

An Introduction to Recursive Partitioning Using the RPART Routines

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1 Introduction

This document is a shortened version of the Mayo Clinic Section of Biostatistics technical report [5] written by Terry Therneau and Beth Atkinson. More complete information about the theory behind the coding can be obtained there. The following pages provide an overview of the methods found in the `rpart` routines, which implement many of the ideas found in the CART (Classification and Regression Trees) book and programs of Breiman, Friedman, Olshen and Stone [1]. Because CART is the trademarked name of a particular software implementation of these ideas, and *tree* has been used for the S-plus routines of Clark and Pregibon~[2] a different acronym — Recursive PARTitioning or `rpart` — was chosen.

The `rpart` programs build classification or regression models of a very general structure using a two stage procedure; the resulting models can be represented as binary trees. The types of endpoints that `rpart` handles includes classifications (such as yes/no), continuous values (such as bone mineral density), poisson counts (such as the number of fractures in Medicare patients), and survival information (time to death/last known contact). The `rpart` library includes tools to model, plot, and summarize the end results.

2 Basic steps needed to use rpart

Step-I. Attach the library so that the functions can be found.

```
library(rpart)
```

Step-II. Decide what type of endpoint you have

- Categorical ==> method == "class"
- Continuous ==> method == "anova"
- Poisson Process/Count ==> method == "poisson"
- Survival ==> method == "exp"

Step-III. Fit the model using the standard Splus modeling language.

```
fit <- rpart(skips ~ Opening + Solder + Mask + PadType  
            + Panel, data=solder, method='poisson')
```

Step-IV. Print a text version of the tree.

```
print(fit)
```

Step-V. Print a summary which examines each node in depth.

```
summary(fit)
```

Step–VI. Plot a standard version of the plot with some basic information.

```
plot(fit)
text(fit,use.n=T)
```

Step–VII. Create a prettier version of the tree.

```
post(fit,file="")
```

3 Rpart model options

This section examines the various options that are available when fitting the model `rpart`. The options are listed below with a brief explanation, then explored further with actual examples.

The central fitting function is `rpart`, whose main arguments are

formula: the model formula, as in `lm` and other S model fitting functions. The right hand side may contain both continuous and categorical (factor) terms. If the outcome y has more than two levels, then categorical predictors must be fit by exhaustive enumeration, which can take a very long time.

data, weights, subset: as for other S models.

method: the type of splitting rule to use. Options at this point are classification, anova, Poisson, and exponential.

parms: a list of method specific optional parameters. Poisson splitting has a single parameter, the coefficient of variation of the prior distribution on the rates (`shrink`). It is used to prevent problems if nodes end up with 0 events. Usually not changed (default=1). For classification, the list can contain any of:

- `prior`– the vector of prior probabilities
- `loss`– the loss matrix
- `split`– the splitting index

The priors must be positive and sum to 1. The loss matrix must have zeros on the diagonal and positive off-diagonal elements. The splitting index can be ‘gini’ or ‘information’.

```

> lmat <- matrix(c(0,4,3,0), nrow=2, ncol=2, byrow=F)
> fit1 <- rpart(Kyphosis ~ Age + Number + Start, data=kyphosis, loss
  method='class', parms=list(split='gini', loss=lmat, shrink=0))
> fit2 <- rpart(Kyphosis ~ Age + Number + Start, data=kyphosis,
  method='class', parms=list(split='gini', prior=
  c(.65,.35)))
> wts <- rep(3, nrow(stagec))
> fit3 <- rpart(Surv(pgtime, pgstat) ~ age + eet + g2+grade+gleason +ploidy,
  stagec, parms=list(shrink=0), method="poisson", weights=wts)

```

na.action: the action for missing values. The default action for rpart is `na.rpart`, this default is not overridden by the `options(na.action)` global option. The default action removes only those rows for which either the response y or *all* of the predictors are missing. This ability to retain partially missing observations is perhaps the single most useful feature of rpart models.

control: a list of control parameters, usually the result of the `rpart.control` function. The list contains

- **minsplit:** The minimum number of observations in a node for which the routine will even try to compute a split. The default is 20. This parameter can save computation time, since smaller nodes are almost always pruned away by cross-validation.
- **minbucket:** The minimum number of observations in a terminal node. This defaults to `minsplit/3`.
- **maxcompete:** It is often useful in the printout to see not only the variable that gave the best split at a node, but also the second, third, etc best. This parameter controls the number that will be printed. It has no effect on computational time, and a small effect on the amount of memory used. The default is 5.
- **xval:** The number of cross-validations to be done. Usually set to zero during exploratory phases of the analysis. A value of 10, for instance, increases the compute time to 11-fold over a value of 0.
- **maxsurrogate:** The maximum number of surrogate variables to retain at each node. (No surrogate that does worse than “go with the majority” is printed or used). Setting this to zero will cut the computation time in half, and set `usesurrogate` to zero. The default is 5. Surrogates give different information than competitor splits. The competitor list asks “which other splits would have as many correct classifications”, surrogates ask “which other splits would classify the same subjects in the same way”, which is a harsher criteria.

- **usesurrogate:** A value of `usesurrogate=2`, the default, splits subjects in the way described previously. This is similar to CART. If the value is 0, then a subject who is missing the primary split variable does not progress further down the tree. A value of 1 is intermediate: all surrogate variables except “go with the majority” are used to send a case further down the tree.
- **cp:** The threshold complexity parameter.


```
> temp <- rpart.control(xval=10, minbucket=2, minsplit=4, cp=0)
> dfit <- rpart(y ~ x, method='class', control=temp)
```

4 Plotting options

This section examines the various options that are available when plotting an `rpart` object. The options are listed below with a brief explanation, then explored further with actual plots.

4.1 `plot.rpart` options

The function `plot.rpart` plots an `rpart` object on the current graphics device. The main arguments to the function are

Required Arguments: arguments you must specify.

tree: a fitted object of class `rpart`, containing a classification, regression, or rate tree.

Optional Arguments: arguments you can add if you wish.

uniform: if `TRUE`, uniform vertical spacing of the nodes is used; this may be less cluttered when fitting a large plot onto a page. The default is to use a non-uniform spacing proportional to the error in the fit.

branch: controls the shape of the branches from parent to child node. Any number from 0 to 1 is allowed. A value of 1 gives square shouldered branches, a value of 0 give V shaped branches, with other values being intermediate.

compress: if `FALSE`, the leaf nodes will be at the horizontal plot coordinates of `1:nleaves` (like `plot.tree`). If `TRUE`, the routine attempts a more compact arrangement of the tree. The compaction algorithm assumes `uniform=T`.

nospace: the amount of extra space between a node with children and a leaf, as compared to the minimal space between leaves. Applies to compressed trees only. The default is the value of `branch`.

margin: an extra percentage of white space to leave around the borders of the tree. (Long labels sometimes get cut off by the default computation).

minbranch: set the minimum length for a branch to `minbranch` times the average branch length. This parameter is ignored if `uniform=T`. Sometimes a split will give very little improvement, or even (in the classification case) no improvement at all. A tree with branch lengths strictly proportional to improvement leaves no room to squeeze in node labels.

4.2 `text.rpart` options

The function `text.rpart` labels the current plot of the tree dendrogram with text. The main arguments to this function are

Required Arguments: arguments you must specify.

x: fitted model object of class `rpart`. This is assumed to be the result of some function that produces an object with the same named components as that returned by the `rpart` function.

Optional Arguments: arguments you can add if you wish.

splits: logical flag. If `TRUE` (default), then the splits in the tree are labeled with the criterion for the split.

label: a column name of `x$frame`; values of this will label the nodes.

