

Data Mining Classification Trees (2)

Ad Feelders

Universiteit Utrecht

September 13, 2018

Basic Tree Construction Algorithm

Construct tree

```
nodelist ← {{training sample}}
```

Repeat

```
    current node ← select node from nodelist
```

```
    nodelist ← nodelist – current node
```

```
    if impurity(current node) > 0
```

```
    then
```

```
        S ← candidate splits in current node
```

```
         $s^* \leftarrow \arg \max_{s \in S} \text{impurity reduction}(s, \text{current node})$ 
```

```
        child nodes ← apply( $s^*$ , current node)
```

```
        nodelist ← nodelist  $\cup$  child nodes
```

```
    fi
```

Until $nodelist = \emptyset$

Overfitting and Pruning

- We continue splitting until all leaf nodes of T contain examples of a single class (i.e. resubstitution error $R(T) = 0$).
- Is this a good tree for predicting the class of new examples?
- Not unless the problem is truly “deterministic”!
- Problem of *overfitting*.

Proposed Solutions

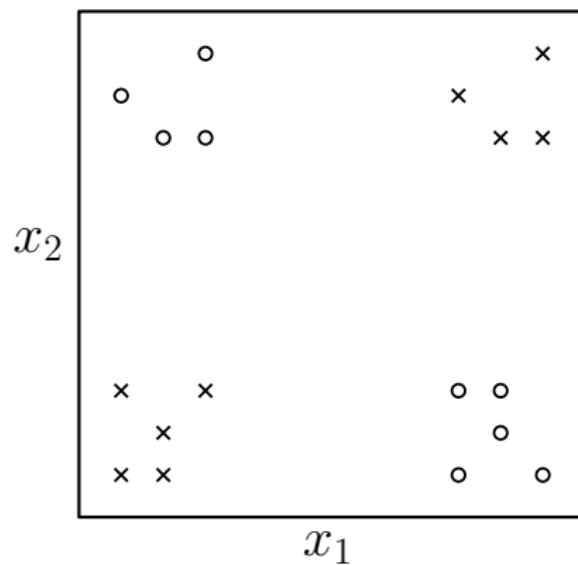
- Stopping Rules: e.g. don't expand a node if the impurity reduction of the best split is below some threshold.
- Pruning: grow a very large tree T_{\max} and merge back nodes.

Note: in the practical assignment we do use a stopping rule based on the `nmin` and `minleaf` parameters.

Stopping Rules

Disadvantage: sometimes you first have to make a weak split to be able to follow up with a good split.

Since we only look one step ahead we may miss the good follow-up split.



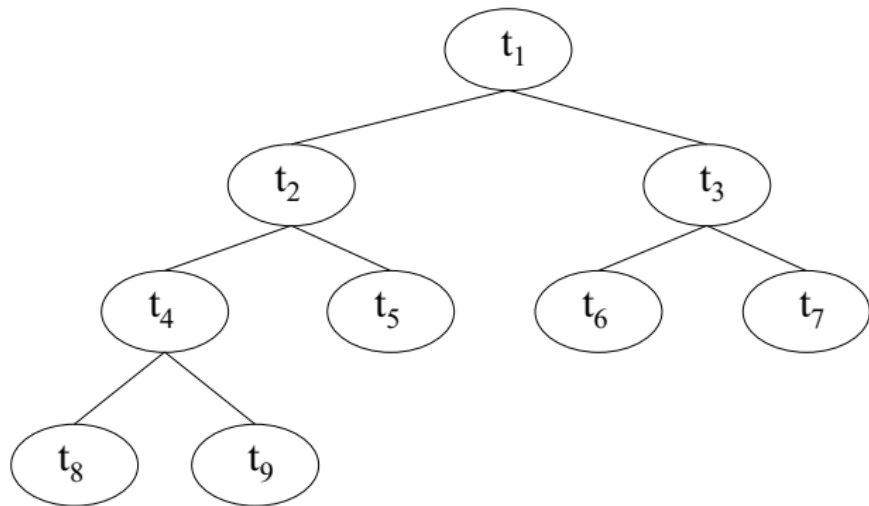
Pruning

- To avoid the problem of stopping rules, we first grow a very large tree on the training sample, and then *prune* this large tree.
- Objective: select the pruned subtree that has lowest *true* error rate.
- Problem: how to find this pruned subtree?

