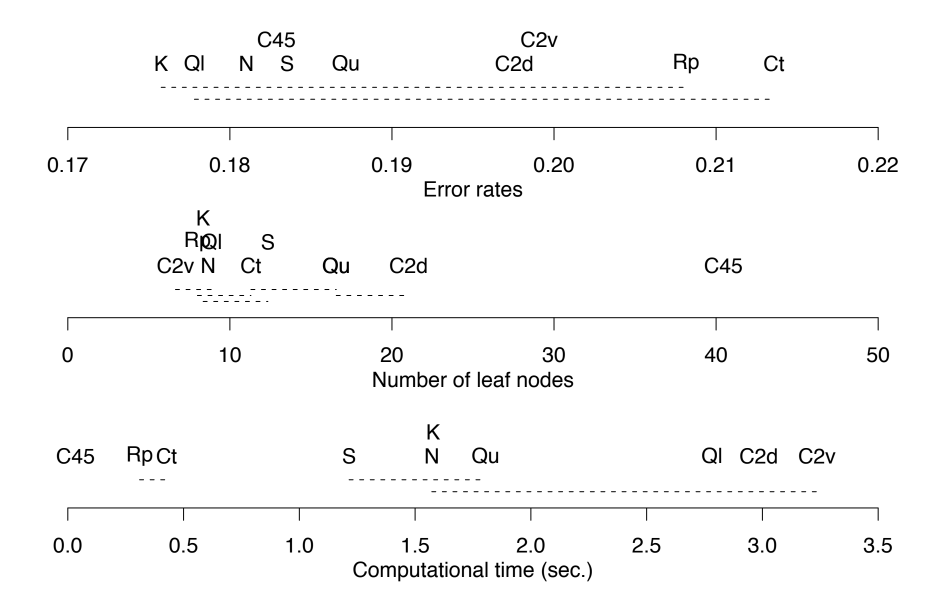
## **Error rates by dataset**



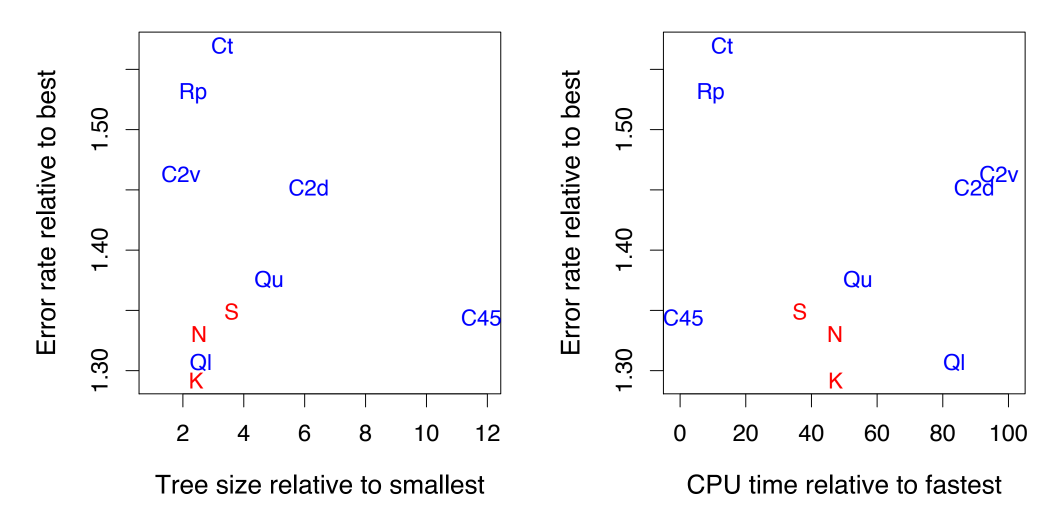
## Number of leaf nodes by dataset



#### **Geometric means over 46 datasets**



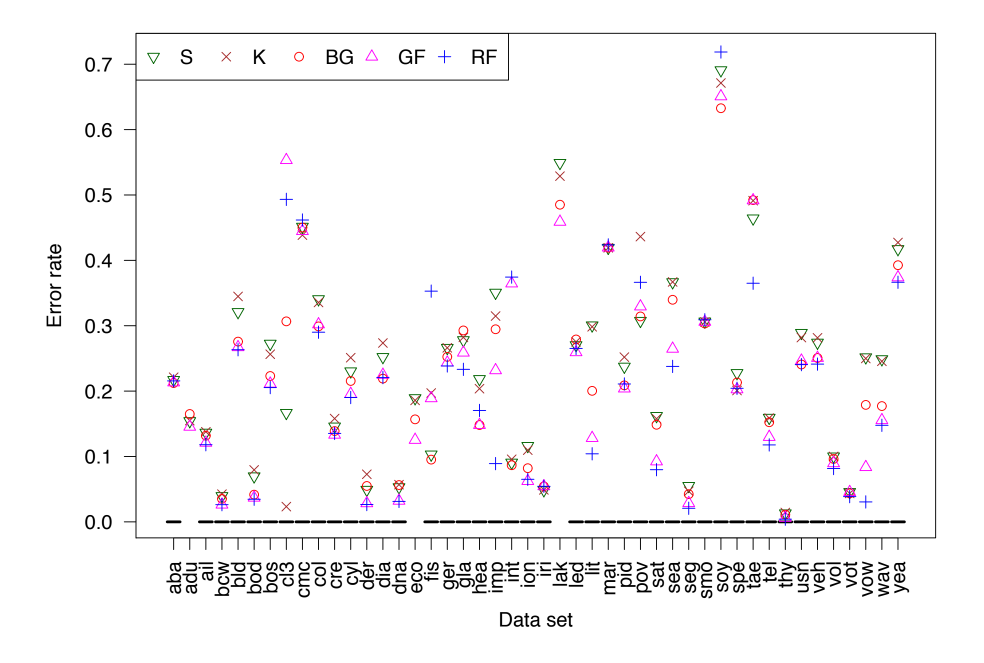
#### Geometric means relative to best for dataset



#### **Tree ensembles**

A tree ensemble uses the majority vote from a collection of tree models to predict the class of an observation

- Bagging (Breiman 1996) creates the ensemble by using bootstrap samples
  of the training data to construct the trees
- Random Forest (RF) employs 500 CART trees, but chooses a random subset of  $\sqrt{K}$  variables to split each node
- Bagged GUIDE (BG) is an ensemble of 100 pruned GUIDE trees, each constructed using the S method from a bootstrap sample
- *GUIDE Forest* (GF) is an ensemble of 500 unpruned GUIDE trees constructed by the S method without interaction and linear splits. As in RF, GF uses a random subset of  $\sqrt{K}$  variables to split each node



### Mean error rates over 43 datasets (Loh, 2009)

Algorithm	S	K	BG	GF	RF
Error rate	0.228	0.231	0.212	0.212	0.206

#### Notes:

- Although the differences in mean error rates are not statistically significant, ensemble methods tend to have 10% or higher higher prediction accuracy than single-tree methods
- RF is inapplicable if categorical variables have more than 32 levels
  - datasets adu and lak have this characteristic
- RF gives an error if the test sample contains class values that do not appear in the training sample
  - dataset eco has this characteristic