FUN: the name of a labeling function, e.g. `text`.

all: Logical. If `TRUE`, all nodes are labeled, otherwise just terminal nodes.

pretty: an integer denoting the extent to which factor levels in split labels will be abbreviated. A value of (0) signifies no abbreviation. A `NULL` (default) signifies using elements of letters to represent the different factor levels.

digits: number of significant digits to include in numerical labels.

use.n: Logical. If `TRUE`, adds to label (`#events level1/ #events level2/etc.` for class, `n` for anova, and `# events/n` for poisson and exp).

fancy: Logical. If `TRUE`, nodes are represented by ellipses (interior nodes) and rectangles (leaves) and labeled by `yval`. The edges connecting the nodes are labeled by left and right splits.

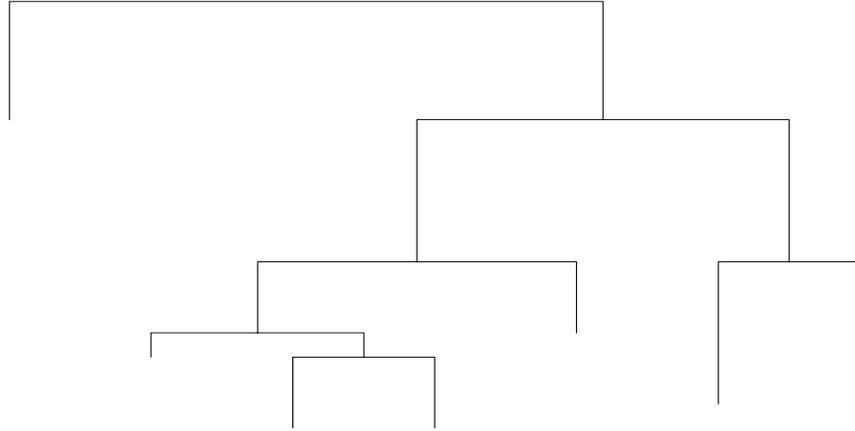


Figure 1: `plot(fit); text(fit)`

fwidth: Relates to option `fancy` and the width of the ellipses and rectangles. If `fwidth < 1` then it is a scaling factor (default = .8). If `fwidth > 1` then it represents the number of character widths (for current graphical device) to use.

fheight: Relates to option `fancy` and the height of the ellipses and rectangles. If `fheight < 1` then it is a scaling factor (default = .8). If `fheight > 1` then it represents the number of character heights (for current graphical device) to use.

4.3 Examples using `plot.rpart` and `text.rpart` together

For simplicity, the same model will be used throughout the subsection. The simplest labelled plot (figure 1) is called by using `plot` and `text` without changing any of the defaults. This is useful for a first look, but sometimes you'll want more information about each of the nodes.

```
> fit <- rpart(progstat ~ age + eet + g2 + grade + gleason + ploidy,
               stagec, control=rpart.control(cp=.025))

> plot(fit)
> text(fit)
```

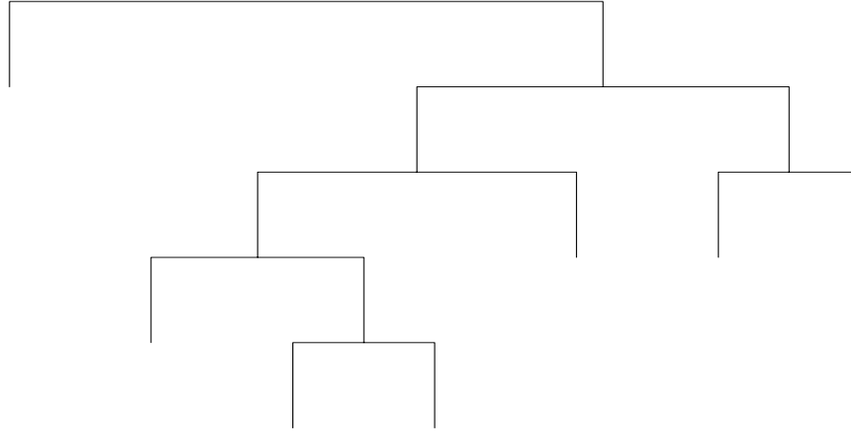


Figure 2: `plot(fit, uniform=T); text(fit,use.n=T,all=T)`

The next plot (figure 2) has uniform stem lengths (`uniform=T`), has extra information (`use.n=T`) specifying number of subjects at each node (here it lists how many are diseased and how many are not diseased), and has labels on all the nodes, not just the terminal nodes (`all=T`).

```
> plot(fit, uniform=T)
> text(fit, use.n=T, all=T)
```

Fancier plots can be created by modifying the `branch` option, which controls the shape of branches that connect a node to its children. The default for the plots is to have square shouldered trees (`branch = 1.0`). This can be taken to the other extreme with no shoulders at all (`branch=0`) as shown in figure 3.

```
> plot(fit, branch=0)
> text(fit, use.n=T)
```

These options can be combined with others to create the plot that fits your particular needs. The default plot may be inefficient in its use of space: the terminal nodes will always lie at x-coordinates of 1,2,... The `compress` option attempts to improve this by overlapping some nodes. It has little effect on figure 4, but in figure 3 it allows the lowest branch to “tuck under” those above. If you want to play around with the spacing with `compress`, try using `nospace` which regulates the space between a terminal node and a split.