Pruning Methods

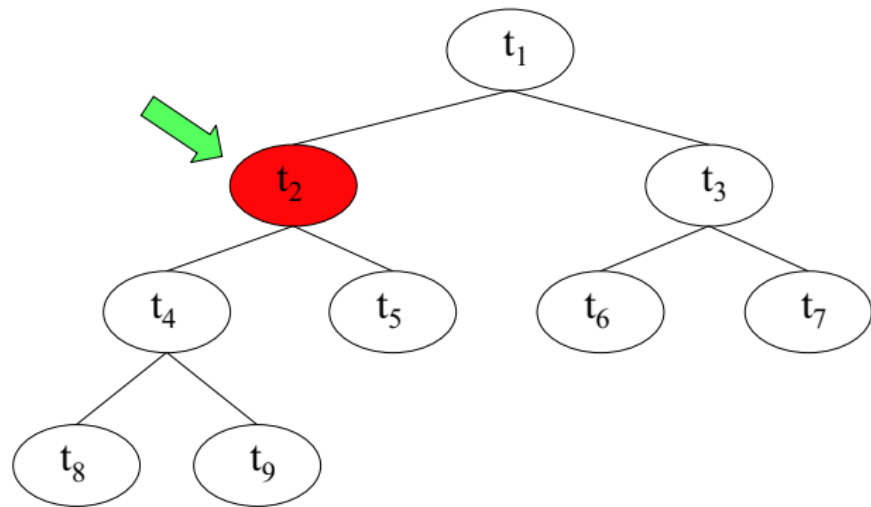
- Cost-complexity pruning (Breiman et al.; CART),
also called *weakest link* pruning.
- Reduced-error pruning (Quinlan)
- Pessimistic pruning (Quinlan; C4.5)
- ...

Terminology: Tree T

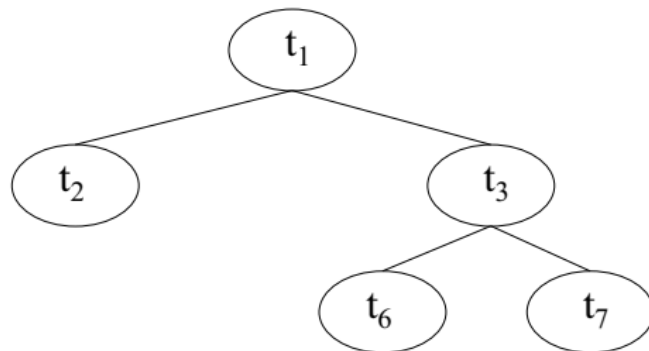


\tilde{T} denotes the collection of leaf nodes of tree T .
 $\tilde{T} = \{t_5, t_6, t_7, t_8, t_9\}, |\tilde{T}| = 5$

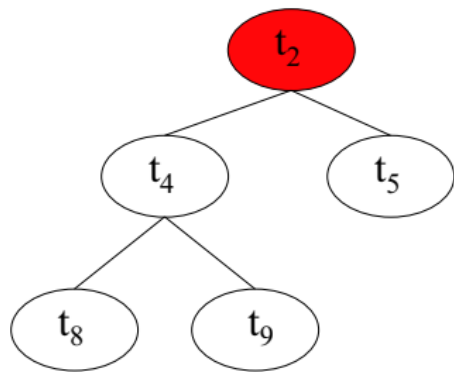
Terminology: Pruning T in node t_2



Terminology: T after pruning in t_2 : $T - T_{t_2}$



Terminology: Branch T_{t_2}



$$\tilde{T}_{t_2} = \{t_5, t_8, t_9\}, |\tilde{T}_{t_2}| = 3$$

Cost-complexity pruning

- The total number of pruned subtrees of a balanced binary tree with ℓ leaves is

$$\lfloor 1.5028369^\ell \rfloor$$

- With just 40 leaf nodes we have approximately 12 million pruned subtrees.
- Exhaustive search not recommended.
- Basic idea of cost-complexity pruning: reduce the number of pruned subtrees we have to consider by selecting the ones that are the “best of their kind” (in a sense to be defined shortly...)

Total cost of a tree

Strike a balance between fit and complexity. Total cost $C_\alpha(T)$ of tree T

$$C_\alpha(T) = R(T) + \alpha|\tilde{T}|$$

Total cost consists of two components:

- resubstitution error $R(T)$, and
- a penalty for the complexity of the tree $\alpha|\tilde{T}|$, ($\alpha \geq 0$).

Note: $R(T) = \frac{\text{number of wrong classifications made by } T}{\text{number of examples in the training sample}}$

Tree with lowest total cost

- Depending on the value of α , different pruned subtrees will have the lowest total cost.
- For $\alpha = 0$ (no complexity penalty) the tree with smallest resubstitution error wins.
- For higher values of α , a less complex tree that makes a few more errors might win.

