# ML01 – Introduction to Machine Learning Tree-based methods

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#### Tree-based methods

- Here we describe tree-based methods for regression and classification.
- These involve recursively segmenting the predictor space into a number of simple regions.
- Since the set of splitting rules used to segment the predictor space can be summarized in a tree, these types of approaches are known as tree-based methods.



#### Tree-based methods

- Tree-based methods are simple and useful for interpretation.
- However they typically are not competitive with the best supervised learning approaches in terms of prediction accuracy.
- Hence we also discuss two methods for combining several trees:
  - Bagging and
  - Random forests.

These methods grow multiple trees which are then combined to yield a single consensus prediction.

 Combining a large number of trees can often result in dramatic improvements in prediction accuracy, at the expense of some loss interpretation.



#### Regression and classification Trees

- The tree-based approach can be applied to both regression and classification problems.
- We first consider regression trees, and then move on to classification/decision trees.



#### Overview

- Introductory example
- 2 Learning a regression tree
  - Tree building process
  - Cost-complexity pruning
- Classification trees
- 4 Combining trees
  - Bagging
  - Random Forests

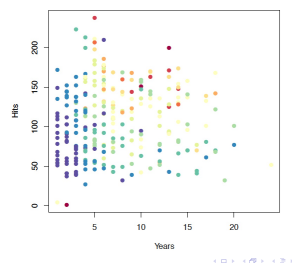


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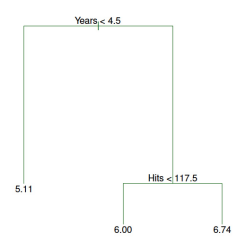
## Baseball salary data

Salary is color-coded from low (blue, green) to high (yellow,red)





#### Regression tree for these data

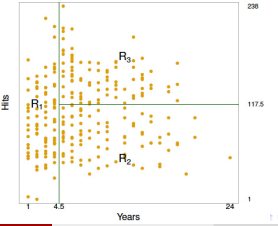


- At a given internal node, the label (of the form  $X_j < s$ ) indicates the left-hand branch emanating from the split, and the right-hand branch corresponds to  $X_j \ge s$ .
- The tree has two internal nodes and three terminal nodes, or leaves. The number in each leaf is the mean of the response for the observations that fall there.



#### Results

Overall, the tree stratifies or segments the players into three regions of predictor space:  $R_1 = \{X \mid Years < 4.5\}, R_2 = \{X \mid Years \ge 4.5, Hits < 117.5\}, and$  $R_3 = \{X \mid Years \ge 4.5, Hits \ge 117.5\}.$ 



## Terminology for trees

- The regions  $R_1$ ,  $R_2$ , and  $R_3$  are known as terminal nodes.
- The points along the tree where the predictor space is split are referred to as internal nodes.
- In the example, the two internal nodes are indicated by the text Years < 4.5 and Hits < 117.5.



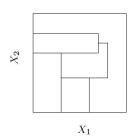
#### **Predictions**

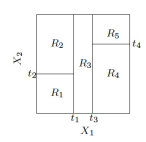
- We predict the response for a given test observation using the mean of the training observations in the region to which that test observation belongs.
- The prediction function is, thus, stepwise constant.
- A five-region example of this approach is shown in the next slide.

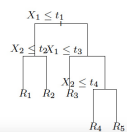


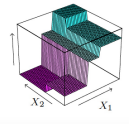


## Example











## Details of previous figure

Top Left: A partition of two-dimensional predictor space that could not result from recursive binary splitting.

Top Right: The output of recursive binary splitting on a two-dimensional example.

Bottom Left: A tree corresponding to the partition in the top right panel.

Bottom Right: A perspective plot of the prediction surface corresponding to that tree.