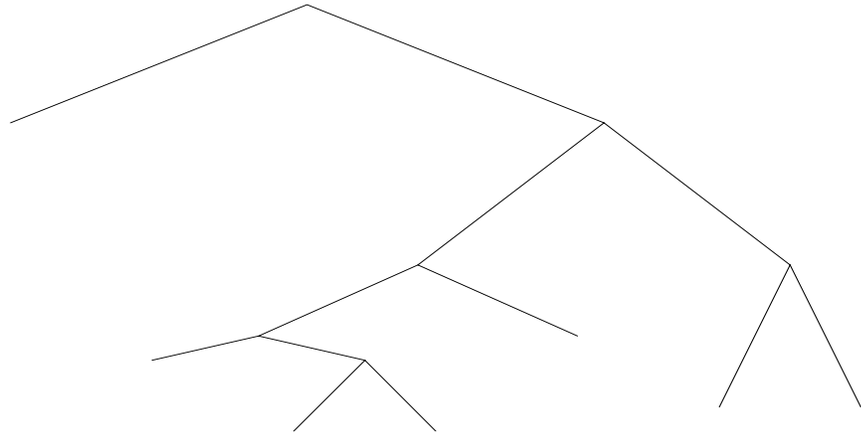


Figure 3: `plot(fit, branch=0); text(fit,use.n=T)`

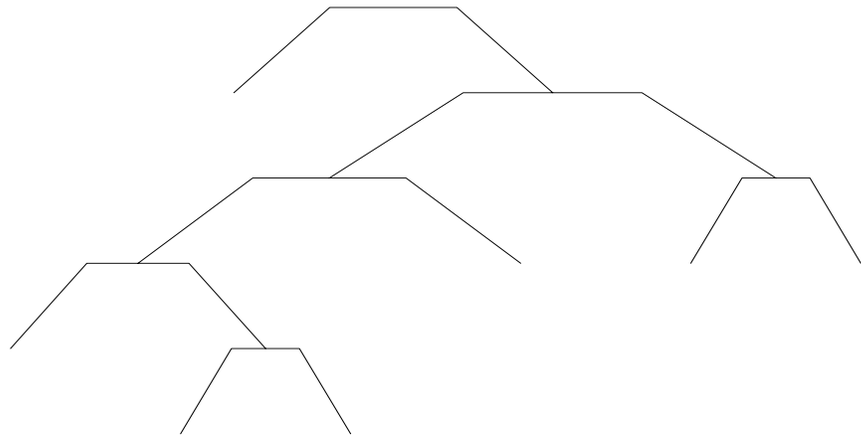


Figure 4: `plot(fit, branch=.4, uniform=T,compress=T)`

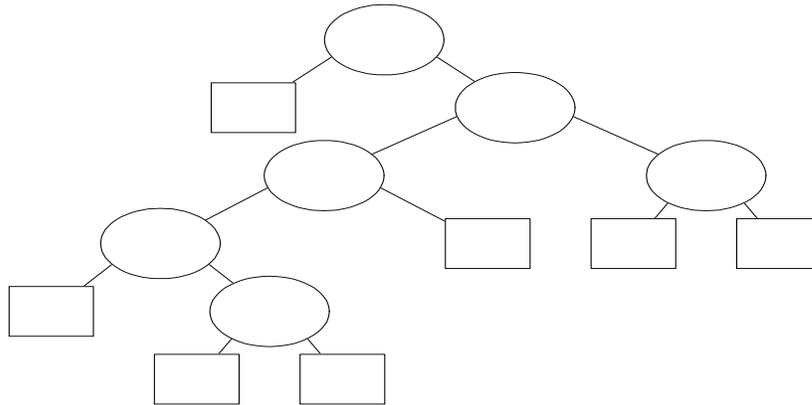


Figure 5: `post(fit)`

```
> plot(fit,branch=.4,uniform=T,compress=T)
> text(fit,all=T,use.n=T)
```

We have combined several of these options into a function called `post.rpart`. Results are shown in figure 5. The code is essentially

```
> plot(tree, uniform = T, branch = 0.2, compress = T, margin = 0.1)
> text(tree, all = T, use.n=T, fancy = T)
```

The `fancy` option of `text` creates the ellipses and rectangles, and moves the splitting rule to the midpoints of the branches. `Margin` shrinks the plotting region slightly so that the `text` boxes don't run over the edge of the plot. The `branch` option makes the lines exit the ellipse at a "good" angle. The call `post(fit)` will create a postscript file `fit.ps` in the current directory. The additional argument `file=""` will cause the plot to appear on the current active device. Note that `post.rpart` is just our choice of options to the functions `plot.rpart` and `text.rpart`. The reader is encouraged to try out other possibilities, as these specifications may not be the best choice in all situations.

5 Other available functions

There are a number of other functions in the `rpart` library. This list describes the all the main `rpart` functions briefly. More details about the functions can be found on the help pages in Splus (ex. `help(snip.rpart)`) or elsewhere in this document.

as.tree Create a tree object from an `rpart` object.

meanvar.rpart Mean-Variance plot for an `rpart` object.

rsq.rpart Plots R-square versus number of splits.

resid.rpart Residuals from an `rpart` object.

predict.rpart Predictions from an `rpart` object.

plot.rpart Plot an `rpart` object.

plotcp Plot the Complexity Parameter (`cp`) for an `rpart` object.

post.rpart Plot an `rpart` object - prettier version.

print.rpart Print a `rpart` object.

printcp Print the Complexity Parameter (`cp`) table for an `rpart` object.

prune.rpart Prune back an `rpart` object given a complexity parameter value ranging from 0 to 1.

snip.rpart Snip subtrees of an `rpart` object.

summary.rpart Summarize an `rpart` object.

text.rpart Place text on a dendrogram.

xpred.rpart Cross-validation for `rpart` object.

There are also some `tree` functions that may work for `rpart` objects, once the function `as.tree` has been applied to it. These functions are

hist.tree Augment a dendrogram with histograms.

rug.tree Augment a dendrogram with a rug.

tile.tree Augment a dendrogram with tiles.

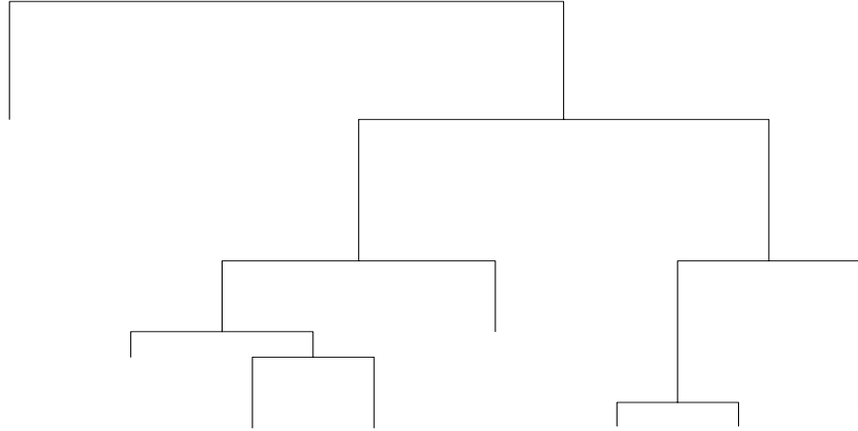


Figure 6: Classification tree for the Stage C data

6 Examples with explanation of output

6.1 Stage C prostate cancer (class method)

This first example is based on a data set of 146 stage C prostate cancer patients [4]. The main clinical endpoint of interest is whether the disease recurs after initial surgical removal of the prostate, and the time interval to that progression (if any). The endpoint in this example is `status`, which takes on the value 1 if the disease has progressed and 0 if not. Later we'll analyze the data using the `exp` method, which will take into account time to progression. A short description of each of the variables is listed below. The main predictor variable of interest in this study was DNA ploidy, as determined by flow cytometry. For diploid and tetraploid tumors, the flow cytometric method was also able to estimate the percent of tumor cells in a G_2 (growth) stage of their cell cycle; $G_2\%$ is systematically missing for most aneuploid tumors.

The variables in the data set are

pgtime	time to progression, or last follow-up free of progression
pgstat	status at last follow-up (1=progressed, 0=censored)
age	age at diagnosis
eet	early endocrine therapy (1=no, 0=yes)
ploidy	diploid/tetraploid/aneuploid DNA pattern
g2	% of cells in G_2 phase
grade	tumor grade (1-4)
gleason	Gleason grade (3-10)

6.1.1 print function

The model is fit by using the `rpart` function. The first argument of the function is a model formula, with the `~` symbol standing for “is modeled as”. The `print` function gives an abbreviated output, as for other S models. The `plot` and `text` command plot the tree and then label the plot, the result is shown in figure 6.

```
> progstat <- factor(stagec$pgstat, levels=0:1, labels=c("No", "Prog"))
> cfit <- rpart(progstat ~ age + eet + g2 + grade + gleason + ploidy,
               data=stagec, method='class')
> print(cfit)
```

```
node), split, n, loss, yval, (yprob)
* denotes terminal node
```

```
1) root 146 54 No ( 0.6301 0.3699 )
 2) grade<2.5 61 9 No ( 0.8525 0.1475 ) *
 3) grade>2.5 85 40 Prog ( 0.4706 0.5294 )
 6) g2<13.2 40 17 No ( 0.5750 0.4250 )
 12) ploidy:diploid,tetraploid 31 11 No ( 0.6452 0.3548 )
    24) g2>11.845 7 1 No ( 0.8571 0.1429 ) *
    25) g2<11.845 24 10 No ( 0.5833 0.4167 )
        50) g2<11.005 17 5 No ( 0.7059 0.2941 ) *
        51) g2>11.005 7 2 Prog ( 0.2857 0.7143 ) *
 13) ploidy:aneuploid 9 3 Prog ( 0.3333 0.6667 ) *
 7) g2>13.2 45 17 Prog ( 0.3778 0.6222 )
 14) g2>17.91 22 8 No ( 0.6364 0.3636 )
    28) age>62.5 15 4 No ( 0.7333 0.2667 ) *
    29) age<62.5 7 3 Prog ( 0.4286 0.5714 ) *
    15) g2<17.91 23 3 Prog ( 0.1304 0.8696 ) *
```

```
> plot(cfit)
> text(cfit)
```

- The creation of a labeled factor variable as the response improves the labeling of the printout.
- We have explicitly directed the routine to treat progstat as a categorical variable by asking for method='class'. (Since progstat is a factor this would have been the default choice). Since no optional classification parameters are specified the routine will use the Gini rule for splitting, prior probabilities that are proportional to the observed data frequencies, and 0/1 losses.
- The child nodes of node x are always numbered $2x$ (left) and $2x + 1$ (right), to help in navigating the printout (compare the printout to figure 6).
- Other items in the list are the definition of the variable and split used to create a node, the number of subjects at the node, the loss or error at the node (for this example, with proportional priors and unit losses this will be the number misclassified), and the predicted class for the node.
- * indicates that the node is terminal.
- Grades 1 and 2 go to the left, grades 3 and 4 go to the right. The tree is arranged so that the branches with the largest “average class” go to the right.

