

Classification.R

Wow

Mon Sep 10 10:14:20 2018

```
#####  
#                               Classification                               #####  
#####  
library(mlbench)  
data(Zoo) #17 attributes, 101 animals, 7 classes = {Mammal, Bird, Reptile,  
Fish, Amphibian, Insect, Mollusc.et.al.}  
head(Zoo)  
  
##           hair feathers  eggs  milk airborne aquatic predator toothed  
## aardvark TRUE      FALSE FALSE TRUE      FALSE  FALSE      TRUE   TRUE  
## antelope TRUE      FALSE FALSE TRUE      FALSE  FALSE      FALSE  TRUE  
## bass     FALSE     FALSE TRUE  FALSE     FALSE   TRUE       TRUE   TRUE  
## bear     TRUE      FALSE FALSE TRUE      FALSE  FALSE      TRUE   TRUE  
## boar     TRUE      FALSE FALSE TRUE      FALSE  FALSE      TRUE   TRUE  
## buffalo  TRUE      FALSE FALSE TRUE      FALSE  FALSE      FALSE  TRUE  
##           backbone breathes venomous  fins legs  tail domestic catsize  
## aardvark TRUE      TRUE      FALSE FALSE  4 FALSE  FALSE  TRUE  
## antelope TRUE      TRUE      FALSE FALSE  4 TRUE   FALSE  TRUE  
## bass     TRUE      FALSE     FALSE TRUE   0 TRUE   FALSE FALSE  
## bear     TRUE      TRUE      FALSE FALSE  4 FALSE  FALSE  TRUE  
## boar     TRUE      TRUE      FALSE FALSE  4 TRUE   FALSE  TRUE  
## buffalo  TRUE      TRUE      FALSE FALSE  4 TRUE   FALSE  TRUE  
##           type  
## aardvark mammal  
## antelope mammal  
## bass     fish  
## bear     mammal  
## boar     mammal  
## buffalo  mammal  
  
summary(Zoo)  
  
##           hair           feathers           eggs           milk  
## Mode :logical Mode :logical Mode :logical Mode :logical  
## FALSE:58      FALSE:81      FALSE:42      FALSE:60  
## TRUE :43      TRUE :20      TRUE :59      TRUE :41  
##  
##  
##  
##           airborne           aquatic           predator           toothed  
## Mode :logical Mode :logical Mode :logical Mode :logical
```

```

## FALSE:77      FALSE:65      FALSE:45      FALSE:40
## TRUE :24      TRUE :36      TRUE :56      TRUE :61
##
##
##
## backbone      breathes      venomous      fins
## Mode :logical  Mode :logical  Mode :logical  Mode :logical
## FALSE:18      FALSE:21      FALSE:93      FALSE:84
## TRUE :83      TRUE :80      TRUE :8       TRUE :17
##
##
##
## legs          tail          domestic      catsize
## Min. :0.000    Mode :logical  Mode :logical  Mode :logical
## 1st Qu.:2.000    FALSE:26      FALSE:88      FALSE:57
## Median :4.000    TRUE :75      TRUE :13      TRUE :44
## Mean :2.842
## 3rd Qu.:4.000
## Max. :8.000
##
##          type
## mammal   :41
## bird     :20
## reptile  : 5
## fish     :13
## amphibian : 4
## insect   : 8
## mollusc.et.al:10

#####
# Recursive Partitioning and Regression Trees (RPART)
# similar to Classification and regression trees (CART)
#####
library(rpart)
tree1 <- rpart(type ~ ., data=Zoo) # predict "type" attribute using all the
rest attributes
tree1 #TRUE =1, FALSE = 0

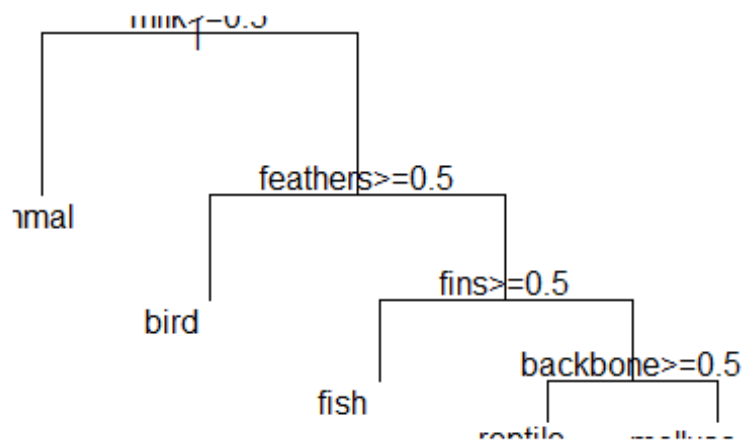
## n= 101
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
## 1) root 101 60 mammal (0.41 0.2 0.05 0.13 0.04 0.079 0.099)
##    2) milk>=0.5 41 0 mammal (1 0 0 0 0 0) *
##    3) milk< 0.5 60 40 bird (0 0.33 0.083 0.22 0.067 0.13 0.17)
##      6) feathers>=0.5 20 0 bird (0 1 0 0 0 0) *
##      7) feathers< 0.5 40 27 fish (0 0 0.12 0.32 0.1 0.2 0.25)