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#### Growing a regression tree

- We now turn to the question of how to grow a regression tree.
- Our data consists of p predictors and a response, for each of n observations: that is,  $(x_i, y_i)$  for i = 1, 2, ..., n, with  $x_i = (x_{i1}, x_{i2}, ..., x_{ip})$ .
- The algorithm needs to automatically decide on
  - 1 The splitting variables and split points
  - 2 What topology (shape) the tree should have.





#### Predicted response in a given region

• Suppose first that we have a partition into M regions  $R_1, R_2, \ldots, R_M$ and we model the response as a constant  $c_m$  in each region:

$$\widehat{f}(x) = \sum_{m=1}^{M} c_m I(x \in R_m)$$

 If we adopt as our criterion minimization of the RSS error  $\sum_{i=1}^{n} (y_i - \hat{f}(x_i))^2$ , it is easy to see that the best  $c_m$  is just the average of  $v_i$  in region  $R_m$ :

$$\widehat{c}_m = \text{Ave}\left\{y_i : x_i \in R_m\right\}.$$





## Top-down, greedy approach

Now, finding the best binary partition in terms of minimum sum of squares is generally computationally infeasible. Hence we proceed with a top-down, greedy approach:

- The approach is top-down because it begins at the top of the tree and then successively splits the predictor space; each split is indicated via two new branches further down on the tree.
- It is greedy because at each step of the tree-building process, the best split is made at that particular step, rather than looking ahead and picking a split that will lead to a better tree in some future step.



## Selecting a splitting variable and split point

• Starting with all of the data, consider a splitting variable  $X_j$  and split point s, and define the pair of half-spaces

$$R_1(j,s) = \{X \mid X_j \le s\} \text{ and } R_2(j,s) = \{X \mid X_j > s\}$$

• Then we seek the splitting variable  $X_j$  and split point s that solve

$$\min_{j,s} \left[ \sum_{x_i \in R_1(j,s)} (y_i - \widehat{c}_1(j,s))^2 + \sum_{x_i \in R_2(j,s)} (y_i - \widehat{c}_2(j,s))^2 \right]$$

with

$$\widehat{c}_1(j,s) = \text{Ave}\{y_i : x_i \in R_1(j,s)\} \text{ and } \widehat{c}_2(j,s) = \text{Ave}\{y_i : x_i \in R_2(j,s)\}$$

• For each splitting variable, the determination of the split point s can be done very quickly and hence by scanning through all of the predictors, determination of the best pair (j,s) is feasible.

#### Recursive building process

- Having found the best split, we partition the data into the two resulting regions and repeat the splitting process on each of the two regions.
- Then this process is repeated on all of the resulting regions.
- How large should we grow the tree? Clearly a very large tree might overfit the data, while a small tree might not capture the important structure. (More on this later).





## Regression trees in R

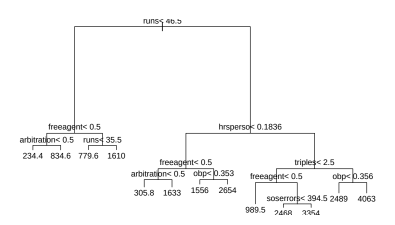
```
library(rpart)
baseball <- read.table("baseball.dat",header=TRUE)
n<-nrow(baseball)

train = sample(n, 2*n/3)
fit<-rpart(salary~.,data=baseball,subset=train,method="anova")
plot(fit)
text(fit,pretty=0,cex=0.8)</pre>
```





## Regression trees in R





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## Tuning the model's complexity

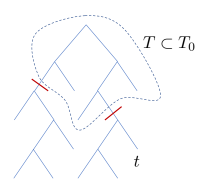
- Tree size is a tuning parameter governing the model's complexity, and the optimal tree size should be adaptively chosen from the data.
- One approach would be to split tree nodes only if the decrease in sum-of-squares due to the split exceeds some threshold. This strategy is too short-sighted, however, since a seemingly worthless split might lead to a very good split below it.
- The preferred strategy is to grow a large tree  $T_0$ , stopping the splitting process only when some minimum node size (say 5) is reached. Then this large tree is pruned using cost-complexity pruning.