6.1.2 printcp function

```
> printcp(cfit)
```

```
Classification tree:
```

```
rpart(formula = progstat ~ age + eet + g2 + grade + gleason + ploidy, data =
      stagec, method = "class")
```

```
Variables actually used in tree construction:
```

```
[1] age    g2     grade  ploidy
```

```
Root node error: 54/146 = 0.36986
```

```
n= 146
```

	CP	nsplit	rel error	xerror	xstd
1	0.104938	0	1.00000	1.0000	0.10802
2	0.055556	3	0.68519	1.1852	0.11103
3	0.027778	4	0.62963	1.0556	0.10916
4	0.018519	6	0.57407	1.0556	0.10916
5	0.010000	7	0.55556	1.0556	0.10916

```
> fitc2 <- prune(cfit,cp=.03)
```

The `cptable` provides a brief summary of the overall fit of the model.

- The table is printed from the smallest tree (no splits) to the largest one (7 splits). We find it easier to compare one tree to another when they start at the same place.
- The number of splits is listed, rather than the number of nodes. The number of nodes is always $1 +$ the number of splits.
- For easier reading, the error columns have been scaled so that the first node has an error of 1. Since in this example the model with no splits must make 54/146 misclassifications, multiply columns 3-5 by 54 to get a result in terms of absolute error. (Computations are done on the absolute error scale, and printed on relative scale).
- The complexity parameter (`cp`) column has been similarly scaled.

Looking at the table, we see that the best tree has 5 terminal nodes (4 splits), based on cross-validation. There is a 1-SE rule used to find the best number of splits which takes the smallest cross validation error (`xerror`), adds the corresponding standard error (`xstd`), and finds the fewest cross-validation error that is smaller than this number. Here the 1-SE rule is $1.0556 + 0.10916$, or 1.16476 . This subtree is extracted with a call to `prune` and is saved in `cfit2`.

6.1.3 summary function

For a more detailed listing of the `rpart` object, we use the `summary` function. It includes the information from the CP table (not repeated below), plus information about each node. It is easy to print a subtree based on a different `cp` value using the `cp` option. Any value between 0.0555 and 0.1049 would produce the same result as is listed below, that is, the tree with 3 splits. Because the printout is long, the `file` option of `summary.rpart` is often useful (prints output directly to a file).

```
> summary(cfit,cp=.06)
```

```
Node number 1: 146 observations,    complexity param=0.1049
  predicted class= No  expected loss= 0.3699
    class counts:  92 54
    probabilities: 0.6301 0.3699
  left son=2 (61 obs) right son=3 (85 obs)
```

```

Primary splits:
  grade < 2.5  to the left,  improve=10.360, (0 missing)
  gleason < 5.5 to the left,  improve= 8.400, (3 missing)
  ploidy splits as LRR,      improve= 7.657, (0 missing)
  g2 < 13.2  to the left,  improve= 7.187, (7 missing)
  age < 58.5 to the right, improve= 1.388, (0 missing)
Surrogate splits:
  gleason < 5.5  to the left,  agree=0.863, adj=0.672, (0 split)
  ploidy splits as LRR,      agree=0.644, adj=0.148, (0 split)
  g2 < 9.945 to the left,  agree=0.630, adj=0.115, (0 split)
  age < 66.5  to the right, agree=0.589, adj=0.016, (0 split)

```

```

Node number 2: 61 observations
  predicted class= No  expected loss= 0.1475
  class counts:  52  9
  probabilities:  0.8525 0.1475

```

```

Node number 3: 85 observations,  complexity param=0.1049
  predicted class= Prog  expected loss= 0.4706
  class counts:  40 45
  probabilities:  0.4706 0.5294
  left son=6 (40 obs) right son=7 (45 obs)

```

```

Primary splits:
  g2 < 13.2  to the left,  improve=2.1780, (6 missing)
  ploidy splits as LRR,      improve=1.9830, (0 missing)
  age < 56.5 to the right, improve=1.6600, (0 missing)
  gleason < 8.5  to the left, improve=1.6390, (0 missing)
  eet < 1.5  to the right, improve=0.1086, (1 missing)
Surrogate splits:
  ploidy splits as LRL,      agree=0.962, adj=0.914, (6 split)
  age < 68.5 to the right, agree=0.608, adj=0.114, (0 split)
  gleason < 6.5  to the left, agree=0.582, adj=0.057, (0 split)
.
.
.

```

- There are 54 progressions (class 1) and 92 non-progressions, so the first node has an expected loss of $54/146 \approx 0.37$. (The computation is this simple only for the default priors and losses).
- Grades 1 and 2 go to the left, grades 3 and 4 to the right. The tree is arranged so that the “more severe” nodes go to the right.

- The improvement is n times the change in impurity index. In this instance, the largest improvement is for the variable `grade`, with an improvement of 10.36. The next best choice is Gleason score, with an improvement of 8.4. The actual values of the improvement are not so important, but their relative size gives an indication of the comparative utility of the variables.
- At node 3, `g2` is missing for 6 observations. 96.2% of the observations that have both ploidy and `g2` agree at their respective splits, hence ploidy is chosen as the best surrogate for `g2`. The `adj` indicates how much is gained over simply choosing a "go with the majority" rule.
- Ploidy is a categorical variable, with values of diploid, tetraploid, and aneuploid, in that order. (To check the order, type `table(stagec$ploidy)`). All three possible splits were attempted: aneuploid+diploid vs. tetraploid, aneuploid+tetraploid vs. diploid, and aneuploid vs. diploid + tetraploid. The best split sends diploid to the right and the others to the left (node 6, see figure (6)).
- For node 3, the primary split variable is missing on 6 subjects. All 6 are split based on the first surrogate, ploidy. Diploid and aneuploid tumors are sent to the left, tetraploid to the right. $76/79 = .962$ of the observations that have both ploidy and `g2` agree at their respective splits, hence ploidy is chosen as the best surrogate for `g2`.

The `adj` indicates how much is gained beyond "go with the majority" ($= 44/79$). `Adj` is calculated as $(76/79 - 44/79)/(1 - 44/79)$.

	<code>g2 < 13.2</code>	<code>g2 > 13.2</code>	NA
Diploid/aneuploid	33	2	5
Tetraploid	1	43	1

6.2 Consumer Report Auto data (anova method)

The `anova` method leads to regression trees; it is the default method if y a simple numeric vector, i.e., not a factor, matrix, or survival object.

The dataset `car.a11` contains a collection of variables from the April, 1990 Consumer Reports; it has 36 variables on 111 cars. Documentation may be found in the S-Plus manual. We will work with a subset of 23 of the variables, created by the first two lines of the example below. We will use `Price` as the response. This data set is a good example of the usefulness of the missing value logic in `rpart`: most of the variables are missing on only 3-5 observations, but only 42/111 have a complete subset.