As it turns out, we can find a nested sequence of pruned subtrees of T_{\max} , such that the trees in the sequence minimize total cost for consecutive intervals of α values.

Smallest minimizing subtree

For any value of α , there exists a smallest minimizing subtree $T(\alpha)$ of T_{\max} that satisfies the following conditions:

- ① $C_\alpha(T(\alpha)) = \min_{T \leq T_{\max}} C_\alpha(T)$
(that is, $T(\alpha)$ minimizes total cost for that value of α).
- ② If $C_\alpha(T) = C_\alpha(T(\alpha))$ then $T(\alpha) \leq T$.
(that is, $T(\alpha)$ is a pruned subtree of all trees that minimize total cost).

Note: $T' \leq T$ means T' is a pruned subtree of T ,
i.e. it can be obtained by pruning T in 0 or more nodes.

Sequence of subtrees

Construct a *decreasing sequence* of pruned subtrees of T_{\max}

$$T_{\max} > T_1 > T_2 > T_3 > \dots > \{t_1\}$$

(where t_1 is the root node of the tree) such that T_k is the smallest minimizing subtree for $\alpha \in [\alpha_k, \alpha_{k+1})$.

Note: From a computational viewpoint, the important property is that T_{k+1} is a pruned subtree of T_k , i.e. it can be obtained by pruning T_k . No backtracking is required.

Decomposition of total cost

Total cost has an additive decomposition over the leaf nodes of a tree:

$$C_\alpha(T) = \sum_{t \in \tilde{T}} R(t) + \alpha$$

$R(t)$ is the number of errors we make in node t if we predict the majority class, divided by the total number of observations in the training sample.

Finding the T_k and corresponding α_k

T_t : branch of T with root node t .

After pruning in t , its contribution to total cost is:

$$C_\alpha(\{t\}) = R(t) + \alpha,$$

The contribution of T_t to the total cost is:

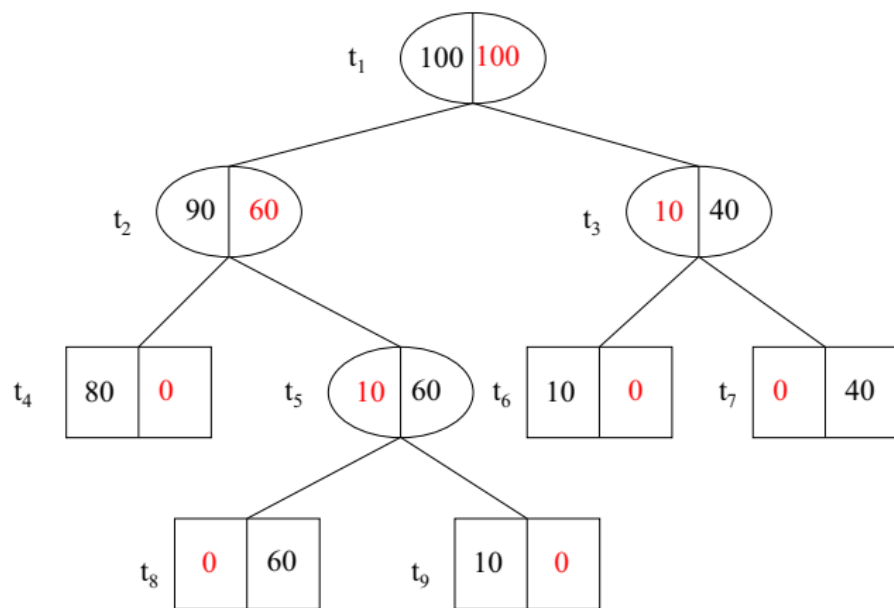
$$C_\alpha(T_t) = R(T_t) + \alpha |\tilde{T}_t|,$$

where $R(T_t) = \sum_{t' \in \tilde{T}_t} R(t')$.

$T - T_t$ becomes better than T when

$$C_\alpha(\{t\}) = C_\alpha(T_t)$$

Computing contributions to total cost of T



$$C_\alpha(\{t_2\}) = R(t_2) + \alpha = \frac{3}{10} + \alpha$$

$$C_\alpha(T_{t_2}) = R(T_{t_2}) + \alpha |\tilde{T}_{t_2}| = \sum_{t' \in \tilde{T}_{t_2}} R(t') + \alpha |\tilde{T}_{t_2}| = 0 + 3\alpha$$

Solving for α

The total cost of T and $T - T_t$ become equal when

$$C_\alpha(\{t\}) = C_\alpha(T_t),$$

At what value of α does this happen?