```

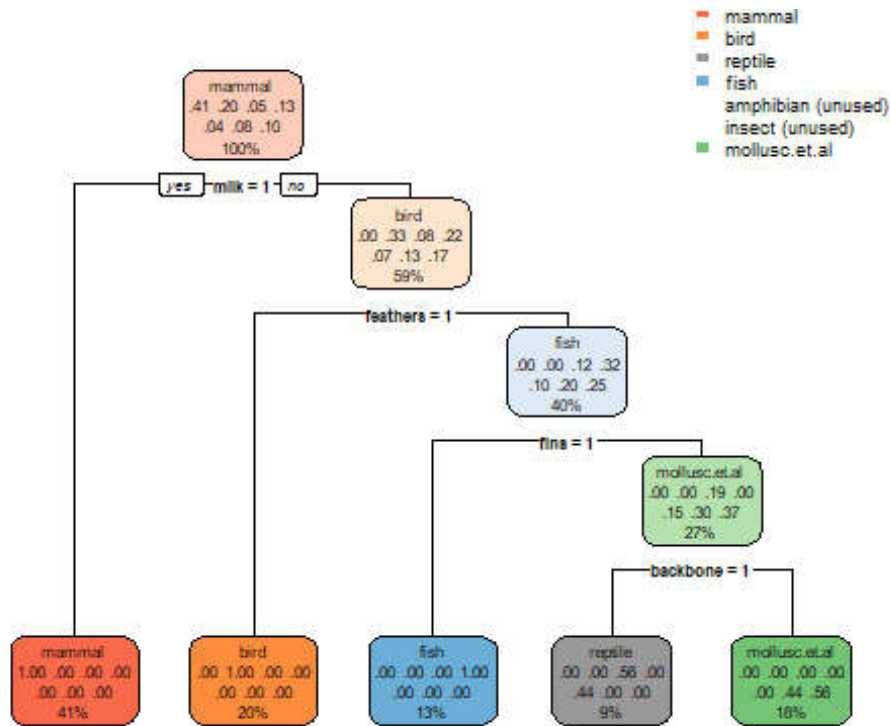
```
##      14) fins>=0.5 13  0 fish (0 0 0 1 0 0 0) *
##      15) fins< 0.5 27 17 mollusc.et.al (0 0 0.19 0 0.15 0.3 0.37)
##      30) backbone>=0.5 9  4 reptile (0 0 0.56 0 0.44 0 0) *
##      31) backbone< 0.5 18  8 mollusc.et.al (0 0 0 0 0 0.44 0.56) *
```

meaning of the first line: 101 = # total animals at that node | 60 = # misclassified animals | mammal = prediction result at that node | (0.41 0.2 0.05 0.13 0.04 0.079 0.099) = classes distribution (calculated from #total)

```
plot(tree1) #see ?plot.rpart
text(tree1)
```



```
# Better plotting
library(rpart.plot)
rpart.plot(tree1)
```



```

# Create a full tree (see: ?rpart.control)
tree2 <- rpart(type ~., data=Zoo, control=rpart.control(minsplit=2,cp=0.01))
# minsplit = the minimum number of observations that must exist in a node in
# order for a split to be attempted.
# cp = complexity parameter. Any split that does not decrease the overall
# lack of fit by a factor of cp is not attempted.
tree2

## n= 101
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
## 1) root 101 60 mammal (0.41 0.2 0.05 0.13 0.04 0.079 0.099)
##    2) milk>=0.5 41 0 mammal (1 0 0 0 0 0) *
##    3) milk< 0.5 60 40 bird (0 0.33 0.083 0.22 0.067 0.13 0.17)
##      6) feathers>=0.5 20 0 bird (0 1 0 0 0 0) *
##      7) feathers< 0.5 40 27 fish (0 0 0.12 0.32 0.1 0.2 0.25)
##        14) fins>=0.5 13 0 fish (0 0 0 1 0 0) *
##        15) fins< 0.5 27 17 mollusc.etal (0 0 0.19 0 0.15 0.3 0.37)
##          30) backbone>=0.5 9 4 reptile (0 0 0.56 0 0.44 0 0)
##            60) aquatic< 0.5 4 0 reptile (0 0 1 0 0 0) *
##            61) aquatic>=0.5 5 1 amphibian (0 0 0.2 0 0.8 0 0)
##              122) eggs< 0.5 1 0 reptile (0 0 1 0 0 0) *
##              123) eggs>=0.5 4 0 amphibian (0 0 0 0 1 0) *
##          31) backbone< 0.5 18 8 mollusc.etal (0 0 0 0 0 0.44 0.56)
##            62) airborne>=0.5 6 0 insect (0 0 0 0 0 1 0) *