6.2.1 print, printcp functions

```
> cars <- car.all[, c(1:12, 15:17, 21, 28, 32:36)]
> cars$Eng.Rev <- as.numeric(as.character(car.all$Eng.Rev2))
> fit3 <- rpart(Price ~ ., data=cars)
> fit3
n=105 (6 observations deleted due to missing)

node), split, n, deviance, yval
  * denotes terminal node

1) root 105 7.118e+09 15810
 2) Disp.<156 70 1.492e+09 11860
   4) Country:Brazil,Japan,Japan/USA,Korea,Mexico,USA 58 4.212e+08 10320
     8) Type:Small 21 5.031e+07 7629 *
     9) Type:Compact,Medium,Sporty,Van 37 1.328e+08 11840 *
   5) Country:France,Germany,Sweden 12 2.707e+08 19290 *
 3) Disp.>156 35 2.351e+09 23700
   6) HP.revs<5550 24 9.803e+08 20390
     12) Disp.<267.5 16 3.960e+08 17820 *
     13) Disp.>267.5 8 2.676e+08 25530 *
   7) HP.revs>5550 11 5.316e+08 30940 *

> printcp(fit3)

Regression tree:
rpart(formula = Price ~ ., data = cars)

Variables actually used in tree construction:
[1] Country Disp. HP.revs Type

Root node error: 7.1183e9/105 = 6.7793e7

n=105 (6 observations deleted due to missing)

      CP nsplit rel error  xerror  xstd
1 0.460146      0  1.00000 1.02413 0.16411
2 0.117905      1  0.53985 0.79225 0.11481
3 0.044491      3  0.30961 0.60042 0.10809
4 0.033449      4  0.26511 0.58892 0.10621
5 0.010000      5  0.23166 0.57062 0.11782
```

Only 4 of 22 predictors were actually used in the fit: engine displacement in cubic inches, country of origin, type of vehicle, and the revolutions for maximum horsepower (the “red line” on a tachometer).

- The relative error is $1 - R^2$, similar to linear regression. The xerror is related to the PRESS statistic. The first split appears to improve the fit the most. The last split adds little improvement to the apparent error.
- The 1-SE rule would choose a tree with 3 splits.
- This is a case where the default cp value of .01 may have overpruned the tree, since the cross-validated error is not yet at a minimum. A rerun with the cp threshold at .002 gave a maximum tree size of 8 splits, with a minimum cross-validated error for the 5 split model.
- For any CP value between 0.46015 and 0.11791 the best model has one split; for any CP value between 0.11791 and 0.04449 the best model is with 3 splits; and so on.
- In the anova method the splitting criteria is $SS_T - (SS_L + SS_R)$, where $SS_T = \sum (y_i - \bar{y})^2$ is the sum of squares for the node, and SS_R, SS_L are the sums of squares for the right and left son, respectively. This is equivalent to choosing the split to maximize the between-groups sum-of-squares in a simple analysis of variance. This rule is identical to the regression option for `tree`.
- A summary statistic or vector, which is used to describe a node. The first element of the vector is considered to be the fitted value. For the anova method this is the mean of the node.
- The prediction error for a new observation, assigned to the node, is $(y_{new} - \bar{y})$.

The `print` command also recognizes the `cp` option, which allows the user to see which splits are the most important.

```
> print(fit3,cp=.10)
n=105 (6 observations deleted due to missing)

node), split, n, deviance, yval
  * denotes terminal node

1) root 105 7.118e+09 15810
 2) Disp.<156 70 1.492e+09 11860
   4) Country:Brazil,Japan,Japan/USA,Korea,Mexico,USA 58 4.212e+08 10320 *
   5) Country:France,Germany,Sweden 12 2.707e+08 19290 *
 3) Disp.>156 35 2.351e+09 23700
   6) HP.revs<5550 24 9.803e+08 20390 *
   7) HP.revs>5550 11 5.316e+08 30940 *
```

The first split on displacement partitions the 105 observations into groups of 70 and 35 (nodes 2 and 3) with mean prices of 11,860 and 23,700. The deviance (corrected sum-of-squares) at these 2 nodes are 1.49×10^9 and 2.35×10^9 , respectively. More detailed summarization of the splits is again obtained by using the function `summary.rpart`.

6.2.2 summary function

```
> summary(fit3, cp=.10)
Node number 1: 105 observations,    complexity param=0.4601
mean=15810, MSE=67790000
left son=2 (70 obs) right son=3 (35 obs)
Primary splits:
  Disp.    < 156  to the left,  improve=0.4601, (0 missing)
  HP       < 154  to the left,  improve=0.4549, (0 missing)
  Tank     < 17.8 to the left,  improve=0.4431, (0 missing)
  Weight   < 2890 to the left,  improve=0.3912, (0 missing)
  Wheel.base < 104.5 to the left, improve=0.3067, (0 missing)
Surrogate splits:
  Weight   < 3095 to the left,  agree=0.914, adj=0.743, (0 split)
  HP       < 139  to the left,  agree=0.895, adj=0.686, (0 split)
  Tank     < 17.95 to the left, agree=0.895, adj=0.686, (0 split)
  Wheel.base < 105.5 to the left, agree=0.857, adj=0.571, (0 split)
  Length   < 185.5 to the left, agree=0.838, adj=0.514, (0 split)

Node number 2: 70 observations,    complexity param=0.1123
mean=11860, MSE=21310000
left son=4 (58 obs) right son=5 (12 obs)
Primary splits:
  Country splits as L-RRLLLLRL, improve=0.5361, (0 missing)
  Tank    < 15.65 to the left,  improve=0.3805, (0 missing)
  Weight  < 2568 to the left,  improve=0.3691, (0 missing)
  Type    splits as R-RLRR,    improve=0.3650, (0 missing)
  HP      < 105.5 to the left,  improve=0.3578, (0 missing)
Surrogate splits:
  Tank    < 17.8 to the left,  agree=0.857, adj=0.167, (0 split)
  Rear.Seating < 28.75 to the left, agree=0.843, adj=0.083, (0 split)
  .
  .
  .
```

- The improvement listed is the percent change in sums of squares (SS) for this split, i.e., $1 - (SS_{right} + SS_{left})/SS_{parent}$.

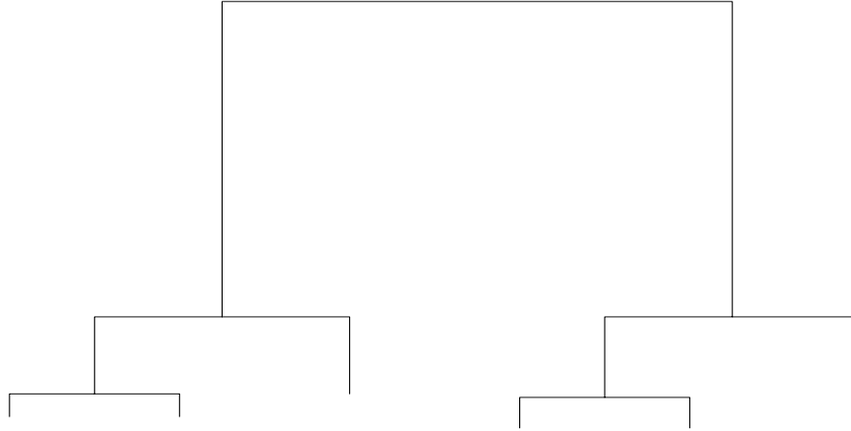


Figure 7: A *anova tree* for the *car.test.frame* dataset. The label of each node indicates the mean *Price* for the cars in that node.