## Computational times (sec.) of GUIDE

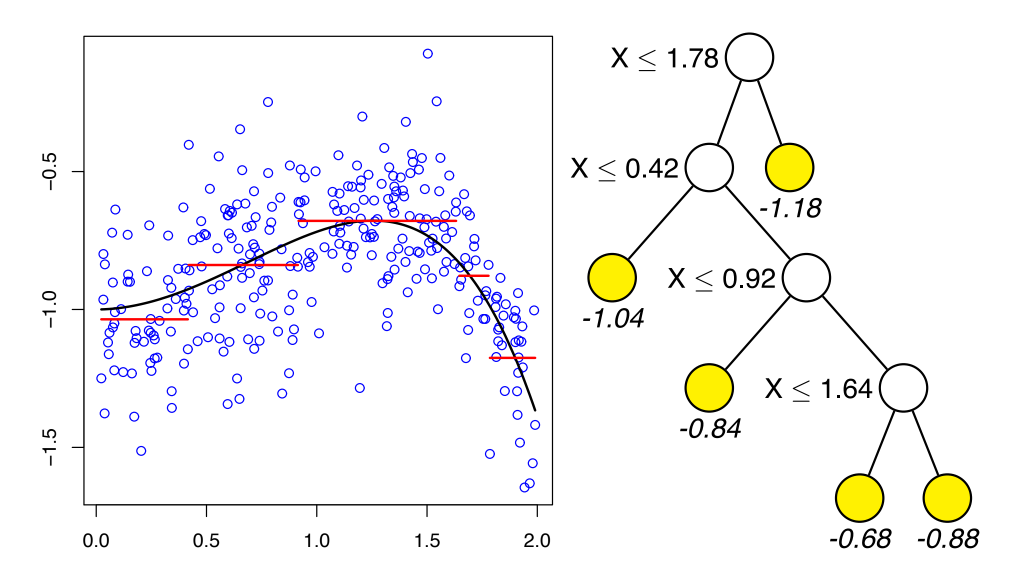
Data	#Obs	#Cat	#Ord	Linux	Win7
Golub	72	0	3,571	2.5	2.8
Adult	32,561	7	6	6.6	7.7
Coil	5,822	2	83	31	36
Arcene	200	0	10,000	71	83
Cover	495,141	2	10	92	106
Gene	1,504	288	17	289	307
Gisette	6,000	0	5,000	403	459
Birthwt	4,268,495	11	12	1933	2198

Computer: 16GB 3.3GHz i3-2120; Compiler: Intel Fortran

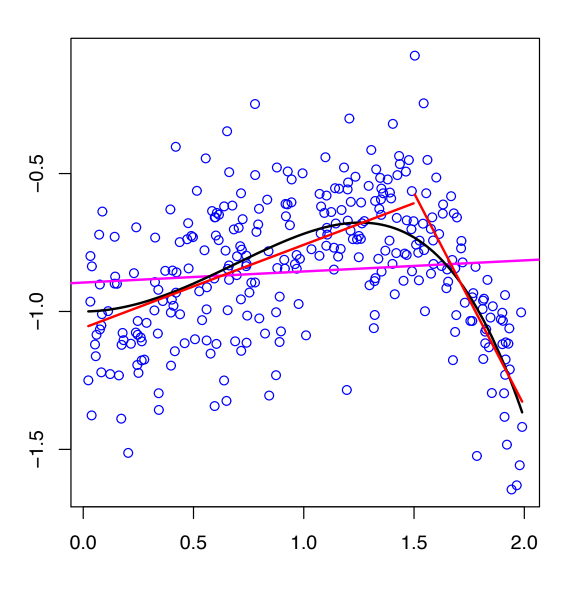
# **CART** regression tree algorithm

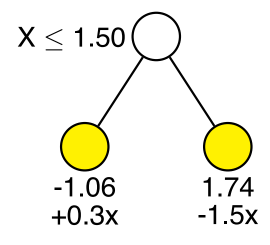
- Fit a constant, the node mean  $\bar{y}$ , at each node
- Use sum of squared deviations  $\sum_i (y_i \bar{y})^2$  as node impurity
- Keep rest of the CART algorithm unchanged

# Piecewise-constant regression model



# Piecewise-linear regression model





# **GUIDE** regression tree models

- Piecewise constant, multiple linear, stepwise linear, best simple polynomial, and best simple ANCOVA
- Least squares, least median of squares, quantile, Poisson, proportional hazards (with censoring), multi-response, and longitudinal data
- Predictor variables can be used for model fitting only, splitting only, or both
- Unbiased variable selection (bootstrap bias correction for linear models)
- Trees pruned with CART method

Ref: Chaudhuri et al. (1994, 1995); Chaudhuri and Loh (2002); Loh (2002, 2006, 2008b); Loh and Zheng (2013)

## Variable roles in GUIDE description files

D: Dependent variable (least-squares, least median of squares, quantile, Poisson, multi-response and longitudinal) or death indicator (proportional hazards)

N: Numerically ordered variable used for fitting and splitting

F: Numerically ordered variable used for fitting only

S: Numerically ordered variable used for splitting only

C: Categorical variable used for splitting only

B: Categorical variable for both for splitting and fitting via dummies

R: Treatment categorical variable for fitting only

W: Weight variable for weighted least squares and case exclusion

T: Survival or observation time (prop. hazards or longitudinal data)

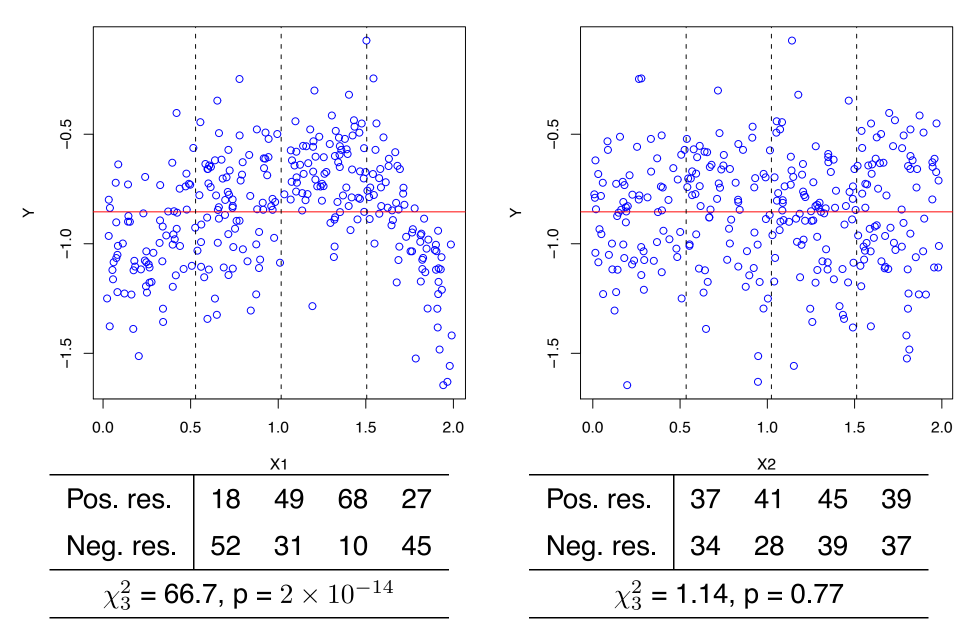
**Z:** Offset variable (Poisson regression)