#### **Notations**

- We define a subtree  $T \subset T_0$  to be any tree that can be obtained by pruning  $T_0$ , that is, collapsing any number of its internal nodes.
- We index terminal nodes by t, with node t representing region  $R_t$ .







#### Cost-complexity criterion

• Let T denote the set of terminal nodes in T. Let

$$n_t = \#\{x_i \in R_t\}, \quad \widehat{c}_t = \frac{1}{n_t} \sum_{x_i \in R_t} y_i,$$

$$Q_t = \frac{1}{n_t} \sum_{x_i \in R_t} (y_i - \widehat{c}_t)^2, \quad C(T) = \sum_{t \in \widetilde{T}} n_t Q_t$$

We define the cost-complexity criterion as

$$C_{\alpha}(T) = C(T) + \alpha |\widetilde{T}|$$





#### Cost-complexity pruning

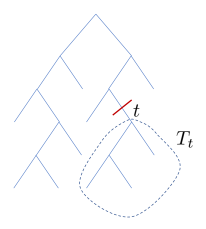
- The idea is to find, for each  $\alpha$ , the subtree  $T(\alpha) \subseteq T_0$  to minimize  $C_{\alpha}(T)$ .
- The tuning parameter  $\alpha \geq 0$  governs the tradeoff between tree size and its goodness of fit to the data. Larger values of  $\alpha$  result in smaller trees  $T(\alpha)$ .
- For  $\alpha = 0$ , the solution is the full tree  $T_0$ .
- Questions:
  - **1** For given  $\alpha$ , how to find a tree that minimizes  $C_{\alpha}(T)$ ?
  - **2** How to choose  $\alpha$ ?





#### Weakest link

- We start from the full tree  $T_0$ .
- For any internal node t, let  $T_t$  be the branch of T with root t.





## Weakest link (continued)

• If we prune  $T_t$ , the cost-complexity criterion becomes smaller if

$$C(t) + \alpha < C(T_t) + \alpha |\widetilde{T}_t| \Leftrightarrow \alpha > \frac{C(t) - C(T_t)}{|\widetilde{T}_t| - 1} = g_0(t)$$

- The weakest link  $t_0$  in  $T_0$  is the node such that  $g_0(t_0) = \min_t g_0(t)$ . Let  $\alpha_1 = g_0(t_0)$ .
- Meaning: if we increase  $\alpha$  starting from 0,  $t_0$  is the first node t such that pruning  $T_t$  improves the cost-complexity criterion.
- Let  $T_1 = T_0 T_{t_0}$ . We again find the weakest link  $t_1$  in  $T_1$ , etc.





## Sequence of optimal trees

• By iterating the above process until the tree is reduced to the root node  $t_{root}$ , we get a decreasing sequence of trees

$$T_0\supset T_1\supset\ldots\supset t_{root},$$

and an increasing sequence of  $\alpha$  values,  $0 = \alpha_0 < \alpha_1 < \alpha_2 < \dots$ 

• We can show that, for all  $k \geq 0$  and all  $\alpha \in [\alpha_k, \alpha_{k+1})$ , the optimum tree  $T(\alpha)$  is equal to  $T_k$ .





#### Choosing $\alpha$

- If we have a lot of data, it is easy to estimate the sum-of-squares error of each subtree in the sequence  $T_0 \supset T_1 \supset \ldots \supset t_{root}$  using a validation set. We choose the tree  $T_k$  with minimum validation error.
- Otherwise, we use cross-validation.