- The weight and displacement are very closely related, as shown by the surrogate split agreement of 91%.
- Not all types are represented in node 2, e.g., there are no representatives from England (the second category). This is indicated by a - in the list of split directions.

6.2.3 plot, text, rsq.rpart functions

```
> plot(fit3)
> text(fit3,use.n=T)
```

As always, a plot of the fit is useful for understanding the *rpart* object. In this plot, we use the option `use.n=T` to add the number of cars in each node. (The default is for only the mean of the response variable to appear). Each individual split is ordered to send the less expensive cars to the left.

Other plots can be used to help determine the best *cp* value for this model. The function `rsq.rpart` plots the jackknifed error versus the number of splits. Of interest is the smallest error, but any number of splits within the “error bars” (1-SE rule) are considered a reasonable number of splits (in this case, 1 or 3 splits seem to be sufficient). As is often true with modelling, simpler is usually better. Another

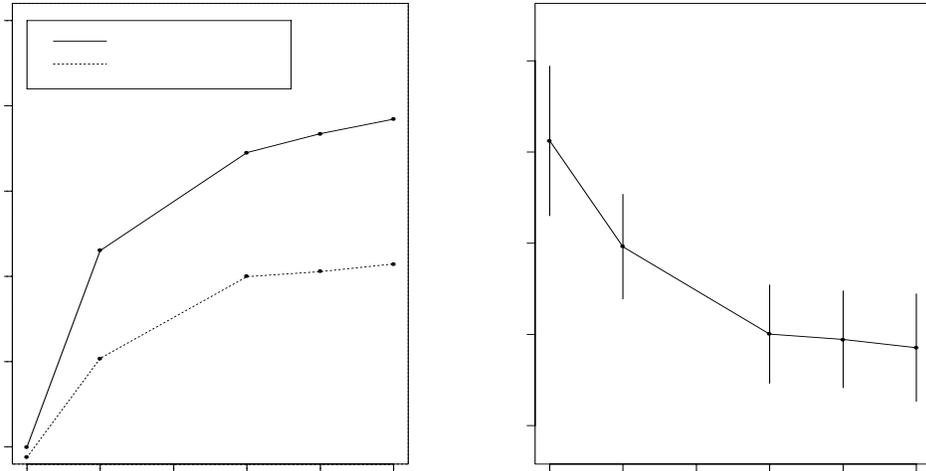


Figure 8: *Both plots were obtained using the function `rsq.rpart(fit3)`. The figure on the left shows that the first split offers the most information. The figure on the right suggests that the tree should be pruned to include only 1 or 2 splits.*

useful plot is the R^2 versus number of splits. The (1 - apparent error) and (1 - relative error) show how much is gained with additional splits. This plot highlights the differences between the R^2 values (figure 9).

Finally, it is possible to look at the residuals from this model, just as with a regular linear regression fit, as shown in the following figure.

```
> plot(predict(fit3),resid(fit3))
> axis(3,at=fit3$frame$yval[fit3$frame$var=='<leaf>'],
      labels=row.names(fit3$frame)[fit3$frame$var=='<leaf>'])
> mtext('leaf number',side=3, line=3)
> abline(h=0)
```

6.3 Solder data (poisson method)

The solder data frame, as explained in the Splus help file, is a design object with 900 observations, which are the results of an experiment varying 5 factors relevant to the wave-soldering procedure for mounting components on printed circuit boards. The response variable, `skips`, is a count of how many solder skips appeared to a visual inspection. The other variables are listed below:

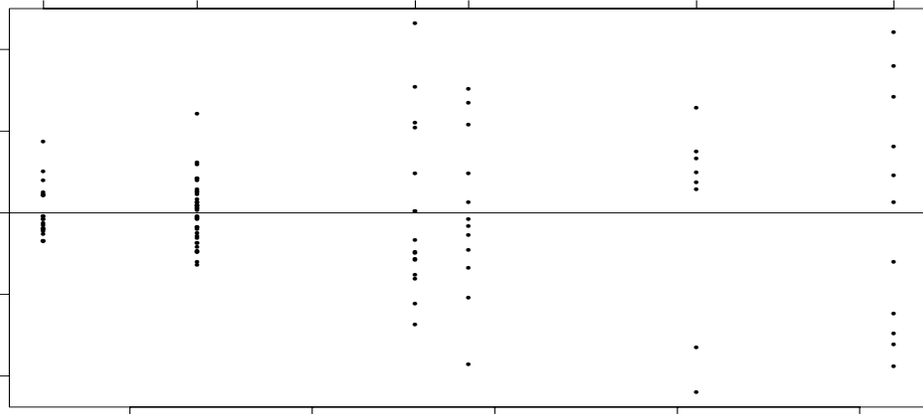


Figure 9: *This plot shows the (observed-expected) cost of cars versus the predicted cost of cars based on the nodes/leaves in which the cars landed. There appears to be more variability in node 7 than in some of the other leaves.*

Opening factor: amount of clearance around the mounting pad (S < M < L)
 Solder factor: amount of solder used (Thin < Thick)
 Mask factor: Type of solder mask used (5 possible)
 PadType factor: Mounting pad used (10 possible)
 Panel factor: panel (1, 2 or 3) on board being counted

In this call, the `rpart.control` options are modified: `maxcompete = 2` means that only 2 other competing splits are listed (default is 4); `cp = .05` means that a smaller tree will be built initially (default is .01). The y variable for Poisson partitioning may be a two column matrix containing the observation time in column 1 and the number of events in column 2, or it may be a vector of event counts alone.

```
fit <- rpart(skips ~ Opening + Solder + Mask + PadType
            + Panel, data=solder, method='poisson',
            control=rpart.control(cp=.05, maxcompete=2))
```

6.3.1 print function

The print command summarizes the tree, as in the previous examples.

```
n= 900

node), split, n, deviance, yval
  * denotes terminal node

1) root 900 8788.0 5.530
 2) Opening:M,L 600 2957.0 2.553
   4) Mask:A1.5,A3,B3 420 874.4 1.033 *
   5) Mask:A6,B6 180 953.1 6.099 *
 3) Opening:S 300 3162.0 11.480
   6) Mask:A1.5,A3 150 652.6 4.535 *
   7) Mask:A6,B3,B6 150 1155.0 18.420 *
```

- The response value is the expected event rate (with a time variable), or in this case the expected number of skips. The values are shrunk towards the global estimate of 5.530 skips/observation.
- The deviance is the same as the null deviance (sometimes called the residual deviance) that you'd get when calculating a Poisson glm model for the given subset of data.
- The splitting rule is based on the likelihood ratio test for two Poisson groups

$$D_{\text{parent}} - (D_{\text{left son}} + D_{\text{right son}})$$

6.3.2 summary function

```
> summary(fit,cp=.10)
```

Call:

```
rpart(formula = skips ~ Opening + Solder + Mask + PadType + Panel, data =
      solder, method = "poisson", control = rpart.control(cp = 0.05,
      maxcompete = 2))
n= 900
```

	CP	nsplit	rel error	xerror	xstd
1	0.3038	0	1.0000	1.0051	0.05248
2	0.1541	1	0.6962	0.7016	0.03299
3	0.1285	2	0.5421	0.5469	0.02544
4	0.0500	3	0.4137	0.4187	0.01962

Node number 1: 900 observations, complexity param=0.3038
events=4977, estimated rate=5.53 , mean deviance=9.765
left son=2 (600 obs) right son=3 (300 obs)

Primary splits:

Opening splits as RLL, improve=2670, (0 missing)
Mask splits as LLRLR, improve=2162, (0 missing)
Solder splits as RL, improve=1168, (0 missing)