X: Excluded variable

# GUIDE variable selection for regression

- 1. Fit a model to the data in the node and obtain the residuals
- 2. Define a "class" variable that equals +1 if residual is positive, -1 otherwise
- 3. Follow GUIDE classification procedure to select a variable to split node

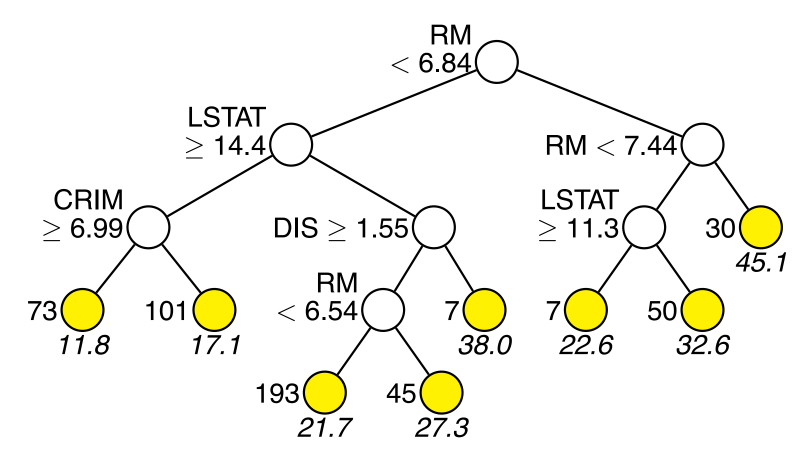
## Split variable selection based on residual patterns



# Selection bias: Boston housing data

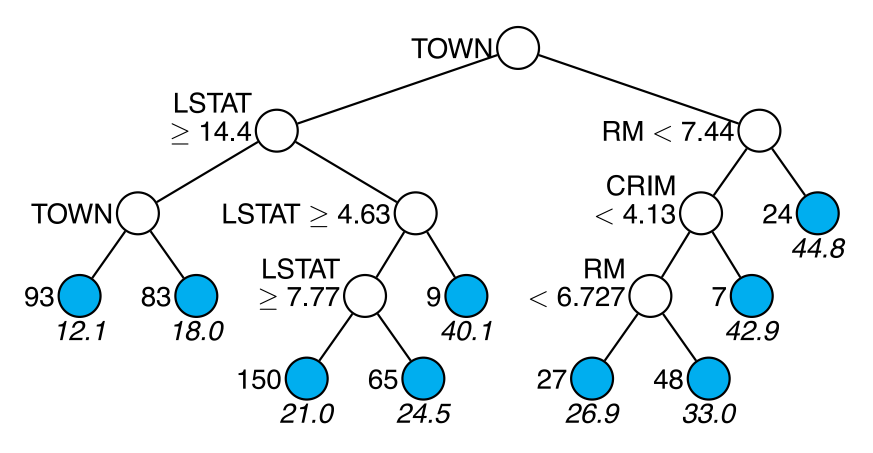
- Categorical variable TOWN has 92 values
- If TOWN is included, RPART has a high chance to select it
  - actually, RPART can search over at most 32 categorical values
  - it is unclear how it deals with TOWN
- GUIDE is much less influenced by the presence of TOWN

#### **RPART tree for MEDV without TOWN**



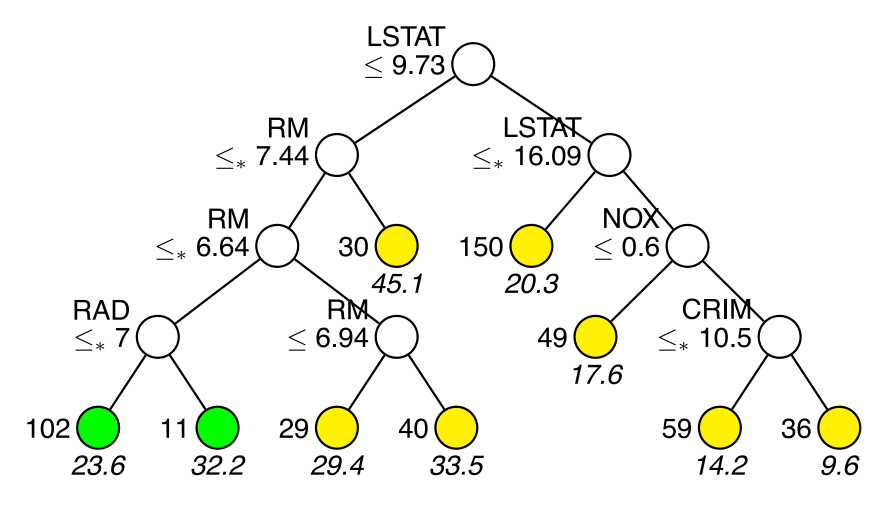
Predicted MEDV values beneath terminal nodes; sample sizes on left

#### **RPART tree for MEDV with TOWN**



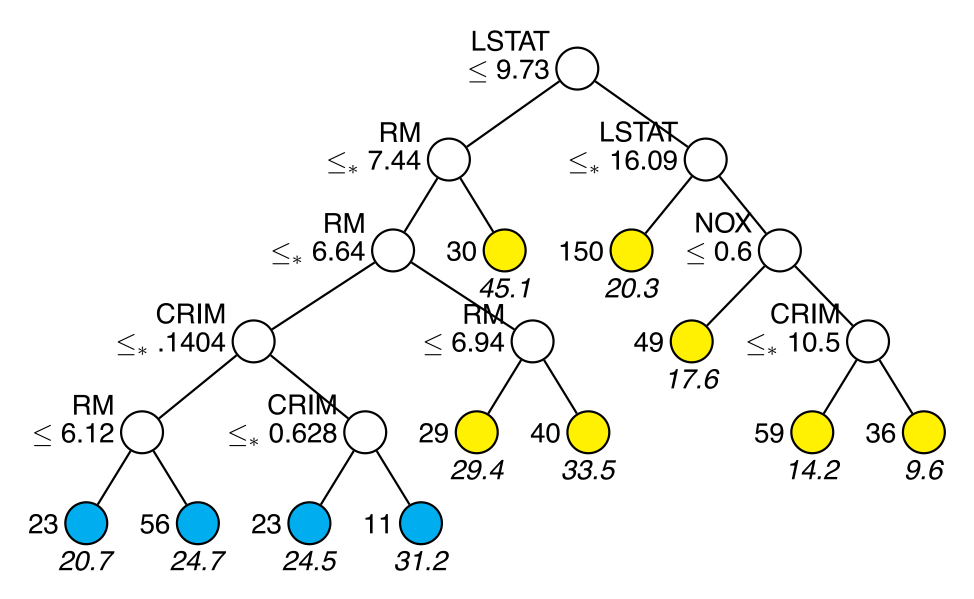
Predicted MEDV values beneath terminal nodes; sample sizes on left