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

Solving for α we get

$$\alpha = \frac{R(t) - R(T_t)}{|\tilde{T}_t| - 1}$$

Computing $g(t)$: the “critical” α value for node t

- For each non-terminal node t we compute its “critical” α value:

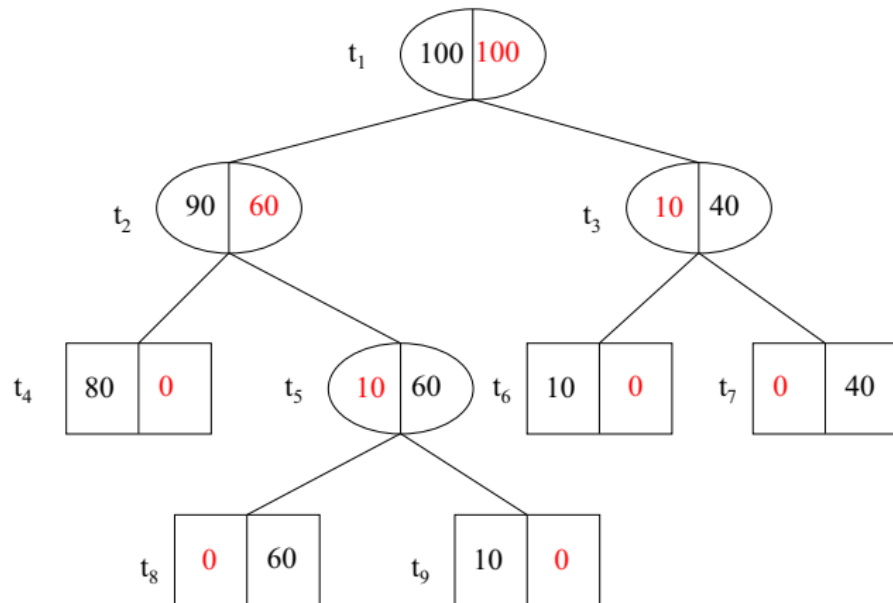
$$g(t) = \frac{R(t) - R(\tilde{T}_t)}{|\tilde{T}_t| - 1}$$

In words:

$$g(t) = \frac{\text{increase in error due to pruning in } t}{\text{decrease in } \# \text{ leaf nodes due to pruning in } t}$$

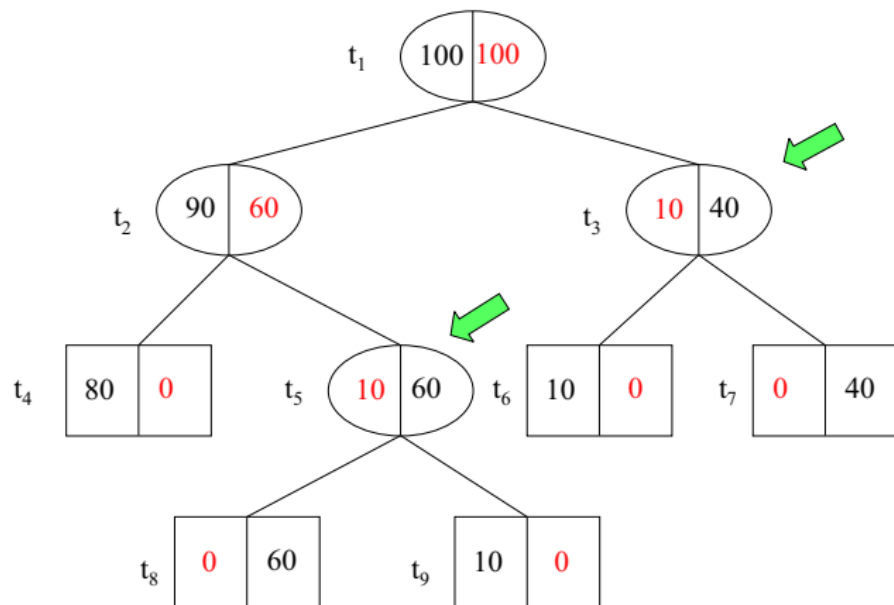
- Subsequently, we prune in the nodes for which $g(t)$ is the smallest (the “weakest links”).
- This process is repeated until we reach the root node.

