Classification and Regression Tree Example

 $Packages\ used$

```
## ade4 mvpart rpart vegan
## 1.6-2 1.6-2 4.1-8 2.0-10
```

Data

Load the rpart package and take a look at data on cars from *Consumer Reports* (price, reliability, mileage, etc.):

```
library(rpart)
summary(cu.summary)
```

(n.b.: do **not** load the mvpart package in this section – it will load a different version of the rpart() function. If you load it by accident, use det ach("package:mvpart") to get rid of it)

Grow tree

If you know you want to use all the variables other than the response variable (Mileage in this case) as predictors, you can use the formula Mileage...

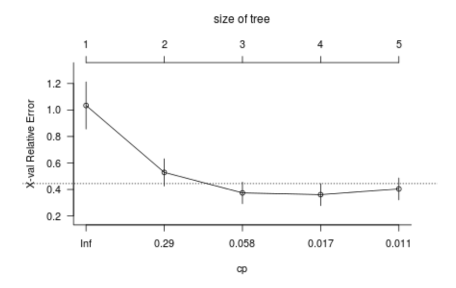
If your data are binary and so you want to build a classification tree rather than a regression tree, use method="class".

Display results (basic information plus a table of "complexity parameter" CP, number of splits, relative error, cross-validation error (xerror) and the standard deviation of the cross-validation error (xstd).

```
printcp(fit)
```

We can plot the cross-validation error as a function of the number of splits:

```
plotcp(fit)
```

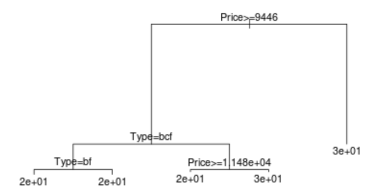


Or get detailed information about the tree:

summary(fit) # detailed summary of splits

We can also plot the tree itself (plot gives just the branches, text adds labels):

plot(fit)
text(fit)

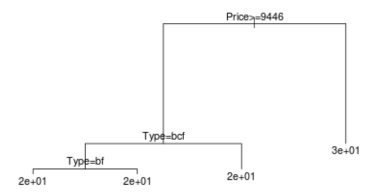


Now we can prune the tree back to its optimal size (based on cross-validation error):

```
best.tree <- which.min(fit$cptable[,"xerror"])
cpval <- fit$cptable[best.tree,"CP"]
pfit <- prune(fit, cp=cpval)</pre>
```

The resulting tree is simpler (only 4 splits; note that all the trees above 3 splits have about the same xerror):

```
plot(pfit)
text(pfit)
```



```
summary(pfit)
```

Pick your own tree size



summary(dfit)

(Code modified from Quick-R: Accessing the Power of R)

Multivariate Regression Tree Example

Data

Data information

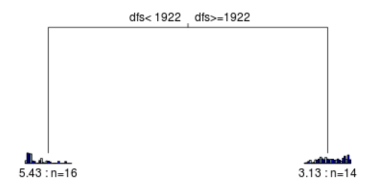
```
library(ade4)
data(doubs)
env <- doubs$env  ## site (row) * env variable (column)
spe <- doubs$fish  ## site (row) * species (column)</pre>
```

Transform response variables

The transformation consists of expressing each fish density as a proportion of the sum of all densities in the analytical unit and taking the square root of the resulting value (Legendre and Gallagher 2001). The square-root portion of the transformation decreases the importance of the most abundant species.

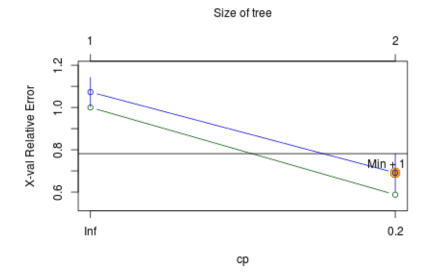
```
library(vegan)
spe.norm <- as.matrix(decostand(spe, "hellinger"))</pre>
```

$Multivariate\ Regression\ Tree$



Error: 0.588 CV Error: 0.692 SE: 0.0902

```
summary(spe.ch.mvpart)
printcp(spe.ch.mvpart)
plotcp(spe.ch.mvpart)
```



Or we can use "pick"

- xv = Selection of tree by cross-validation:
 - "1se" gives best tree within one SE of the overall best,
 - "min" the best tree
 - "pick" pick the tree size interactively,
 - "none" no cross-validation.
- xval = Number of cross-validations or vector defining cross-validation groups (here we use as many rows there are in the dataset (leave-one-out cross-validation) because it is a small dataset)
- xvmult = Number of multiple cross-validations.
- which = Which split labels and where to plot them: 1=centered, 2 = left, 3 = right and 4 = both.

(Modified R Code from Numerical Ecology with R, Borcard et al. 2012)