#### Cross-validation

- Using the whole training set, we get a sequence of trees,  $T_0 \supset T_1 \supset \ldots \supset t_{root}$ , where  $T_k$  is the best tree for  $\alpha_k \leq \alpha < \alpha_{k+1}$ .
- For  $k = 0, 1, 2, \ldots$ , set  $\beta_k = \sqrt{\alpha_k \alpha_{k+1}}$
- Assume we use *K*-fold cross validation: we partition the training data in *K* subsets of approximately equal size.
- We construct K sequences of trees by leaving each of K subsets out and building the trees using the K-1 remaining subsets. Let  $T_0^{(r)} \supset T_1^{(r)} \supset \ldots \supset t_{root}^{(r)}$  be the sequence of trees obtained by leaving subset r out.
- Compute the cross-validated error  $C_{cv}(T_k)$  using the trees  $T^{(r)}(\beta_k)$ , r = 1, ..., K.
- Select the tree  $T_k$  corresponding to the minimum cross-validated error.



```
fit<-rpart(salary~.,data=baseball,subset=train,method="anova",
control = rpart.control(xval = 10, minbucket = 2, cp = 0))
printcp(fit)
plotcp(fit)</pre>
```





```
> printcp(fit)
Regression tree:
rpart(formula = salary ~ .. data = baseball, subset = train.
   method = "anova", control = rpart.control(xval = 10, minbucket = 2,
       cp = 0)
Variables actually used in tree construction:
 [1] arbitration average
                               doubles
                                                        freeagent
                                            errors
                                                                     hits
 [7] hitspererror hitsperso
                               homeruns
                                            hrsperso
                                                                      rbis
                                                         obp
[13] rbisperso
                  runs
                               runsperso
                                            sbsobp
                                                         sbsruns
                                                                      sos
[19] soserrors
                 triples
                              walks
                                            walksperso
Root node error: 364623518/224 = 1627784
n= 224
          CP nsplit rel error xerror
  3.6931e-01
                   0 1.000000 1.01101 0.112712
2 1.2284e-01
                     0.630689 0.70087 0.080434
  6.0838e-02
                     0.507850 0.68227 0.081909
  4.2082e-02
                  4 0.386175 0.62816 0.078994
  3.6111e-02
                  5 0.344093 0.62059 0.086438
  3.3694e-02
                    0.307982 0.59791 0.085306
  2.5680e-02
                  7 0.274289 0.57487 0.084283
  2.1447e-02
                  8 0.248609 0.58795 0.084931
9 1.6639e-02
                  9 0.227162 0.57559 0.085300
10 1.5463e-02
                 10 0.210522 0.56992 0.084713
11 1.5064e-02
                 11 0.195059 0.56094 0.086454
12 1.4133e-02
                 12 0.179996 0.61223 0.101958
13 1.3472e-02
                     0.165862 0.60748 0.099526
```

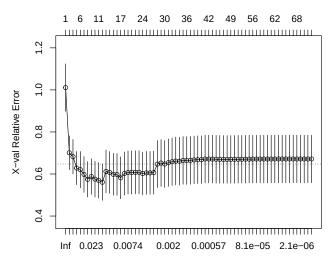


0.152390 0.59866 0.099545

14 1.2588e-02

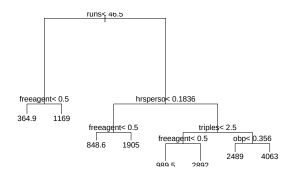
15 1.2339e-02

#### size of tree





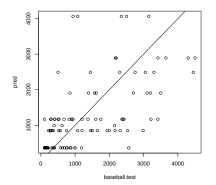
```
pruned_tree<-prune(fit,cp=2.5680e-02)
plot(pruned_tree)
text(pruned_tree,pretty=0)</pre>
```





## Prediction with a regression tree in R

```
yhat=predict(pruned_tree,newdata=baseball[-train,])
baseball.test=baseball[-train, "salary"]
plot(baseball.test,yhat)
abline(0,1)
```





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#### Classification trees

- If the response is a categorical variable (factor) taking c values indexed by  $1, 2, \ldots, c$ , we have a classification problem.
- The only changes needed in the tree-growing algorithm concern the criteria for splitting nodes and pruning the tree.
- In regression, each split aims at obtaining nodes with small sample variance

$$Q_t = \frac{1}{n_t} \sum_{x_i \in R_t} (y_i - \widehat{c}_t)^2.$$

• In classification, each split will aim at obtaining nodes as "pure" as possible (a node is pure if it contains observations from only one class). We need to define a suitable impurity measure.