Computing $g(t)$: the “critical” α value for node t



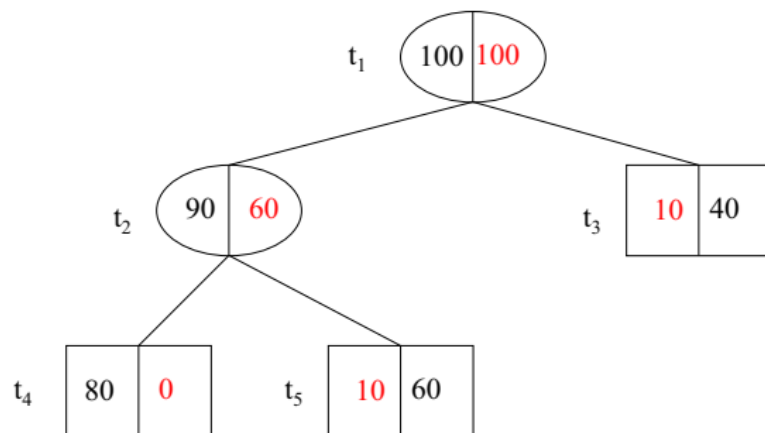
$$g(t_1) = \frac{1}{8}, g(t_2) = \frac{3}{20}, g(t_3) = \frac{1}{20}, g(t_5) = \frac{1}{20}.$$

Finding the weakest links



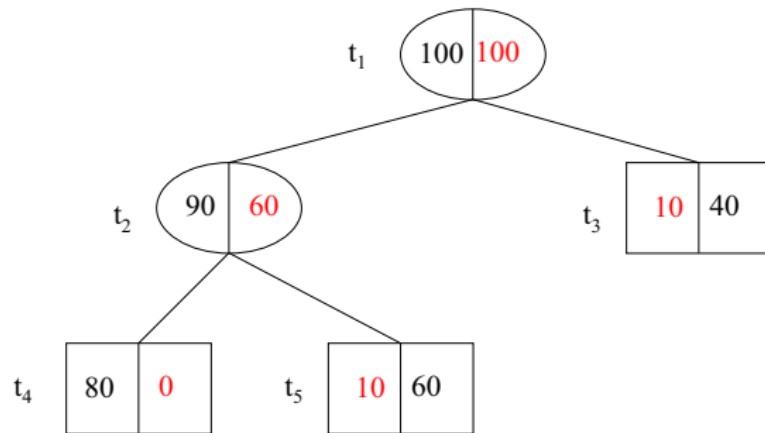
$$g(t_1) = \frac{1}{8}, g(t_2) = \frac{3}{20}, g(t_3) = \frac{1}{20}, g(t_5) = \frac{1}{20}.$$

Pruning in the weakest links



By pruning the weakest links we obtain the next tree in the sequence.

Repeating the same procedure



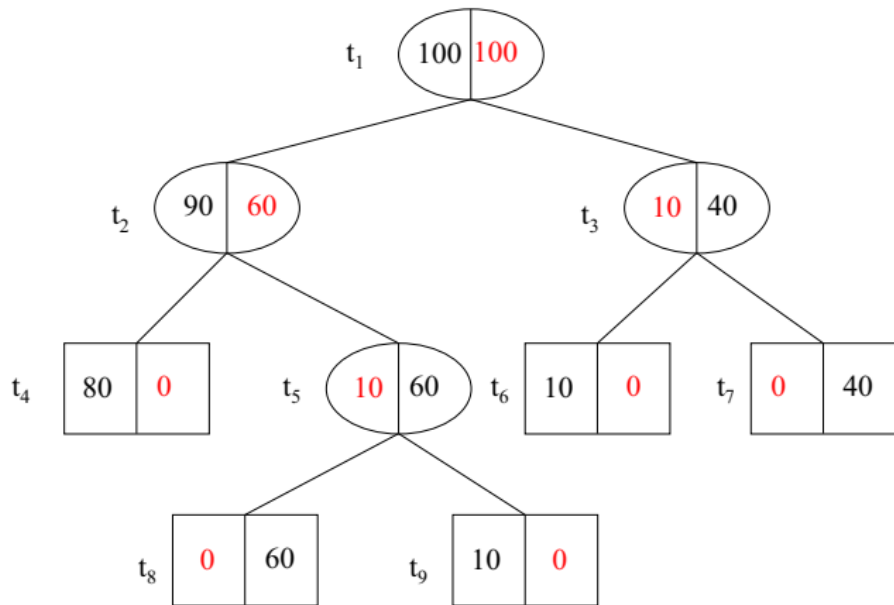
$$g(t_1) = \frac{2}{10}, g(t_2) = \frac{1}{4}.$$