```

```
##          63) airborne< 0.5 12 2 mollusc.et.al (0 0 0 0 0 0.17 0.83)
##          126) predator< 0.5 4 2 insect (0 0 0 0 0 0.5 0.5)
##          252) legs>=3 2 0 insect (0 0 0 0 0 1 0) *
##          253) legs< 3 2 0 mollusc.et.al (0 0 0 0 0 0 1) *
##          127) predator>=0.5 8 0 mollusc.et.al (0 0 0 0 0 0 1) *
```

```
rpart.plot(tree2)
```

```
## Warning: All boxes will be white (the box.palette argument will be
## ignored) because
## the number of classes predicted by the model 7 is greater than
## length(box.palette) 6.
## To silence this warning use box.palette=0 or trace=-1.
```

```
# Use the model to predict training records
```

```
# (See ?predict.rpart)
```

```
pred1 <- predict(tree1, Zoo) #return probability (as default)
```

```
pred1
```

	mammal	bird	reptile	fish	amphibian	insect	mollusc.et.al
aardvark	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
antelope	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
bass	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
bear	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
boar	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
buffalo	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
calf	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
carp	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
catfish	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
cavy	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
cheetah	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
chicken	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
chub	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
clam	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
crab	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
crayfish	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
crow	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
deer	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
dogfish	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
dolphin	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
dove	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
duck	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
elephant	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
flamingo	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
flea	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
frog.1	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000
frog.2	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000
fruitbat	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
giraffe	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
girl	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
gnat	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556