#### Notations

• In a node t, let

$$\widehat{p}_{tk} = \frac{1}{n_t} \sum_{x_i \in R_t} I(y_i = k)$$

be the proportion of class k observations in node t.

- We have  $\sum_{k=1}^{c} \widehat{p}_{tk} = 1$ .
- We classify the observations in node t to the majority class in node t:

$$k(t) = \arg \max_{k} \widehat{p}_{tk},$$





### Impurity measures

• Different measures  $Q_t$  of node impurity include the following: Misclassification error:

$$Q_t^{\mathsf{mis}} = rac{1}{n_t} \sum_{x_i \in R_t} I(y_i 
eq k(t)) = 1 - \widehat{
ho}_{tk(t)}$$

Gini index:

$$Q_t^{\mathsf{Gini}} = \sum_{k=1}^c \widehat{p}_{tk} (1 - \widehat{p}_{tk})$$

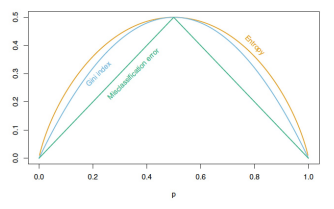
Entropy:

$$Q_t^{\mathsf{ent}} = -\sum_{k=1}^c \widehat{p}_{tk} \log \widehat{p}_{tk}$$

• All three criteria are equal to 0 when  $\widehat{p}_{tk} = 1$  for some k, and are maximum when  $\widehat{p}_{tk} = 1/c$  for all k.

### Comparison between impurity measures

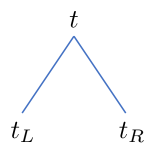
• Plot as a function of the proportion p of one class in the case c=2:



• All three are similar, but entropy and the Gini index are differentiable, and hence more amenable to numerical optimization.

## Selecting the best split

- Consider a node t with size  $n_t$  with impurity  $Q_t$ .
- For some variable j and split point s, we split t in two nodes, t<sub>1</sub> and  $t_R$ , with sizes  $n_{t_l}$  and  $n_{t_R}$ , and with impurities  $Q_{t_l}$  and  $Q_{t_R}$ .







## Selecting the best split

• The average decrease of impurity is

$$\Delta(j,s) = Q_t - \left(\frac{n_{t_L}}{n_t}Q_{t_L} + \frac{n_{t_R}}{n_t}Q_{t_R}\right)$$

- ullet If  $Q_t$  is the entropy, then  $\Delta(j,s)$  is interpreted as an information gain.
- We select at each step the splitting variable j and the split point s that maximizes  $\Delta(j,s)$  or, equivalently, that minimizes the average impurity

$$\frac{n_{t_L}}{n_t}Q_{t_L} + \frac{n_{t_R}}{n_t}Q_{t_R}$$





## Example: Heart data

- A retrospective sample of males in a coronary heart disease (CHD) high-risk region of the Western Cape, South Africa.
- There are roughly two controls per positive case of CHD.
- Variables:
  - sbp: systolic blood pressure
  - tobacco: cumulative tobacco (kg)
  - Idl: low density lipoprotein cholesterol
  - adiposity
  - famhist: family history of heart disease (Present, Absent)
  - typea: type-A behavior
  - obesity
  - alcohol: current alcohol consumption
  - age: age at onset
  - chd: response, coronary heart disease