#### **GUIDE tree for MEDV without TOWN**



Predicted MEDV values beneath terminal nodes; sample sizes on left

#### **GUIDE tree for MEDV with TOWN**



Predicted MEDV values beneath terminal nodes; sample sizes on left

# Converting categorical variables to dummy variables is undesirable

- 1. Transform each X into a 0-1 dummy vector  $(U_1, \ldots, U_c)$
- 2. Use  $U_1, \ldots, U_c$  as predictors in model
- 3. Resulting ANCOVA model
  - (a) uses up many degrees of freedom if c is large
  - (b) forces all non-categorical predictors to have constant slope coefficients for all values of categorical predictors
- 4. Splits of the form  $U_i = 0$  vs.  $U_i = 1$  translates to unappealing "singleton" splits of the form X = a vs.  $X \neq a$

#### GUIDE treatment of categorical predictors

- Categorical predictors can be used for splitting and/or model fitting;
   ANCOVA models tend to yield shorter trees
- Splits are on subsets of categories

#### Naive variable selection for piecewise-linear model

- 1. Fit a linear model to the **n** and **f**-variables in the node and obtain residuals
- 2. For each **s** and **n**-variable X:
  - (a) Divide cases into three or four groups
  - (b) Cross-tab data with signs of residuals as rows and groups as columns
  - (c) Compute a Wilson-Hilferty  $\chi_1^2$ -value
- 3. Do the same for each **c**-variable, using categories to form columns of table
- 4. Select the variable with the largest  $\chi_1^2$  value

#### Selection bias in linear fit

- Residuals uncorrelated with n-predictors, but not with c and s-variables
- $\chi^2$  tests for **n**-variables are less significant than those for **c** and **s**-variables

### Simulation experiment

Predictors	Independent	Weakly dependent	Strongly dependent
$X_1$	T	T	T
$X_2$	W	W	W
$X_3$	Z	T + W + Z	W + 0.1Z
$X_4$	$C_5$	$\lfloor UC_{10}/2 \rfloor + 1$	$\lfloor UC_{10}/2 \rfloor + 1$
$X_5$	$C_{10}$	$C_{10}$	$C_{10}$

- $C_k$  is k-category taking values  $\{1, 2, \dots, k\}$  with equal probabilities
- T is non-categorical uniformly distributed variable on  $\{\pm 1, \pm 3\}$
- ullet U is uniform U(0,1); W is exponential with mean 1; Z is N(0,1)
- $C_k$ , U, T, W, and Z are mutually independent
- [.] is the greatest integer function

# Selection probabilities for piecewise linear model when Y is independent of predictors

		Independent $X_i$		Weakly depend. $X_i$		Strongly depend. $X_i$	
$X_{i}$	Type	Uncorr.	Corr.	Uncorr.	Corr.	Uncorr.	Corr.
$X_1$	n	0	.202	0	.181	0	.197
$X_2$	n	0	.217	0	.228	0	.214
$X_3$	S	.352	.203	.288	.134	.313	.121
$X_4$	C	.307	.178	.360	.238	.360	.256
$X_5$	C	.341	.200	.352	.219	.327	.212

### Bootstrap bias correction: basic idea

- Since chi-squared values of **n**-variables are stochastically smaller, scale them with a multiple  $\gamma>1$
- Estimate  $\gamma$  with the bootstrap: randomly permute the Y values and find the  $\gamma$  that yields equal selection probabilities

## **GUIDE** regression in a nutshell

- 1. Fit a model to the node and use residual signs to form two classes
- 2. Apply GUIDE classification to select a variable to split node
- 3. If selected variable is due to a marginal test:
  - X is n or s: Search all splits of form  $X \leq c$  to minimize sum of deviances X is b or c:
    - (a) If 9 or fewer unique X values, search exhaustively
    - (b) Otherwise apply GUIDE classification to the two-class problem
- 4. If selected variables are due to an interaction test, use GUIDE classification to select variable and split set
  - See p.118 for Steps 3(b) and 4.

### GUIDE approach to missing values for regression

- 1. A "missing" category is created for each categorical variable for split selection
- 2. For each split on an ordered variable, missing values are sent to the left or right node, depending on which one reduces node impurity more. The split that sends all missing values to one node and all nonmissing to the other is also considered.
- 3. For piecewise constant models, only cases complete in the **d**, **w**, **t**, and **z** variables are used for split selection and model fitting
- 4. For all other models, fitting is restricted to cases complete in the  $\mathbf{n}$  and  $\mathbf{f}$  variables; the node Y mean is fitted to the other cases
- 5. Bootstrap bias-correction is performed for multiple linear models only

# Quantile regression example: Which colleges are the most expensive?

- Data on 1134 U.S. colleges and universities for year 1995 from *U. S. News* & World Report (http://lib.stat.cmu.edu/)
- Response variable is out-of-state tuition
- Goal: Identify the top 10% most expensive colleges, after allowing for various explanatory variables

# **Explanatory variables for college data**

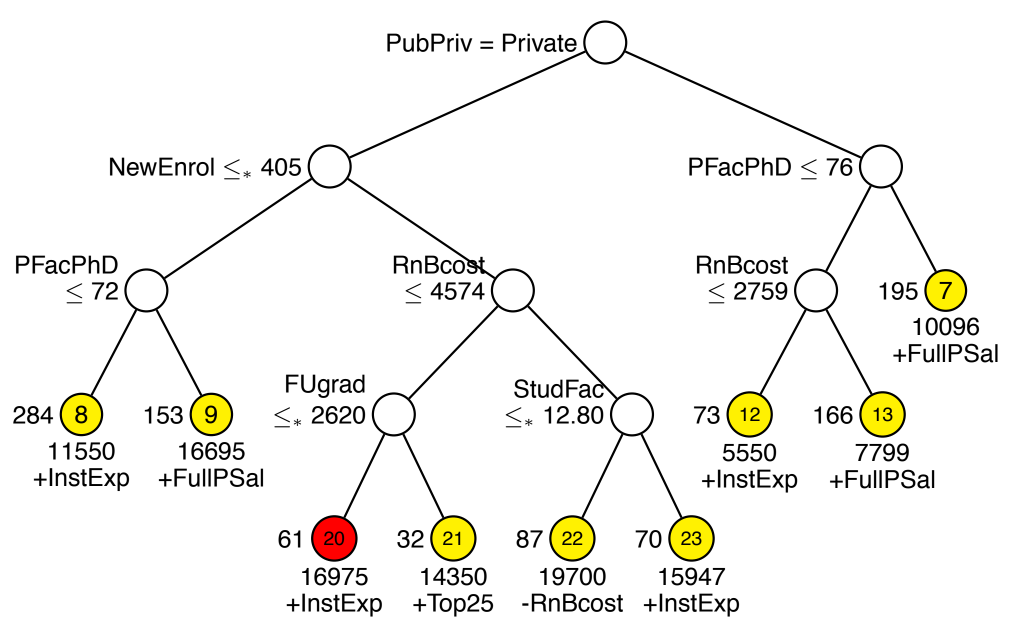
Name	Description	#Missing
PubPriv	Public or private college (binary)	0
CombSAT	Average Combined SAT score	471
AppsRec	Number of applications received	9
AppsAcc	Number of applicants accepted	9
NewEnrol	Number of new students enrolled	5
Top10	Percent new students from top 10% of H.S. class	183
Top25	Percent new students from top 25% of H.S. class	155
FUgrad	Number of fulltime undergraduates	3