## goat	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## gorilla	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## gull	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## haddock	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
## hamster	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## hare	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## hawk	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## herring	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
## honeybee	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## housefly	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## kiwi	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## ladybird	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## lark	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## leopard	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## lion	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## lobster	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## lynx	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## mink	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## mole	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## mongoose	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## moth	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## newt	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000
## octopus	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## opossum	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## oryx	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## ostrich	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## parakeet	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## penguin	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## pheasant	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## pike	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
## piranha	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
## pitviper	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000
## platypus	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## polecat	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## pony	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## porpoise	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## puma	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## pussycat	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## raccoon	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## reindeer	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## rhea	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## scorpion	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## seahorse	0	0	0.0000000	1	0.0000000	0.0000000	0.0000000
## seal	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## sealion	1	0	0.0000000	0	0.0000000	0.0000000	0.0000000
## seasnake	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000
## seawasp	0	0	0.0000000	0	0.0000000	0.4444444	0.5555556
## skimmer	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## skua	0	1	0.0000000	0	0.0000000	0.0000000	0.0000000
## slowworm	0	0	0.5555556	0	0.4444444	0.0000000	0.0000000

```
## slug      0      0 0.0000000      0 0.0000000 0.4444444      0.5555556
## sole      0      0 0.0000000      1 0.0000000 0.0000000      0.0000000
## sparrow   0      1 0.0000000      0 0.0000000 0.0000000      0.0000000
## squirrel  1      0 0.0000000      0 0.0000000 0.0000000      0.0000000
## starfish  0      0 0.0000000      0 0.0000000 0.4444444      0.5555556
## stingray  0      0 0.0000000      1 0.0000000 0.0000000      0.0000000
## swan      0      1 0.0000000      0 0.0000000 0.0000000      0.0000000
## termite   0      0 0.0000000      0 0.0000000 0.4444444      0.5555556
## toad      0      0 0.5555556      0 0.4444444 0.0000000      0.0000000
## tortoise  0      0 0.5555556      0 0.4444444 0.0000000      0.0000000
## tuatara   0      0 0.5555556      0 0.4444444 0.0000000      0.0000000
## tuna      0      0 0.0000000      1 0.0000000 0.0000000      0.0000000
## vampire   1      0 0.0000000      0 0.0000000 0.0000000      0.0000000
## vole      1      0 0.0000000      0 0.0000000 0.0000000      0.0000000
## vulture   0      1 0.0000000      0 0.0000000 0.0000000      0.0000000
## wallaby   1      0 0.0000000      0 0.0000000 0.0000000      0.0000000
## wasp      0      0 0.0000000      0 0.0000000 0.4444444      0.5555556
## wolf      1      0 0.0000000      0 0.0000000 0.0000000      0.0000000
## worm      0      0 0.0000000      0 0.0000000 0.4444444      0.5555556
## wren      0      1 0.0000000      0 0.0000000 0.0000000      0.0000000
```

```
pred2 <- predict(tree1, Zoo, type="class") #return a vector of classes
pred2
```

```
##      aardvark      antelope      bass      bear      boar
##      mammal      mammal      fish      mammal      mammal
##      buffalo      calf      carp      catfish      cavy
##      mammal      mammal      fish      fish      mammal
##      cheetah      chicken      chub      clam      crab
##      mammal      bird      fish mollusc.et.al mollusc.et.al
##      crayfish      crow      deer      dogfish      dolphin
## mollusc.et.al      bird      mammal      fish      mammal
##      dove      duck      elephant      flamingo      flea
##      bird      bird      mammal      bird mollusc.et.al
##      frog.1      frog.2      fruitbat      giraffe      girl
##      reptile      reptile      mammal      mammal      mammal
##      gnat      goat      gorilla      gull      haddock
## mollusc.et.al      mammal      mammal      bird      fish
##      hamster      hare      hawk      herring      honeybee
##      mammal      mammal      bird      fish mollusc.et.al
##      housefly      kiwi      ladybird      lark      leopard
## mollusc.et.al      bird mollusc.et.al      bird      mammal
##      lion      lobster      lynx      mink      mole
##      mammal mollusc.et.al      mammal      mammal      mammal
##      mongoose      moth      newt      octopus      opossum
##      mammal mollusc.et.al      reptile mollusc.et.al      mammal
##      oryx      ostrich      parakeet      penguin      pheasant
##      mammal      bird      bird      bird      bird
##      pike      piranha      pitviper      platypus      polecat
##      fish      fish      reptile      mammal      mammal
```

```
##      pony      porpoise      puma      pussycat      raccoon
##      mammal      mammal      mammal      mammal      mammal
##      reindeer      rhea      scorpion      seahorse      seal
##      mammal      bird mollusc.et.al      fish      mammal
##      sealion      seasnake      seawasp      skimmer      skua
##      mammal      reptile mollusc.et.al      bird      bird
##      slowworm      slug      sole      sparrow      squirrel
##      reptile mollusc.et.al      fish      bird      mammal
##      starfish      stingray      swan      termite      toad
## mollusc.et.al      fish      bird mollusc.et.al      reptile
##      tortoise      tuatara      tuna      vampire      vole
##      reptile      reptile      fish      mammal      mammal
##      vulture      wallaby      wasp      wolf      worm
##      bird      mammal mollusc.et.al      mammal mollusc.et.al
##      wren
##      bird
## Levels: mammal bird reptile fish amphibian insect mollusc.et.al
```

```
pred3 <- predict(tree1, Zoo, type="vector") #return a vector of numbers
representing classes
pred3
```

```
## aardvark antelope      bass      bear      boar      buffalo      calf      carp
##      1      1      4      1      1      1      1      4
## catfish      cavy      cheetah      chicken      chub      clam      crab      crayfish
##      4      1      1      2      4      7      7      7
##      crow      deer      dogfish      dolphin      dove      duck      elephant      flamingo
##      2      1      4      1      2      2      1      2
##      flea      frog.1      frog.2      fruitbat      giraffe      girl      gnat      goat
##      7      3      3      1      1      1      7      1
## gorilla      gull      haddock      hamster      hare      hawk      herring      honeybee
##      1      2      4      1      1      2      4      7
## housefly      kiwi      ladybird      lark      leopard      lion      lobster      lynx
##      7      2      7      2      1      1      7      1
##      mink      mole      mongoose      moth      newt      octopus      opossum      oryx
##      1      1      1      7      3      7      1      1
## ostrich      parakeet      penguin      pheasant      pike      piranha      pitviper      platypus
##      2      2      2      2      4      4      3      1
## polecat      pony      porpoise      puma      pussycat      raccoon      reindeer      rhea
##      1      1      1      1      1      1      1      2
## scorpion      seahorse      seal      sealion      seasnake      seawasp      skimmer      skua
##      7      4      1      1      3      7      2      2
## slowworm      slug      sole      sparrow      squirrel      starfish      stingray      swan
##      3      7      4      2      1      7      4      2
## termite      toad      tortoise      tuatara      tuna      vampire      vole      vulture
##      7      3      3      3      4      1      1      2
## wallaby      wasp      wolf      worm      wren
##      1      7      1      7      2
```



```

# Create confusion matrix
confusion_table <- table(Zoo$type, pred2)
confusion_table

##           pred2
##           mammal bird reptile fish amphibian insect mollusc.et.al
## mammal           41    0      0    0           0      0           0
## bird              0   20      0    0           0      0           0
## reptile           0    0      5    0           0      0           0
## fish              0    0      0   13           0      0           0
## amphibian         0    0      4    0           0      0           0
## insect            0    0      0    0           0      0           8
## mollusc.et.al     0    0      0    0           0      0          10

# Evaluate number of animals that were predicted correctly using tree1
correct <- sum(diag(confusion_table))
correct

## [1] 89

# Evaluate the training errors which is the number of misclassification
errors committed on training records.
error <- sum(confusion_table)-correct
error

## [1] 12

# Evaluate the accuracy
accuracy <- correct/(correct+error)
accuracy

## [1] 0.8811881

#####
# Create function for accuracy
accuracy <- function(truth, prediction) { #2 inputs: a truth vector and a
prediction vector
  tbl <- table(truth, prediction) #create confusion matrix
  sum(diag(tbl))/sum(tbl) #calculate accuracy
}

# Apply "accuracy" function
accuracy(Zoo$type, pred2)

## [1] 0.8811881

# Training error of the full tree
accuracy(Zoo$type, predict(tree2, Zoo, type="class"))

## [1] 1