Going back to the root

t_1	100	100
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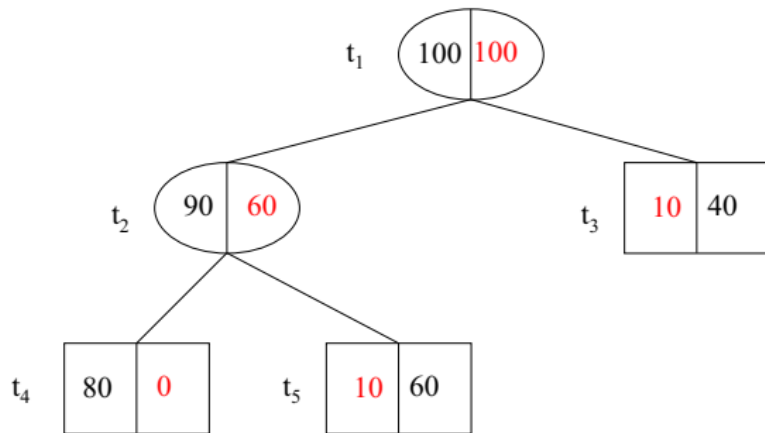
We have arrived at the root so we're done.

The best tree for $\alpha \in [0, \frac{1}{20})$



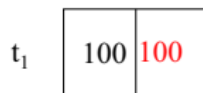
The big tree is the best for values of α below $\frac{1}{20}$.

The best tree for $\alpha \in [\frac{1}{20}, \frac{2}{10})$



When α reaches $\frac{1}{20}$ this tree becomes the best.

The best tree for $\alpha \in [\frac{2}{10}, \infty)$



When α reaches $\frac{2}{10}$ the root wins and we're done.

Compute Pruning Sequence

$T_1 \leftarrow T(\alpha = 0); \alpha_1 \leftarrow 0; k \leftarrow 1$

While $T_k > \{t_1\}$ do

For all non-terminal nodes $t \in T_k$

$$g_k(t) \leftarrow \frac{R(t) - R(T_{k,t})}{|\tilde{T}_{k,t}| - 1}$$

$$\alpha_{k+1} \leftarrow \min_t g_k(t)$$

Visit the nodes in top-down order and prune

whenever $g_k(t) = \alpha_{k+1}$ to obtain T_{k+1}

$$k \leftarrow k + 1$$

od

Note: $T_{k,t}$ is the branch of T_k with root node t ,
and T_k is the pruned tree in iteration k .

Algorithm to compute T_1 from T_{\max}

If we don't continue splitting until all nodes are pure,
then $T_1 = T(\alpha = 0)$ may not be the same as T_{\max} .

Compute T_1 from T_{\max}

$T' \leftarrow T_{\max}$

Repeat

Pick any pair of terminal nodes ℓ and r

with common parent t in T'

such that $R(t) = R(\ell) + R(r)$, and set

$T' \leftarrow T' - T_t$ (i.e. prune T' in t)

Until no more such pair exists

$T_1 \leftarrow T'$

Selection of the final tree: using a test set

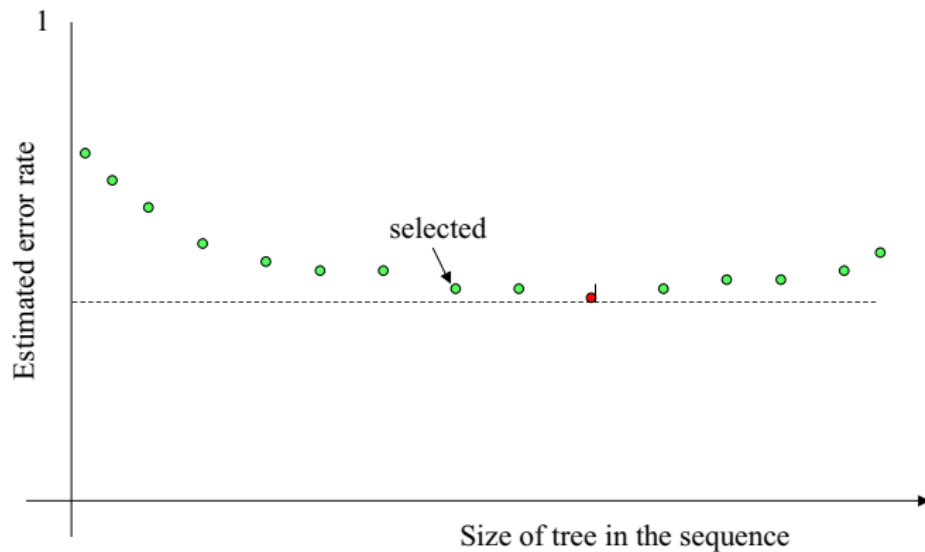
Pick the tree T from the sequence with the lowest error rate $R^{ts}(T)$ on the test set.