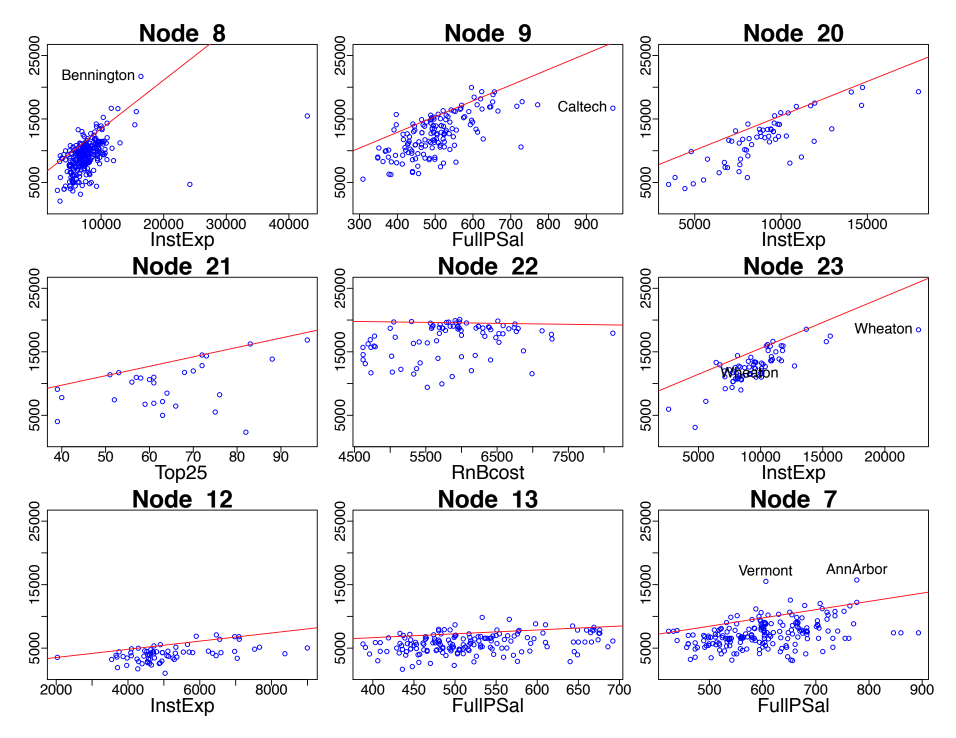
### Explanatory variables for college data (cont'd)

Name	Description	#Missing
RnBcost	Room and board costs	57
PFacPhD	Percent of faculty with Ph.D.'s	29
StudFac	Student/faculty ratio	2
InstExp	Instructional expenditure per student	24
GradRate	Graduation rate	69
Type	College type (I: doctoral, IIA: master, or IIB: bachelor)	0
FullPSal	Average salary—full professors (in \$100's)	61
NFullProf	Number of full professors	0

513 cases with complete observations

# GUIDE simple linear 90th-percentile tree for out-of-state tuition





# Subgroup identification for differential treatment effects: an approach to personalized medicine

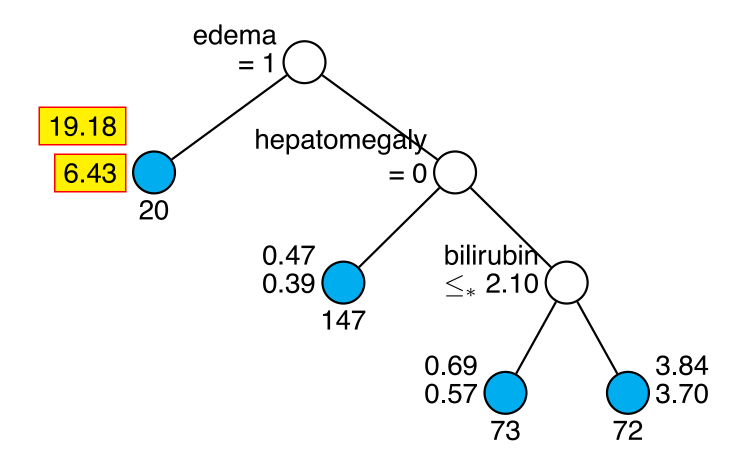
- A piecewise linear model is required for detection of treatment effects
- Piecewise constant models are ineffective because splitting on the treatment variable is useless
- Solution: use the treatment variable as the only linear predictor (after converting to dummy vector)
- Use all other variables for splitting
- Ref: Loh et al. (2014)

# Example: primary biliary cirrhosis (PBC) of the liver (Fleming and Harrington, 2005)

- Randomized placebo controlled trial for the drug D-penicillamine
- 312 PBC patients, referred to Mayo Clinic during 1974–84
- Response variable is number of days between registration and the earlier of death, liver transplantation, or study analysis time in July, 1986

1	age	days
2	sex	0=male, 1=female
3	presence of ascites	0=no 1=yes
4	presence of hepatomegaly	0=no 1=yes
5	presence of spiders	0=no 1=yes
		0=no edema and no diuretic therapy for edema
6	presence of edema	0.5 = edema present w/o diuretics, or edema resolved by diuretics
		1 = edema despite diuretic therapy
7	serum bilirubin	mg/dl
8	serum cholesterol	mg/dl
9	albumin	gm/dl
10	urine copper	ug/day
11	alkaline phosphatase	U/liter
12	SGOT	U/ml
13	triglicerides	amg/dl
14	platelets	per cubic ml / 1000
15	prothrombin	time in seconds
16	histologic stage of disease	1, 2, 3, 4, 5

#### **GUIDE** model for differential treatment effects



- Relative risks of death (drug, upper; placebo, lower) on left of nodes
- Sample sizes beneath nodes

## Longitudinal data (Loh and Zheng, 2013)

- 1. Treat each data point as a curve (trajectory)
- 2. Fit a mean curve (lowess or smoothing spline) to data in the node
- 3. Group trajectories into classes according to shapes relative to mean curve
- 4. For each X variable, find p-value of chi-squared test of class vs. X
- 5. Select X with smallest p-value to split node
- 6. For each split point, fit a mean curve to each child node
- 7. Select the split that minimizes sum of squared deviations (normalized if desired) of trajectories from mean curves in two child nodes
- 8. Stop splitting when sample size in node is too small
- 9. Prune the tree using cross-validation