```

```
#####
# Use rpart on a subset of attributes
tree3 <- rpart(type ~ hair+feathers+eggs+milk, data=Zoo) # predict "type"
attribute using only hair,feathers,eggs,mik attributes
tree3

## n= 101
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
## 1) root 101 60 mammal (0.41 0.2 0.05 0.13 0.04 0.079 0.099)
##   2) milk>=0.5 41 0 mammal (1 0 0 0 0 0) *
##   3) milk< 0.5 60 40 bird (0 0.33 0.083 0.22 0.067 0.13 0.17)
##     6) feathers>=0.5 20 0 bird (0 1 0 0 0 0) *
##     7) feathers< 0.5 40 27 fish (0 0 0.12 0.32 0.1 0.2 0.25) *

#####
# Use rpart on a subset of records
tree4 <- rpart(type ~ ., data=Zoo, subset = c(1:50)) # predict "type"
attribute using only first 50 records
tree4

## n= 50
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
## 1) root 50 27 mammal (0.46 0.18 0 0.14 0.04 0.1 0.08)
##   2) eggs< 0.5 23 0 mammal (1 0 0 0 0 0) *
##   3) eggs>=0.5 27 18 bird (0 0.33 0 0.26 0.074 0.19 0.15)
##     6) feathers>=0.5 9 0 bird (0 1 0 0 0 0) *
##     7) feathers< 0.5 18 11 fish (0 0 0 0.39 0.11 0.28 0.22) *

#####
# Methods to evaluate the performance of a classifier
#####
# 1. Hold-out: Create a decision tree using a training set and test a tree on
a test set
n_train <- as.integer(nrow(Zoo)*0.67) #2/3 of the records will be used as a
training set and 1/3 of the records will be used as a test set
train_id <- sample(1:nrow(Zoo), n_train)
train <- Zoo[train_id,]
test <- Zoo[-train_id, -17]
test_type <- Zoo[-train_id, 17]

# Create a decision tree
tree <- rpart(type ~., data=train,control=rpart.control(minsplit=2,cp=0.01))
tree
```

```

## n= 67
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
##  1) root 67 40 mammal (0.4 0.19 0.03 0.13 0.045 0.075 0.12)
##    2) milk>=0.5 27 0 mammal (1 0 0 0 0 0 0) *
##    3) milk< 0.5 40 27 bird (0 0.32 0.05 0.22 0.075 0.12 0.2)
##      6) feathers>=0.5 13 0 bird (0 1 0 0 0 0 0) *
##      7) feathers< 0.5 27 18 fish (0 0 0.074 0.33 0.11 0.19 0.3)
##    14) fins>=0.5 9 0 fish (0 0 0 1 0 0 0) *
##    15) fins< 0.5 18 10 mollusc.et.al (0 0 0.11 0 0.17 0.28 0.44)
##      30) airborne< 0.5 14 6 mollusc.et.al (0 0 0.14 0 0.21 0.071
0.57)
##      60) backbone>=0.5 5 2 amphibian (0 0 0.4 0 0.6 0 0)
##    120) aquatic< 0.5 2 0 reptile (0 0 1 0 0 0 0) *
##    121) aquatic>=0.5 3 0 amphibian (0 0 0 0 1 0 0) *
##    61) backbone< 0.5 9 1 mollusc.et.al (0 0 0 0 0 0.11 0.89)
##    122) predator< 0.5 3 1 mollusc.et.al (0 0 0 0 0 0.33 0.67)
##    244) legs>=3 1 0 insect (0 0 0 0 0 1 0) *
##    245) legs< 3 2 0 mollusc.et.al (0 0 0 0 0 0 1) *
##    123) predator>=0.5 6 0 mollusc.et.al (0 0 0 0 0 0 1) *
##    31) airborne>=0.5 4 0 insect (0 0 0 0 0 1 0) *

# Training error
accuracy(train$type, predict(tree, train, type="class"))

## [1] 1

# Generalization error
accuracy(test_type, predict(tree, test, type="class"))

## [1] 0.9705882

#####
# 2. Random Subsampling: It repeats holdout method several times.
k <- 10 #number of hold-out (could be changed)

# Do each hold-out
accs <- vector(mode="numeric") #create an empty numeric vector
for(i in 1:k){
  n_train <- as.integer(nrow(Zoo)*0.67)
  train_id <- sample(1:nrow(Zoo), n_train)
  train <- Zoo[train_id,]
  test <- Zoo[-train_id, -17]
  test_type <- Zoo[-train_id, 17]
  tree <- rpart(type ~.,
data=train,control=rpart.control(minsplit=2,cp=0.01))
  accs[i] <- accuracy(test_type, predict(tree, test, type="class"))
}
accs