This is an *estimate* of the true error rate $R^*(T)$ of T .

The standard error of this estimate is

$$SE(R^{ts}) = \sqrt{\frac{R^{ts}(1 - R^{ts})}{n_{test}}}$$

Selection of the final tree: the 1-SE rule



1-SE rule: select the smallest tree with R^{ts} within one standard error of the minimum.

Selection of the final tree: using cross-validation

- When the data set is relatively small, it is a bit of a waste to set aside part of the data for testing.
- A way to avoid this problem is to use *cross-validation*.

v-fold cross-validation (general)

Let C be a complexity parameter of a learning algorithm (like α in the classification tree algorithm). To select the best value of C from a range of values c_1, \dots, c_m we proceed as follows.

- ① Divide the data into v groups G_1, \dots, G_v .
- ② For each value c_i of C
 - ① For $j = 1, \dots, v$
 - ① Train with $C = c_i$ on all data *except* group G_j .
 - ② Predict on group G_j .
 - ② Compute the CV prediction error for $C = c_i$.
- ③ Select the value c^* of C with the smallest CV prediction error.
- ④ Train on the complete training sample with $C = c^*$

Using cross-validation: Step 1

Construct a tree on the full data set, and compute $\alpha_1, \alpha_2, \dots, \alpha_K$ and $T_1 > T_2 > \dots > T_K$.

Estimate the error of a tree T_k from this sequence as follows.

Set

$$\beta_1 = 0,$$

$$\beta_2 = \sqrt{\alpha_2 \alpha_3},$$

$$\beta_3 = \sqrt{\alpha_3 \alpha_4},$$

$$\dots,$$

$$\beta_{K-1} = \sqrt{\alpha_{K-1} \alpha_K},$$

$$\beta_K = \infty.$$

β_k is typical value for $[\alpha_k, \alpha_{k+1})$.

Using cross-validation: Step 2

Divide the data set into v groups G_1, G_2, \dots, G_v (of approximately equal size) and for each group G_j

- ① Build a tree on all data except G_j , and determine the smallest minimizing subtrees $T^{(j)}(\beta_1), T^{(j)}(\beta_2), \dots, T^{(j)}(\beta_K)$ for this reduced data set.
- ② Compute the error of $T^{(j)}(\beta_k)$ ($k = 1, \dots, K$) on G_j .

Using cross-validation: Step 3

- ① For each β_k , sum the errors of $T^{(j)}(\beta_k)$ over G_j ($j = 1, \dots, v$).
- ② Let β_h be the one with the lowest overall error.
Select T_h as the best tree.
- ③ Use the error rate computed with cross-validation as an estimate of its error rate.

Remark: Alternatively, we could again use the 1-SE rule in the final step to select the final tree from the sequence.

Using cross-validation: Step 1

Tree sequence constructed on *full* data set:

- T_1 is the best tree for $\alpha \in [0, \frac{1}{20})$.
- T_2 is the best tree for $\alpha \in [\frac{1}{20}, \frac{2}{10})$.
- T_3 is the best tree for $\alpha \in [\frac{2}{10}, \infty)$.

Set

$\beta_1 = 0$, value corresponding to T_1

$\beta_2 = \sqrt{\frac{1}{20} \frac{2}{10}} = \frac{1}{10}$, value corresponding to T_2

$\beta_3 = \infty$, value corresponding to T_3 (root).

Using cross-validation: Step 2

Divide the data set in $v = 4$ groups G_1, G_2, G_3, G_4 of size 50 each.

First CV-run

- ① Build a tree on all data except G_1 , and determine the smallest minimizing subtrees $T^{(1)}(0)$, $T^{(1)}(\frac{1}{10})$ and $T^{(1)}(\infty)$.
- ② Compute the error of those trees on G_1 .

Repeat this procedure for G_2 , G_3 and G_4 .

Using cross-validation: Step 3

CV-run	$\beta_1 = 0$	$\beta_2 = \frac{1}{10}$	$\beta_3 = \infty$
1	20	10	25
2	18	8	25
3	22	9	25
4	20	13	25
Total	80	40	100

β_2 wins (40 errors), so T_2 gets selected.

We estimate the error rate of T_2 at 20%.