### **Example: CD4 counts from an AIDS study**

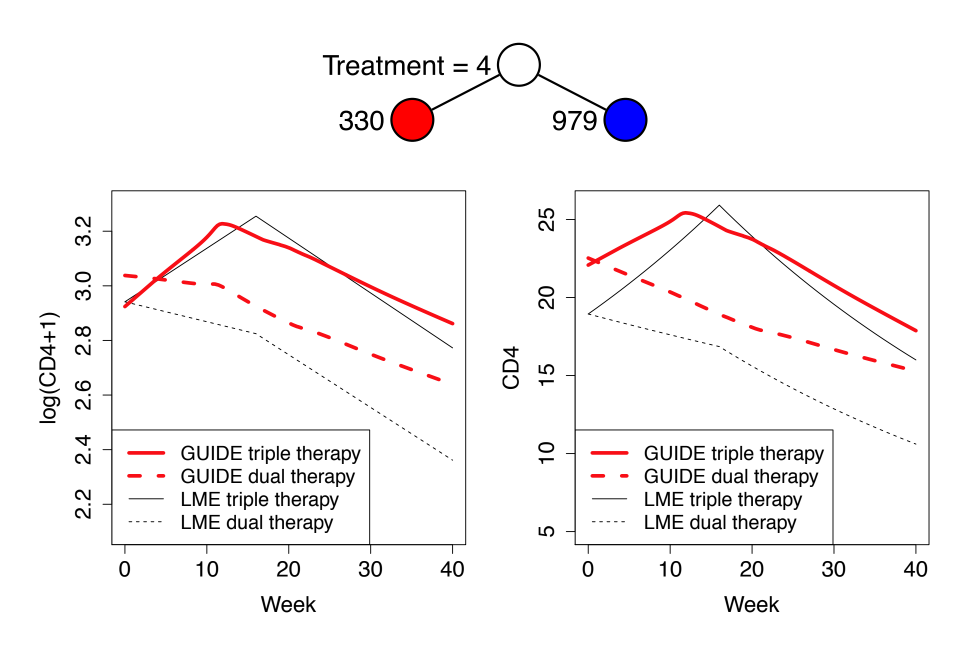
- Randomized, double-blind, study of 1309 AIDS patients with advanced immune suppression (Fitzmaurice et al., 2004)
- Four dual or triple combinations of HIV-1 reverse transcriptase inhibitors:
  - **1:** 600mg *zidovudine* alternating monthly with 400mg *didanosine* (dual therapy)
  - 2: 600mg zidovudine + 2.25mg zalcitabine (dual therapy)
  - **3:** 600mg *zidovudine* + 400mg *didanosine* (dual therapy)
  - **4:** 600mg *zidovudine* + 400mg *didanosine* + 400mg *nevirapine* (triple therapy)
- CD4 counts at baseline and at 8-week intervals during 40-week follow-up
- Observations during follow-up varied from 1–9, with median of 4 due to:
   (i) mistiming and (ii) missingness from skipped visits and dropout
- Response variable is log(CD4 counts + 1); covariates are age and gender

### Fitzmaurice et al. (2004) linear mixed effects model

$$E(Y_{ij} | b_i) = \beta_1 + \beta_2 t_{ij} + \beta_3 (t_{ij} - 16)_+ + \beta_4 I(\mathsf{Trt} = 4) \times t_{ij}$$
  
+  $\beta_5 I(\mathsf{Trt} = 4) \times (t_{ij} - 16)_+ + b_{1i} + b_{2i} t_{ij} + b_{3i} (t_{ij} - 16)_+$ 

- 1.  $Y_{ij} = \log(CD4_{ij} + 1)$  for subject i at time  $t_{ij}$
- 2. All fixed effects significant (p < 0.005)
- 3. Significant difference in rate of change from baseline to week 16 between dual and triple therapies
- 4. No sig. difference in rate of change from week 16 to 40 between groups
- 5. Substantial within and between-patient variability (large random effects)

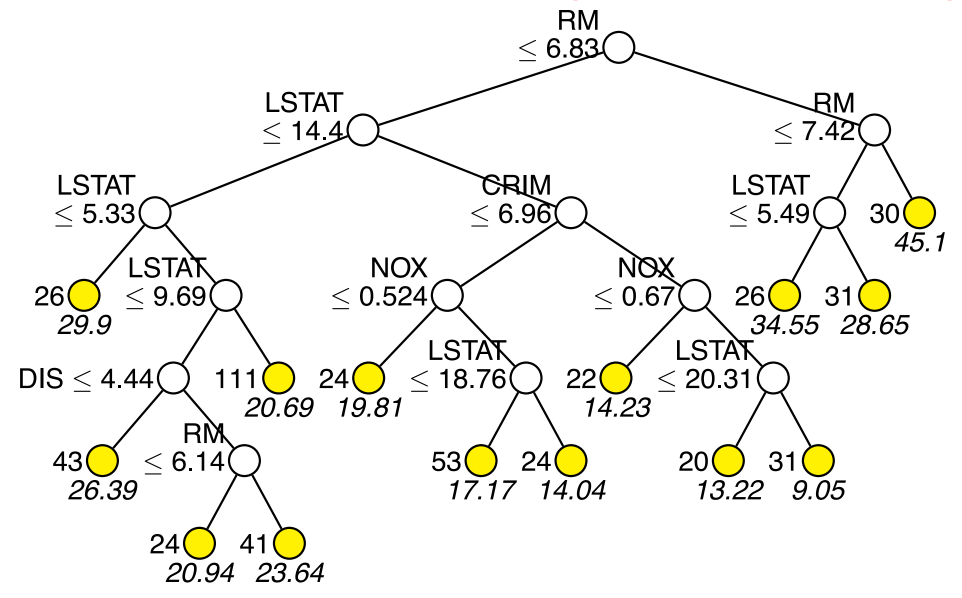
#### **GUIDE** regression tree for AIDS data



# MOB: Model-based recursive partitioning (Zeileis et al., 2008)

- 1. Fit a model once to data in the current node.
- 2. Assess whether parameter estimates are stable with respect to each split variable, using Bonferroni-adjusted p-values.
- 3. If minimum p-value is sufficiently small, select the most unstable variable and split the node into two. Otherwise stop.

### MOB for MEDV without TOWN (crashes with TOWN)



Predicted MEDV values beneath terminal nodes; sample sizes on left

### M5 regression tree (Quinlan, 1992)

1. Grow large tree: Grow a piecewise-constant tree using as reduction in error

$$\frac{m}{n} \left\{ SD(t) - \frac{n_L SD(t_L) + n_R SD(t_R)}{n} \right\}$$

where node t (with sample size n) is split into  $t_L$  and  $t_R$  (with sample sizes  $n_L$ ,  $n_R$ ), SD(t) is the sample standard deviation of the cases in t, and m is the number of non-missing values in the split variable

- 2. Fit linear models: After the tree is grown, fit a multiple linear regression model to the cases in each node t, using as regressors only the variables that are selected for splitting in subtree  $T_t$
- 3. Estimate error: Estimate the prediction error of each node t with

$$\mathsf{Err}(t) = \frac{\sum_{i} |y_i - \hat{y}_i|}{n} \times \frac{n + \nu}{n - \nu}$$

where  $\nu$  is the number of fitted parameters and n is the sample size in t

4. Simplify linear models: Use backward stepwise regression to reduce the number of regressors in each node

- 5. Prune tree: Starting from the bottom, remove branch  $T_t$  if  $Err(T_t) \ge Err(t)$
- 6. Smooth predicted values: Let  $t^*$  be the parent node of t. Given a case, let its predicted value at t and  $t^*$  be  $\hat{y}$  and  $\hat{y}^*$ . The smoothed predicted value is

$$\hat{y}^{**} = (n\hat{y} + k\hat{y}^*)/(n+k)$$

where k is a constant (default value 15). Repeat all the way up to root node.