```

```

## [1] 0.9411765 0.9705882 0.9411765 0.8529412 0.9411765 0.9411765 0.9411765
## [8] 0.9705882 1.0000000 0.9411765

# Report the average accuracy
mean(accs)

## [1] 0.9441176

#####
# 3. Crossvalidation
# 3.1 k-fold cross-validation
k <- 10 #number of folds (could be changed)
index <- 1:nrow(Zoo)
index <- sample(index) # shuffle index
fold <- rep(1:k, each=nrow(Zoo)/k)[1:nrow(Zoo)] #make a repeat vector to be
the same size as index vector
folds <- split(index, fold) # split(x,f) = split divides the data in the
vector x into the groups defined by f.
folds

## $`1`
## [1] 33 84 52 58 45 78 54 87 60 24
##
## $`2`
## [1] 10 2 67 80 95 51 13 71 69 36
##
## $`3`
## [1] 26 21 59 85 37 83 7 34 77 100
##
## $`4`
## [1] 93 30 62 81 15 65 43 44 91 28
##
## $`5`
## [1] 73 98 70 5 27 76 64 99 79 90
##
## $`6`
## [1] 72 68 23 56 50 38 48 3 40 6
##
## $`7`
## [1] 16 46 101 8 86 14 39 88 18 1
##
## $`8`
## [1] 66 11 94 82 47 53 22 92 12 32
##
## $`9`
## [1] 4 9 25 57 89 49 42 75 63 35
##
## $`10`
## [1] 74 29 17 31 96 97 55 41 20 19

```

```

# Do each fold
accs <- vector(mode="numeric") #create an empty numeric vector
for(i in 1:length(folds)) {
  tree <- rpart(type ~., data=Zoo[-folds[[i]],],
control=rpart.control(minsplit=2,cp=0.01))
  accs[i] <- accuracy(Zoo[folds[[i]],]$type, predict(tree, Zoo[folds[[i]],],
type="class"))
}
accs

## [1] 1.0 1.0 0.9 0.9 0.8 1.0 1.0 0.9 0.8 1.0

# Report the average accuracy
mean(accs)

## [1] 0.93

#####
# 3.2 Leave-one-out
accs <- vector(mode="numeric") #create an empty numeric vector
for(i in 1:nrow(Zoo)) {
  tree <- rpart(type ~., data=Zoo[-i,],
control=rpart.control(minsplit=2,cp=0.01))
  accs[i] <- accuracy(Zoo[i,]$type, predict(tree, Zoo[i,], type="class"))
}
accs

## [1] 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 0 1 1 1 1 1 1 1 1
1
## [36] 1 1 1 1 1 1 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
1
## [71] 1 1 0 1 1 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 0 1 0 1 1 1 1 1 1 1 1

# Report the average accuracy
mean(accs)

## [1] 0.9306931

#####
# 4. Bootstrap
k <- 10 #number of rounds (could be changed)
n_train <- as.integer(nrow(Zoo)*0.67) #2/3 of the records will be used as a
training set and 1/3 of the records will be used as a test set
train_id <- sample(1:nrow(Zoo), n_train)
test <- Zoo[-train_id, -17]
test_type <- Zoo[-train_id, 17]

# Do each round
accs <- vector(mode="numeric") #create an empty numeric vector
for(i in 1:k) {
  btrain_id <- sample(train_id, n_train, replace = TRUE) #Bootstrap sampling
for training set

```

```

btrain <- Zoo[btrain_id,]
tree <- rpart(type ~., data=btrain,
control=rpart.control(minsplit=2,cp=0.01))
accs[i] <- accuracy(test_type, predict(tree, test, type="class"))
}
accs

## [1] 0.9411765 0.9411765 0.9117647 0.9411765 0.7941176 0.8823529 0.8529412
## [8] 0.8529412 0.7941176 0.9117647

# Report the average accuracy
mean(accs)

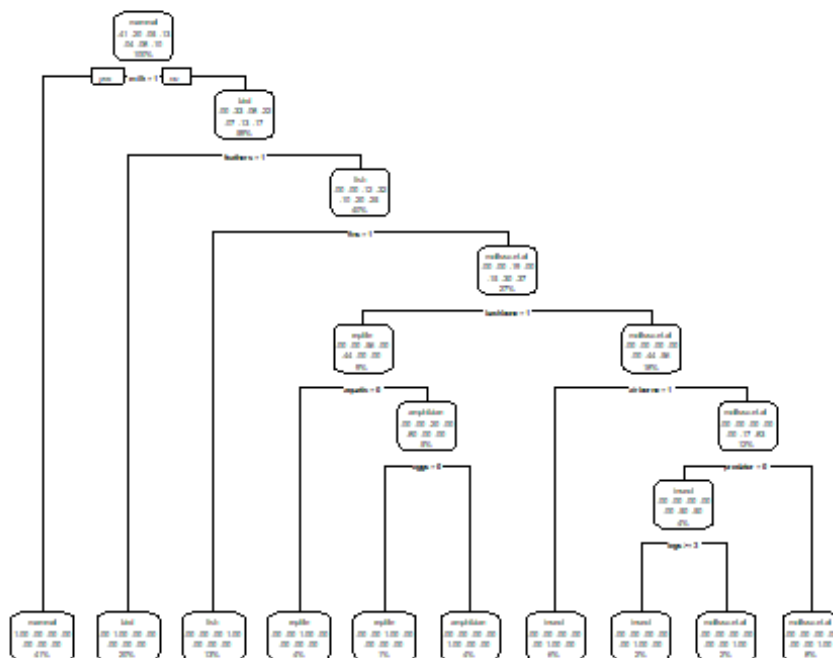
## [1] 0.8823529

#####
# Use caret package for easier model building and evaluation
#####
# See http://caret.r-forge.r-project.org/
# https://cran.r-project.org/web/packages/caret/caret.pdf

# Use multi-core to make R faster
library(foreach)
library(iterators)
library(parallel)
library(doParallel)
registerDoParallel()

# Evaluation with caret (train normally tries to tune cp for rpart). By
# setting tuneLength 0 and tuneGrid fixed the value to 0.01, rpart is run with
# no tuning
# See ?trainControl for method options
library(ggplot2)