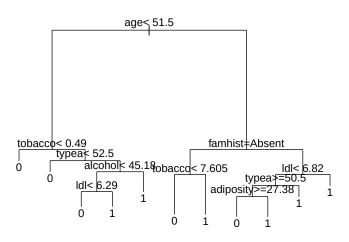


# Tree growing in R

```
heart<-read.table(file = "SAheart.data", sep=", ", header=T,
           row.names=1)
n<-nrow(heart)
train = sample(n, 2*n/3)
fit <- rpart(chd ~ ., data = heart, method="class",</pre>
           subset=train, parms = list(split = 'gini'))
plot(fit,margin = 0.05)
text(fit,pretty=0,cex=0.8)
plotcp(fit)
```

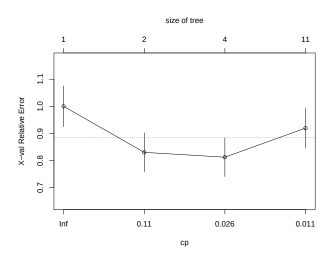


#### Tree





### Cross-validation error







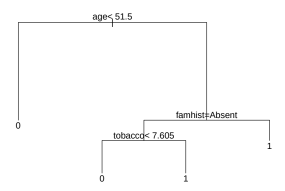
## Pruning

```
pruned_tree<-prune(fit,cp=0.026)
plot(pruned_tree,margin = 0.05)
text(pruned_tree,pretty=0)</pre>
```





### Pruned tree





#### Test error rate estimation

```
yhat=predict(pruned_tree,newdata=heart[-train,],type='class')
y.test=heart[-train,"chd"]
table(y.test,yhat)
err<-1-mean(y.test==yhat)</pre>
```

Confusion matrix:

```
prediction
true class
        95 11
        31
            17
```

Test error rate: 0.27





## Advantages and disadvantages of trees

- Trees can be displayed graphically, and are easily interpreted even by a non-expert (especially if they are small).
- Trees can easily handle qualitative predictors without the need to create dummy variables.
- Unfortunately, trees generally do not have the same level of predictive accuracy as some of the other modern regression and classification approaches.
- However, by aggregating many decision trees, the predictive performance of trees can be substantially improved.
- We will see two combination methods, both based on the same resampling technique: the bootstrap.

ML01 - Tree-based methods



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### Bagging

- Bootstrap aggregation, or bagging, is a general-purpose procedure for reducing the variance of a statistical learning method; we introduce it here because it is particularly useful and frequently used in the context of decision trees.
- Recall that given a set of n independent observations  $X_1, \ldots, X_n$ , each with variance  $\sigma^2$ , the variance of the mean  $\overline{X}$  of the observations is given by  $\sigma^2/n$ .
- In other words, averaging a set of observations reduces variance. Of course, this is not practical because we generally do not have access to multiple training sets.





## Bagging – continued

- Instead, we can the bootstrap technique: to generate a bootstrap training set, we sample n observations with replacement from the original training set.
- For instance, if n = 5, the sample is (0.5, 1.2, -0.3, 0.8, -1.4), we have the following B=3 bootstrap samples:

$$(-0.30, 0.50, 0.50, 0.80, -0.30)$$
  
 $(0.8, -0.3, 0.5, 0.8, -0.3)$   
 $(1.20, 1.20, 0.80, 0.80, 1.20)$ 

• In the bagging approach we generate B different bootstrap training sets and we train B decision trees, one from each bootstrap dataset.



## Bagging regression and classification trees

For regression problems, we average all the predictions to obtain

$$\widehat{f}_{bag}(x) = \frac{1}{B} \sum_{b=1}^{B} \widehat{f}^{*b}(x)$$

where  $\hat{f}^{*b}(x)$  is the prediction at x for the b-th tree.

- For classification trees:
  - For each test observation, we record the class predicted by each of the B trees, and take a majority vote: the overall prediction is the majority class among the B predictions.
  - If we are interested in the posterior probabilities, we can rather average the class proportions in the terminal nodes.