### Categorical predictors in M5

- Each categorical variable is converted to a vector of binary variables
- Suppose categorical variable X takes values  $X_1, X_2, \dots, X_c$ .
  - 1. Order the X values by their sample mean Y-values
  - 2. Denote the ordered values by  $X'_1, X'_2, \dots, X'_c$
  - 3. Create binary variables  $U_1, U_2, \ldots, U_{c-1}$  such that

$$U_i = \begin{cases} 0 & \text{if } X \in \{X_1', \dots, X_i'\} \\ 1 & \text{otherwise} \end{cases}$$

- 4. Replace X by  $(U_1, U_2, \dots, U_{c-1})$
- The conversion is usually carried out only at the root node

### Missing values in M5

**Training data:** Use the Y variable to form a surrogate split: Compare the Y-value of the observation with the mean of the Y-values in the two subnodes

**Test data:** Replace missing values with means from the training sample in the node

M5 is implemented in Witten et al. (2011) as M5'

# Empirical comparison of regression algorithms (Loh et al., 2007)

#### 15 algorithms

- 10 regression tree methods
- 3 ensemble (bagged) methods
- 2 spline methods

#### 52 datasets

- Training sample size from 96 to 21,252
- Number of ordered predictor variables from 1 to 28
- Number of categorical variables from 0 to 6
- Number of variables in model fitting from 3 to 104

### 15 regression algorithms

GUIDE piecewise simple linear (G1)

GUIDE piecewise simple quadratic (G2)

GUIDE piecewise simple cubic (G3)

GUIDE piecewise multiple linear (Gm)

GUIDE piecewise stepwise linear (Gs)

GUIDE stepwise pairs (Gp)

GUIDE simple ancova (Ga)

Generalized additive model (gam)

Multivariate adaptive splines (mars)

M5 piecewise constant (mc)

M5 piecewise multiple linear (mm)

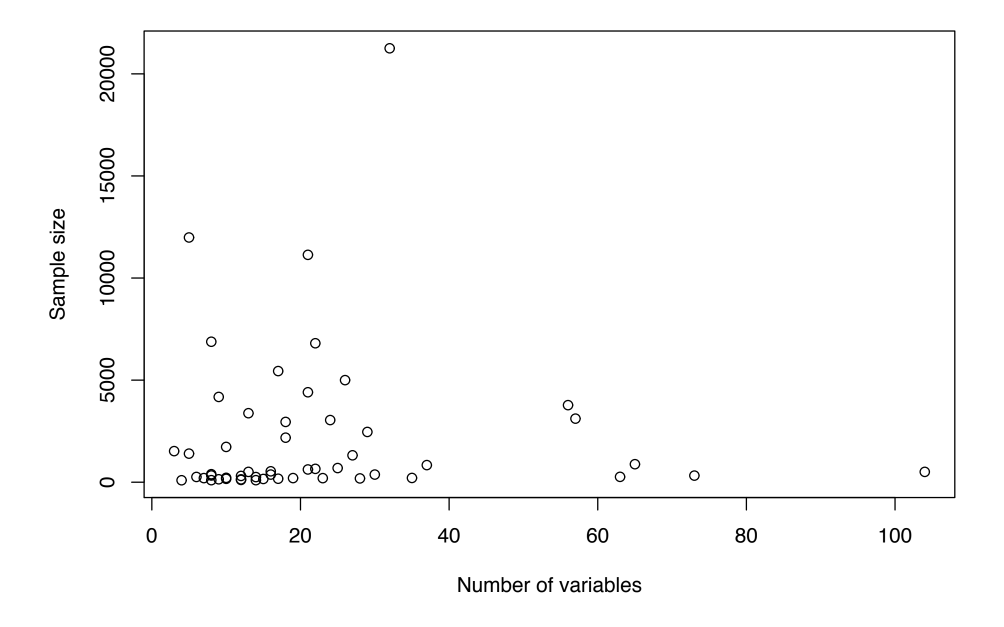
Bagged M5 constant (mcb)

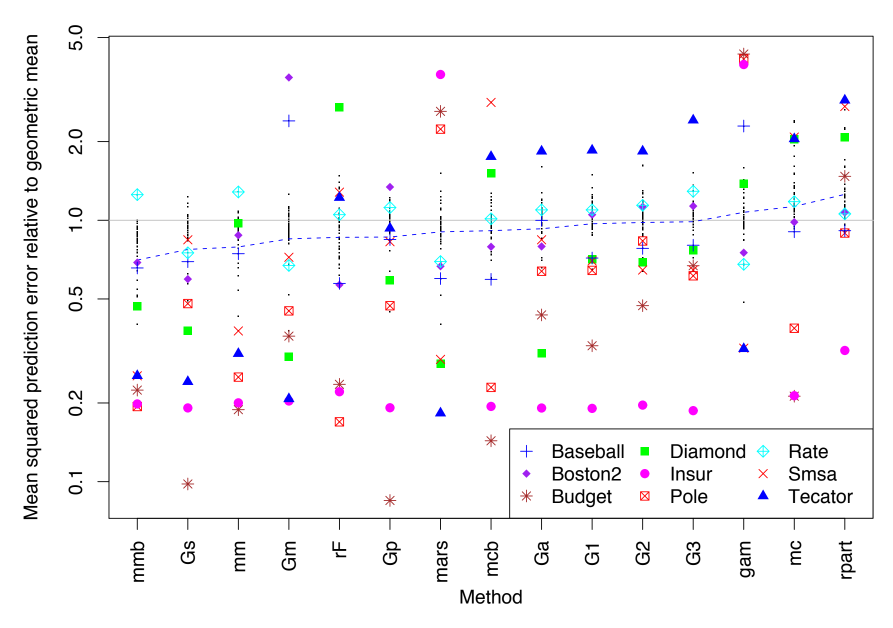
Bagged M5 multiple linear (mmb)

CART clone (rpart)

Random forest (rF)

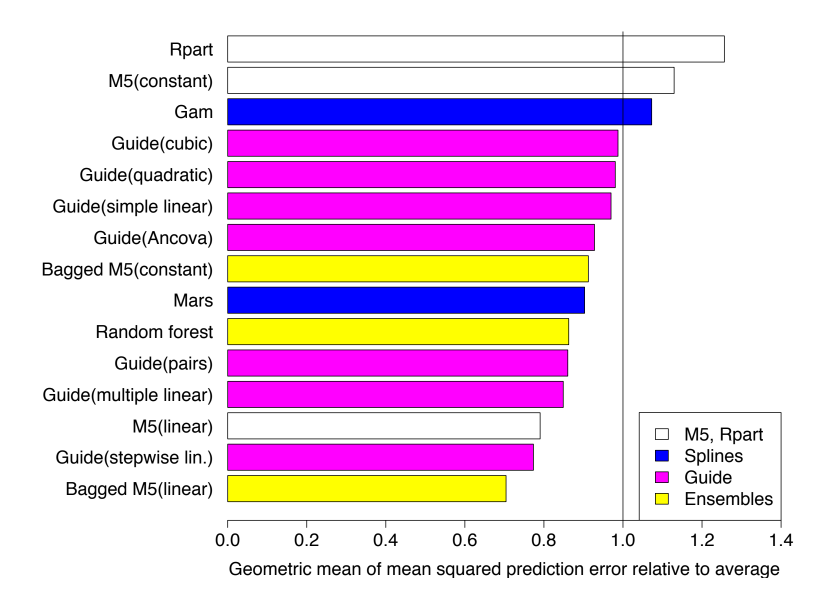
### Characteristics of 52 datasets (no missing values)