```



```
library(lattice)
library(caret)
library(e1071)
# 10 folds crossvalidation
fit <- train(Zoo[,-17], Zoo[,17],
             method = "rpart",
             control=rpart.control(minsplit=2),
             tuneGrid=data.frame(cp=0.01),
             trControl = trainControl(method = "cv", number = 10), #10 is
number of folds
             tuneLength=0)
# if you get the error message -> reinstall package caret
fit

## CART
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.etal'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 91, 90, 91, 91, 92, 90, ...
## Resampling results:
##
## Accuracy Kappa
```

```

##    0.9407071  0.9220195
##
## Tuning parameter 'cp' was held constant at a value of 0.01

# plot tree
rpart.plot(fit$finalModel)

## Warning: All boxes will be white (the box.palette argument will be
## ignored) because
## the number of classes predicted by the model 7 is greater than
## length(box.palette) 6.
## To silence this warning use box.palette=0 or trace=-1.

# Leave-one-out
fit <- train(Zoo[, -17], Zoo[, 17], method = "rpart",
             control=rpart.control(minsplit=2),
             tuneGrid=data.frame(cp=0.01),
             trControl = trainControl(method = "LOOCV"),
             tuneLength=0)

fit

## CART
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
## 'mollusc.et.al'
##
## No pre-processing
## Resampling: Leave-One-Out Cross-Validation
## Summary of sample sizes: 100, 100, 100, 100, 100, 100, ...
## Resampling results:
##
## Accuracy   Kappa
## 0.9306931  0.9086445
##
## Tuning parameter 'cp' was held constant at a value of 0.01

# Bootstrap
fit <- train(Zoo[, -17], Zoo[, 17], method = "rpart",
             control=rpart.control(minsplit=2),
             tuneGrid=data.frame(cp=0.01),
             trControl = trainControl(method = "boot", number = 10), #10 is
number of resampling iterations
             tuneLength=0)

fit

## CART
##
## 101 samples
## 16 predictor

```



```
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Bootstrapped (10 reps)
## Summary of sample sizes: 101, 101, 101, 101, 101, 101, ...
## Resampling results:
##
## Accuracy Kappa
## 0.9264073 0.9020965
##
## Tuning parameter 'cp' was held constant at a value of 0.01

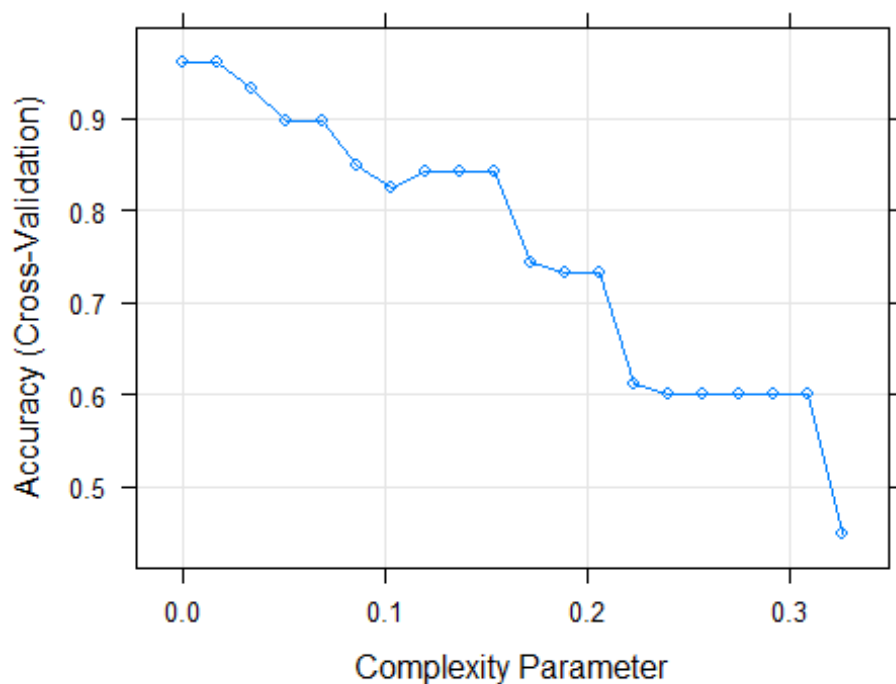
#####
# Finding the best model
#####
# Create training and test sets using createDataPartition function in caret
package
inTrain <- createDataPartition(y=Zoo$type, p = 0.75, list=FALSE)
training <- Zoo[inTrain,]
testing <- Zoo[-inTrain,]

# Find the best model
fit <- train(training[, -17], training[, 17], method = "rpart",
             control=rpart.control(minsplit=2),
             trControl = trainControl(method = "cv", number = 10),
             tuneLength=20) #vary cp (complexity parameter) for 20 values and
pick the one that give the highest accuracy
fit

## CART
##
## 77 samples
## 16 predictors
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 69, 69, 69, 71, 71, 68, ...
## Resampling results across tuning parameters:
##
## cp Accuracy Kappa
## 0.00000000 0.9621032 0.9483716
## 0.01716247 0.9621032 0.9486347
## 0.03432494 0.9329365 0.9079680
## 0.05148741 0.8968254 0.8622192
## 0.06864989 0.8968254 0.8622192
## 0.08581236 0.8496032 0.8019751
## 0.10297483 0.8246032 0.7711475
```

```
## 0.12013730 0.8412698 0.7951475
## 0.13729977 0.8412698 0.7951475
## 0.15446224 0.8412698 0.7951475
## 0.17162471 0.7436508 0.6597031
## 0.18878719 0.7325397 0.6456406
## 0.20594966 0.7325397 0.6418891
## 0.22311213 0.6113095 0.4623883
## 0.24027460 0.6001984 0.4445758
## 0.25743707 0.6001984 0.4445758
## 0.27459954 0.6001984 0.4445758
## 0.29176201 0.6001984 0.4445758
## 0.30892449 0.6001984 0.4445758
## 0.32608696 0.4474206 0.1136667
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was cp = 0.01716247.
```

```
plot(fit)
```



```
# Use the best model
pred <- predict(fit, testing)
pred
```