Node number 2: 600 observations, complexity param=0.1285
events=1531, estimated rate=2.553 , mean deviance=4.928
left son=4 (420 obs) right son=5 (180 obs)

Primary splits:

Mask splits as LLRLR, improve=1129.0, (0 missing)
Opening splits as RRL, improve= 250.8, (0 missing)
Solder splits as RL, improve= 219.8, (0 missing)

Node number 3: 300 observations, complexity param=0.1541
events=3446, estimated rate=11.48 , mean deviance=10.54
left son=6 (150 obs) right son=7 (150 obs)

Primary splits:

Mask splits as LLRRR, improve=1354.0, (0 missing)
Solder splits as RL, improve= 976.9, (0 missing)
PadType splits as RRRRLRLRL, improve= 313.2, (0 missing)

Surrogate splits:

Solder splits as RL, agree=0.6, adj=0.2, (0 split)

Node number 4: 420 observations
events=433, estimated rate=1.033 , mean deviance=2.082

Node number 5: 180 observations
events=1098, estimated rate=6.099 , mean deviance=5.295

Node number 6: 150 observations
events=680, estimated rate=4.535 , mean deviance=4.351

Node number 7: 150 observations
events=2766, estimated rate=18.42 , mean deviance=7.701

- The improvement is $\text{Deviance}_{\text{parent}} - (\text{Deviance}_{\text{left}} + \text{Deviance}_{\text{right}})$, which is the likelihood ratio test for comparing two Poisson samples.
- The cross-validated error has been found to be overly pessimistic when describing how much the error is improved by each split. This is likely an effect of the boundary effect mentioned earlier, but more research is needed.

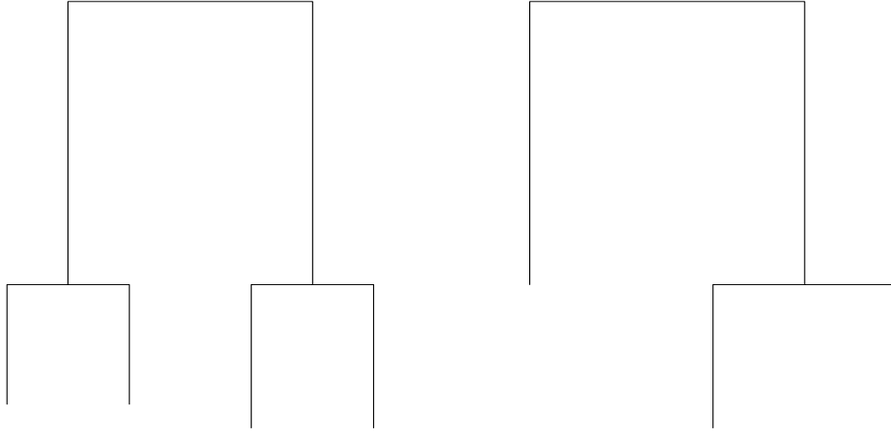


Figure 10: *The first figure shows the solder data, fit with the `poisson` method, using a `cp` value of 0.05. The second figure shows the same fit, but with a `cp` value of 0.15. The function `prune.rpart` was used to produce the smaller tree.*

- The variation `xstd` is not as useful, given the bias of `xerror`.

6.3.3 `plot`, `text`, `prune` functions

```
> plot(fit)
> text(fit,use.n=T)

> fit.prune <- prune(fit,cp=.15)
> plot(fit.prune)
> text(fit.prune,use.n=T)
```

The `use.n=T` option specifies that number of events / total N should be listed along with the predicted rate (number of events/person-years). The function `prune` trims the tree fit to the `cp` value 0.15. The same tree could have been created by specifying `cp = .15` in the original call to `rpart`.

6.4 Stage C Prostate cancer (survival method)

One special case of the Poisson model is of particular interest for medical consulting (such as the authors do). Assume that we have survival data, i.e., each subject has

either 0 or 1 event. Further, assume that the time values have been pre-scaled so as to fit an exponential model. That is, stretch the time axis so that a Kaplan-Meier plot of the data will be a straight line when plotted on the logarithmic scale. An approximate way to do this is

```
temp <- coxph(Surv(time, status) ~1)
newtime <- predict(temp, type='expected')
```

and then do the analysis using the `newtime` variable. (This replaces each time value by $\Lambda(t)$, where Λ is the cumulative hazard function).

A slightly more sophisticated version of this which we will call *exponential scaling* gives a straight line curve for $\log(\text{survival})$ under a parametric exponential model. The only difference from the approximate scaling above is that a subject who is censored between observed death times will receive “credit” for the intervening interval, i.e., we assume the baseline hazard to be linear between observed deaths. If the data is pre-scaled in this way, then the Poisson model above is equivalent to the *local full likelihood* tree model of LeBlanc and Crowley [3]. They show that this model is more efficient than the earlier suggestion of Therneau et. al. [6] to use the martingale residuals from a Cox model as input to a regression tree (anova method). Exponential scaling or `method='exp'` is the default if `y` is a `Surv` object.

Let us again return to the stage C cancer example. Besides the variables explained previously we will use `pgtime`, which is time to tumor progression.

```
> fit <- rpart(Surv(pgtime, pgstat) ~ age + eet + g2 + grade +
gleason + ploidy, data=stagec)
> print(fit)
```

```
n= 146
```

```
node), split, n, deviance, yval
* denotes terminal node
```

```
1) root 146 195.30 1.0000
 2) grade<2.5 61 44.98 0.3617
  4) g2<11.36 33 9.13 0.1220 *
  5) g2>11.36 28 27.70 0.7341
    10) gleason<5.5 20 14.30 0.5289 *
    11) gleason>5.5 8 11.16 1.3140 *
 3) grade>2.5 85 125.10 1.6230
  6) age>56.5 75 104.00 1.4320
    12) gleason<7.5 50 66.49 1.1490
      24) g2<13.475 25 29.10 0.8817 *
      25) g2>13.475 25 36.05 1.4080
```

```

50) g2>17.915 14 18.72 0.8795 *
51) g2<17.915 11 13.70 2.1830 *
13) gleason>7.5 25 34.13 2.0280
26) g2>15.29 10 11.81 1.2140 *
27) g2<15.29 15 19.36 2.7020 *
7) age<56.5 10 15.52 3.1980 *

```

```

> plot(fit, uniform=T, branch=.4, compress=T)
> text(fit, use.n=T)

```

Note that the primary split on grade is the same as when status was used as a dichotomous endpoint, but that the splits thereafter differ.

```

> summary(fit, cp=.02)
Call:
rpart(formula = Surv(pptime, pstat) ~ age + eet + g2 + grade + gleason +
      ploidy, data = stagec)
n= 146

```

	CP	nsplit	rel error	xerror	xstd
1	0.12913	0	1.0000	1.0060	0.07389
2	0.04169	1	0.8709	0.8839	0.07584
3	0.02880	2	0.8292	0.9271	0.08196
4	0.01720	3	0.8004	0.9348	0.08326
5	0.01518	4	0.7832	0.9647	0.08259
6	0.01271	5	0.7680	0.9648	0.08258
7	0.01000	8	0.7311	0.9632	0.08480

```

Node number 1: 146 observations, complexity param=0.1291
events=54, estimated rate=1, mean deviance=1.338
left son=2 (61 obs) right son=3 (85 obs)

```

Primary splits:

```

grade < 2.5 to the left, improve=25.270, (0 missing)
gleason < 5.5 to the left, improve=21.630, (3 missing)
ploidy splits as LRR, improve=14.020, (0 missing)
g2 < 13.2 to the left, improve=12.580, (7 missing)
age < 58.5 to the right, improve= 2.796, (0 missing)

```

Surrogate splits:

```

gleason < 5.5 to the left, agree=0.863, adj=0.672, (0 split)
ploidy splits as LRR, agree=0.644, adj=0.148, (0 split)
g2 < 9.945 to the left, agree=0.630, adj=0.115, (0 split)
age < 66.5 to the right, agree=0.589, adj=0.016, (0 split)

```

```

.
.