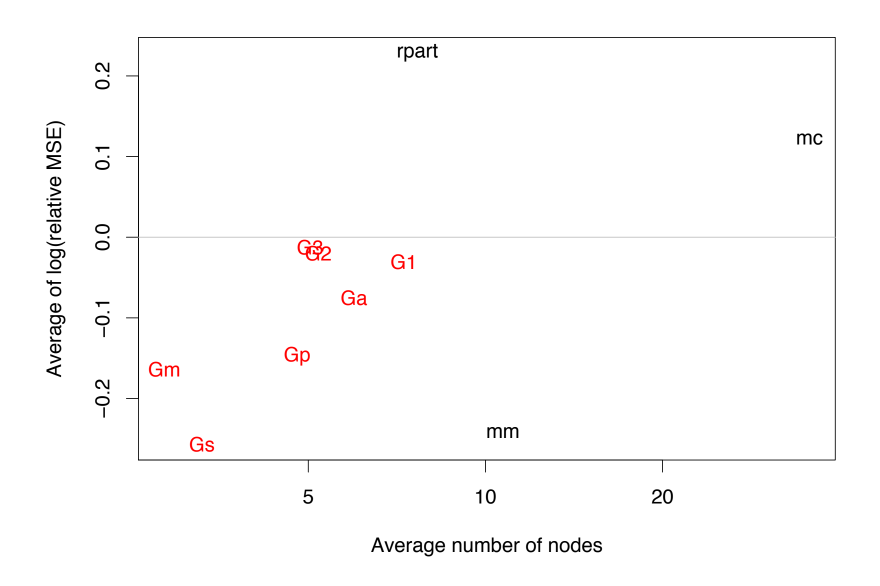


Means and medians joined by dashed and solid lines, respectively

#### **Relative MSE for 52 datasets**



#### **Prediction error vs tree size over 52 datasets**



### Some notations and definitions for asymptotics

- Let  $\mathcal{X}$  be M-dimensional Euclidean space.
- Given a fixed integer  $M_1$ , let  $\mathcal{B}$  be the collection of all polyhedra in  $\mathcal{X}$  having at most  $M_1$  faces. These sets can be described as the solutions to at most  $M_1$  inequalities, each inequality having the form  $b_1x_1 + \ldots + b_Mx_M \leq c$  (or < c).
- If  $M_1 \geq 2M$ ,  $\mathcal{B}$  includes all boxes in  $\mathcal{X}$  of the form

$$B = \{(x_1, \dots, x_M) : x_1 \in I_1, \dots, x_M \in I_M\}$$

where  $I_1, \ldots, I_M$  are open, closed, half-open, or half-closed intervals.

- Let  $X \in \mathcal{X}$  and  $\{(X_i, Y_i) : i = 1, ..., N\}$  be a random sample with the same distribution as (X, Y).
- Given  $N \ge 1$  and  $t \in \mathcal{X}$ , define  $\eta_N(t) = \{i : X_i \in t, 1 \le i \le N\}$ .
- Let  $\tilde{T}_N$  be a partition of  $\mathcal{X}$  into a finite number of disjoint sets, all of which are in  $\mathcal{B}$ , with  $\tilde{T}_N$  independent of (X,Y).

- Let  $\tau_N$  denote the partition function corresponding to  $\tilde{T}_N$ , so that  $\tau_N(x)$  is the set  $t \in \tilde{T}_N$  containing x.
- Let  $\delta(t) = \sup_{x,x' \in t} |x x'|$  be the diameter of t, where |x| is Euclidean distance.
- Let  $D_N(x) = \delta(\tau_N(x))$  be the diameter of the set  $t \in \tilde{T}_N$  containing x.
- Let  $d_N(x) = \bar{y}_N(\tau_N(x))$  be the estimate of the regression function  $d_B$ , where

$$\bar{y}_N(t) = \sum_{i \in \eta_N(t)} Y_i / |\eta_N(t)|.$$

- Let  $p_N(t) = N^{-1}|\{i: X_i \in t, 1 \le i \le N\}|$  be the empirical distribution of X.
- Let  $k_N$  be nonnegative constants such that

$$p_N(t) \ge k_N \log(N)/N$$
 for  $N \ge 1$  and  $t \in \tilde{T}_N$ .

# Bayes risk consistency of piecewise-constant regression models (Breiman et al. 1984)

**Theorem.** Suppose that  $E|Y|^q < \infty$  for some  $1 \le q < \infty$  and that

$$k_N \to \infty \text{ and } D_N(X) \stackrel{P}{\to} 0 \text{ as } N \to \infty.$$
 (1)

Let  $d_B(x) = E(Y | X = x)$ . Then  $E|d_N(X) - d_B(X)|^q \to 0$ .

Given any function d on  $\mathcal{X}$ , let  $R(d) = E[Y - d(X)]^2$  denote the mean squared error of d(X).

**Theorem.** Suppose that  $EY^2 < \infty$  and that condition (1) holds. Then  $\{d_N\}$  is *risk consistent*, i.e.,  $ER(d_N) \to R(d_B)$  as  $N \to \infty$ .

## Asymptotic uniform consistency (Kim et al., 2007)

Given X=x, let Y have mean f(x). Suppose f(x) is continuous in a compact rectangle C and there is a>0 such that

$$\sup_{x \in C} E\{\exp(a|Y - f(x)|) \mid X = x\} < \infty$$

Let  $T_n$  be the regression tree based on training sample size n,  $m_n$  = minimum node sample size, and  $\delta(t) = \sup_{x,z \in t} \|x - z\|$  be the diameter of node t

Assume that as  $n \to \infty$ ,

- 1.  $(\log n)/m_n \stackrel{P}{\rightarrow} 0$
- **2.**  $\sup_{t \in T_n} \delta(t) \stackrel{P}{\to} 0$
- 3. Minimum eigenvalue of node design matrices is bounded from 0 in probability

Let  $\hat{f}(x)$  be the regression estimate at x. Then

$$\sup_{x \in C} |\hat{f}(x) - f(x)| \stackrel{P}{\to} 0$$

#### **Conclusions**

- Parametric models are often constrained by range restrictions, missing values, distributional assumptions, and number and variety of variables.
- Tree models do not have such constraints.
- When the assumptions hold, parametric models are often more accurate.
   But when the assumptions are wrong, the results can be very misleading.
- Parametric models depend on statistical inference for model selection.
   Statistical inference is treacherous when there are many variables.
- Statistical inference is irrelevant to tree models, for which model selection is automatic.
- Tree models can supplement parametric models by validating the assumptions and suggesting alternative functional forms.
- Tree models allow high-level visualization of multivariate data through the tree structures and low-level visualization through plots of terminal nodes.
- Tree models are not necessarily unique. If variable selection is unbiased, each model gives a truthful description of the data.

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