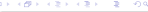




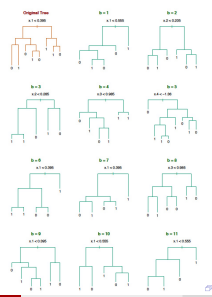
### Example

- We generated a sample of size n = 30, with two classes and p = 5 predictors, each having a standard Gaussian distribution with pairwise correlation 0.95.
- The response Y was generated according to  $Pr(Y=1 \mid x_1 \leq 0.5) = 0.2$ ,  $Pr(Y=1 \mid x_1 > 0.5) = 0.8$ . The Bayes error is 0.2.
- A test sample of size 2000 was also generated from the same population.
- We fit classification trees to the training sample and to each of 200 bootstrap samples. No pruning was used.



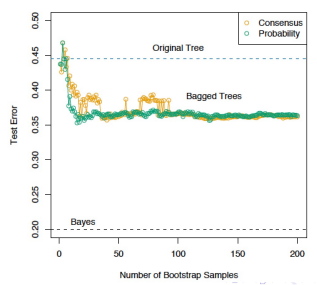


# Bagged decision trees





### Error curves





## Out-of-Bag Error Estimation

- It turns out that there is a very straightforward way to estimate the test error of a bagged model.
- Recall that the key to bagging is that trees are repeatedly fit to bootstrapped subsets of the observations. One can show that on average, each bagged tree makes use of around two-thirds of the observations
- The remaining one-third of the observations not used to fit a given bagged tree are referred to as the out-of-bag (OOB) observations.
- We can predict the response for the *i*-th observation using each of the trees in which that observation was OOB. This will yield around B/3predictions for the *i*-th observation, which we average.





#### Overview

- Introductory example
- 2 Learning a regression tree
  - Tree building process
  - Cost-complexity pruning
- Classification trees
- Combining trees
  - Bagging
  - Random Forests





#### Random Forests

- Random forests provide an improvement over bagged trees by way of a small tweak that decorrelates the trees. This reduces the variance when we average the trees.
- As in bagging, we build a number of decision trees on bootstrapped training samples.
- But when building these decision trees, each time a split in a tree is considered, a random selection of m predictors is chosen as split candidates from the full set of p predictors. The split is allowed to use only one of those m predictors.
- A fresh selection of m predictors is taken at each split, and typically we choose  $m \approx \sqrt{p}$ .
- When m = p, random forests boil down to bagging.





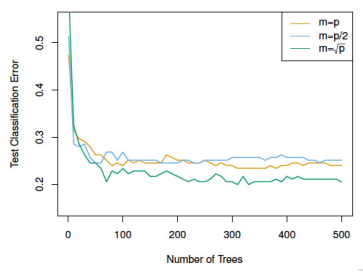
### Example: gene expression data

- We applied random forests to a high-dimensional biological data set consisting of expression measurements of 500 genes measured on tissue samples from 349 patients.
- Each of the patient samples belongs to one of c = 15 classes: either normal or one of 14 different types of cancer.
- We randomly divided the observations into a training and a test set, and applied random forests to the training set for three different values of the number of splitting variables m.





# Results: gene expression data



### Variable importance

- Bagging improves prediction accuracy at the expense of interpretability.
- Although the collection of bagged trees is much more difficult to interpret than a single tree, one can obtain an overall summary of the importance of each predictor.
- There are several ways to measure the importance of a variable.





### Variable importance measures

Mean decrease in accuracy: For each tree, the prediction error on the OOB portion of the data is recorded (error rate for classification, MSE for regression). Then the same is done after randomly permuting each predictor variable. The difference between the two are then averaged over all trees.

Mean decrease in node impurity: total decrease in node impurities from splitting on the variable, averaged over all trees. For classification, the node impurity is measured by the Gini index. For regression, it is measured by RSS.



## Bagging in R

Confusion matrix:

	prediction	
true class	1	2
1	81	13
2	38	22

• Test error rate: 0.33



#### Random forests in R

Confusion matrix:

	prediction	
true class	1	2
1	84	10
2	36	24

Test error rate: 0.30



# Variable importance plot