```

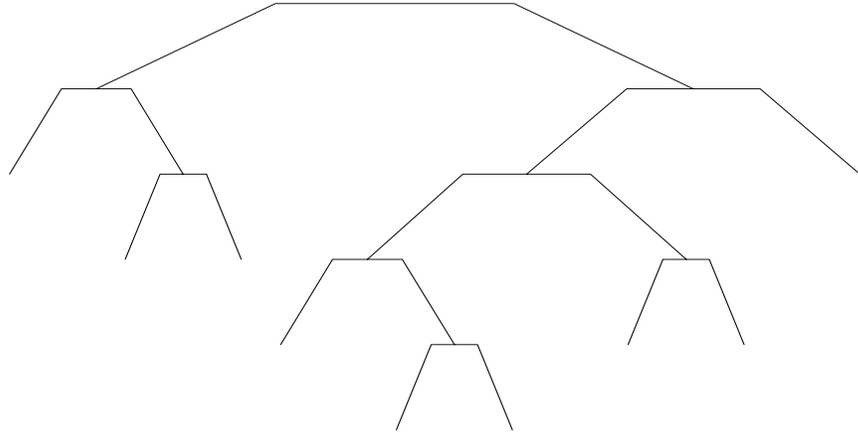


Figure 11: *The prostate cancer data as a survival tree*

.

Suppose that we wish to simplify this tree, so that only four terminal nodes remain. Looking at the table of complexity parameters, we see that `prune(fit, cp=.015)` would give the desired result. It is also possible to trim the figure interactively using `snip.rpart`. Point the mouse on a node and click with the left button to get some simple descriptives of the node. Double-clicking with the left button will ‘remove’ the sub-tree below, or one may click on another node. Multiple branches may be snipped off one by one; clicking with the right button will end interactive mode and return the pruned tree.

```
> plot(fit)
> text(fit,use.n=T)
> fit2 <- snip.rpart(fit)

node number: 6  n= 75
  response= 1.432013 ( 37 )
  Error (dev) = 103.9783

> plot(fit2)
> text(fit2,use.n=T)
```

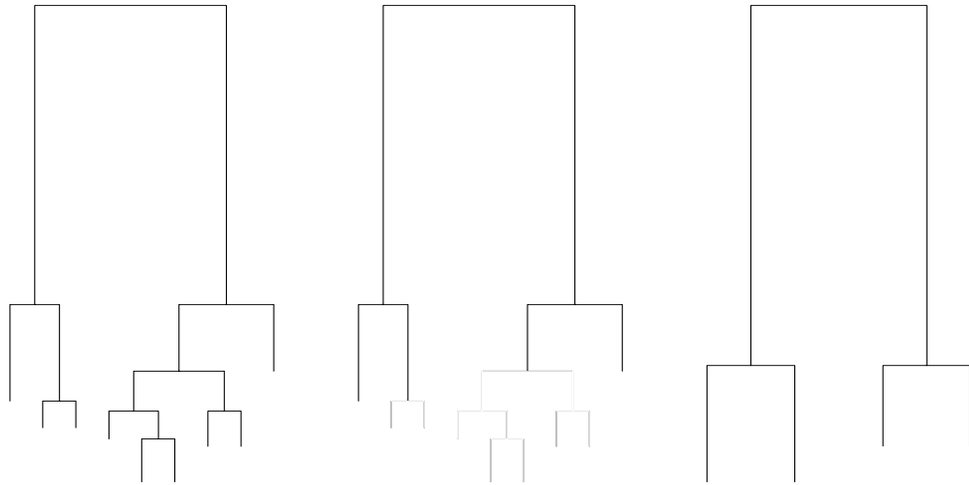


Figure 12: *An illustration of how `snip.rpart` works. The full tree is plotted in the first panel. After selecting node 6 with the mouse (double clicking on left button), the subtree disappears from the plot (shown in the second panel). Finally, the new subtree is redrawn to use all available space and it is labelled.*

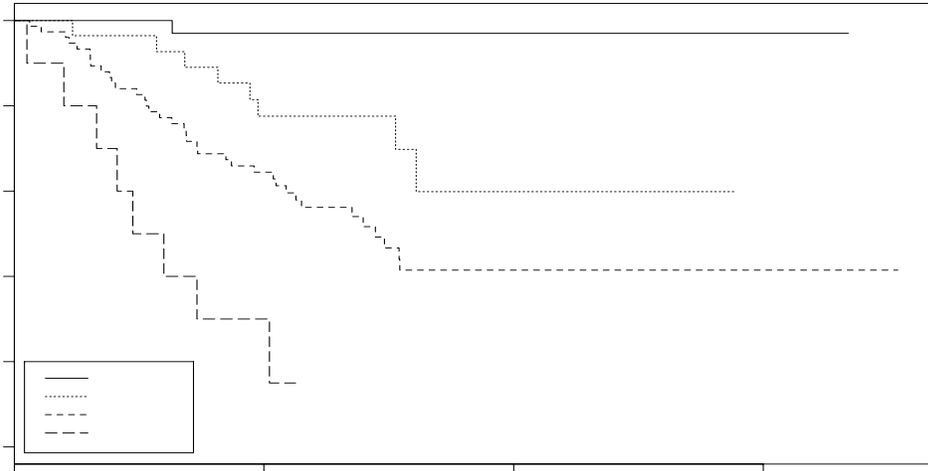


Figure 13: *Survival plot based on snipped rpart object. The probability of tumor progression is greatest in node 7, which has patients who are younger and have a higher initial tumor grade.*

For a final summary of the model, it can be helpful to plot the probability of survival based on the final bins in which the subjects landed. To create new variables based on the rpart groupings, use `where`. The nodes of `fit2` above are shown in the right hand panel of figure 12: node 4 has 1 event, 33 subjects, grade = 1-2 and $g2 < 11.36$; node 5 has 8 events, 28 subjects, grade = 1-2 and $g2 > 11.36$; node 6 has 37 events, 75 subjects, grade = 3-4, age > 56.5 ; node 7 has 8 events, 10 subjects, grade = 3-4, age < 56.5 . Patients who are younger and have a higher initial grade tend to have more rapid progression of disease.

```
> newgrp <- fit2$where
> plot(survfit(Surv(pgtime,pgstat) ~ newgrp, data=stagec),
      mark.time=F, lty=1:4)
> title(xlab='Time to Progression',ylab='Prob Progression')
> legend(.2,.2, legend=paste('node',c(4,5,6,7)), lty=1:4)
```

References

- [1] L. Breiman, J.H. Friedman, R.A. Olshen, , and C.J Stone. *Classification and Regression Trees*. Wadsworth, Belmont, Ca, 1983.

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- [3] M. LeBlanc and J Crowley. Relative risk trees for censored survival data. *Biometrics*, 48:411–425, 1992.
- [4] O. Nativ, Y. Raz, H.Z. Winkler, Y. Hosaka, E.T. Boyle, T.M. Therneau, G.M. Farrow, R.P. Meyers, H. Zincke, and M.M Lieber. Prognostic value of flow cytometric nuclear DNA analysis in stage C prostate carcinoma. *Surgical Forum*, pages 685–687, 1988.
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- [6] T.M. Therneau, Grambsch P.M., and T.R. Fleming. Martingale based residuals for survival models. *Biometrika*, 77:147–160, 1990.