Building Trees in R: Rpart

Pima Indians Diabetes Database from the UCI ML Repository

1. Number of times pregnant
2. Plasma glucose concentration a 2 hours in an oral glucose tolerance test
3. Diastolic blood pressure (mm Hg)
4. Triceps skin fold thickness (mm)
5. 2-Hour serum insulin (mu U/ml)
6. Body mass index (weight in kg/(height in m)²)
7. Diabetes pedigree function
8. Age (years)
9. Class variable (0 or 1)

Class Distribution: (class value 1 is interpreted as "tested positive for diabetes")

Class	Value	Number of instances
0		500
1		268

Building Trees in R: Rpart

```
> pima.dat[1:5,]
   npreg plasma bp triceps serum  bmi pedigree age class
1      6    148  72      35     0 33.6    0.627  50     1
2      1     85  66      29     0 26.6    0.351  31     0
3      8    183  64      0     0 23.3    0.672  32     1
4      1     89  66      23    94 28.1    0.167  21     0
5      0    137  40      35   168 43.1    2.288  33     1

> library(rpart)
> pima.tree <- rpart(class ~ ., data=pima.dat, cp=0, minbucket=1, minsplit=2, method="class")
> printcp(pima.tree)
```

Classification tree:

```
rpart(formula = class ~ ., data = pima.dat, method = "class",
      cp = 0, minbucket = 1, minsplit = 2)
```

Variables actually used in tree construction:

```
[1] age      bmi      bp      npreg      pedigree plasma      serum      triceps
```

Root node error: 268/768 = 0.34896

n= 768

cptable: the pruning sequence

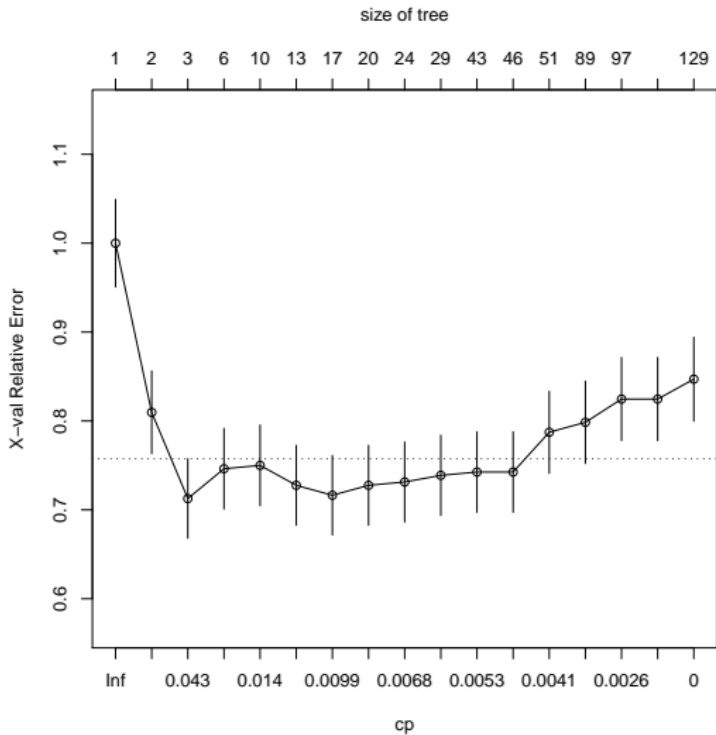
	CP	nsplit	rel error	xerror	xstd
1	0.2425373	0	1.000000	1.000000	0.049288
2	0.1044776	1	0.757463	0.80970	0.046558
3	0.0174129	2	0.652985	0.71269	0.044698
4	0.0149254	5	0.600746	0.74627	0.045381
5	0.0130597	9	0.541045	0.75000	0.045454
6	0.0111940	12	0.492537	0.72761	0.045007
7	0.0087065	16	0.447761	0.71642	0.044776
8	0.0074627	19	0.421642	0.72761	0.045007
9	0.0062189	23	0.391791	0.73134	0.045083
10	0.0055970	28	0.358209	0.73881	0.045233
11	0.0049751	42	0.272388	0.74254	0.045307
12	0.0044776	45	0.257463	0.74254	0.045307
13	0.0037313	50	0.235075	0.78731	0.046159
14	0.0027985	88	0.093284	0.79851	0.046360
15	0.0024876	96	0.070896	0.82463	0.046814
16	0.0018657	109	0.037313	0.82463	0.046814
17	0.0000000	128	0.000000	0.84701	0.047184

CP is α divided by the resubstitution error in the root node.

Example: tree with 2 splits is best for $CP \in [0.0174129, 0.1044776]$.

Tree with 2 splits has cross-validation error of $0.34896 \times 0.71269 = 0.2487$.

Plot of pruning sequence



Selecting the best tree

```
> pima.pruned <- prune(pima.tree, cp=0.02)  
> post(pima.pruned)
```