## [1]	mammal	mammal	fish	mollusc.et.al	bird
## [6]	mammal	bird	fish	insect	mollusc.et.al
## [11]	mammal	mammal	amphibian	bird	fish
## [16]	mammal	mammal	mammal	mammal	bird

```
## [21] amphibian      bird          mollusc.et.al mammal
## Levels: mammal bird reptile fish amphibian insect mollusc.et.al

# Confusion matrix (incl. confidence interval)
confusionMatrix(data = pred, testing$type)

## Confusion Matrix and Statistics
##
##              Reference
## Prediction      mammal bird reptile fish amphibian insect mollusc.et.al
## mammal           10    0      0    0          0      0          0
## bird              0    5      0    0          0      0          0
## reptile           0    0      0    0          0      0          0
## fish              0    0      0    3          0      0          0
## amphibian         0    0      1    0          1      0          0
## insect            0    0      0    0          0      1          0
## mollusc.et.al     0    0      0    0          0      1          2
##
## Overall Statistics
##
##              Accuracy : 0.9167
##              95% CI : (0.73, 0.9897)
##      No Information Rate : 0.4167
##      P-Value [Acc > NIR] : 4.315e-07
##
##              Kappa : 0.8889
##  Mcnemar's Test P-Value : NA
##
## Statistics by Class:
##
##              Class: mammal Class: bird Class: reptile Class: fish
## Sensitivity           1.0000          1.0000          0.00000          1.000
## Specificity           1.0000          1.0000          1.00000          1.000
## Pos Pred Value         1.0000          1.0000              NaN          1.000
## Neg Pred Value         1.0000          1.0000          0.95833          1.000
## Prevalence             0.4167          0.2083          0.04167          0.125
## Detection Rate         0.4167          0.2083          0.00000          0.125
## Detection Prevalence   0.4167          0.2083          0.00000          0.125
## Balanced Accuracy       1.0000          1.0000          0.50000          1.000
##              Class: amphibian Class: insect Class: mollusc.et.al
## Sensitivity           1.00000          0.50000          1.00000
## Specificity           0.95652          1.00000          0.95455
## Pos Pred Value         0.50000          1.00000          0.66667
## Neg Pred Value         1.00000          0.95652          1.00000
## Prevalence             0.04167          0.08333          0.08333
## Detection Rate         0.04167          0.04167          0.08333
## Detection Prevalence   0.08333          0.04167          0.12500
## Balanced Accuracy       0.97826          0.75000          0.97727
```

```
#####
# Other classification methods in caret package
#####
# See http://topepo.github.io/caret/train-models-by-tag.html

# Create fixed sampling scheme (10-folds)
train <- createFolds(Zoo$type, k=10)

test <- Zoo[1:50,-17]
test_type <- Zoo[1:50,17]

#####
# 1. Recursive Partitioning and Regression Trees (RPART)
# similar to Classification and regression trees (CART)
# Tuning parameters: cp (Complexity Parameter)

library(rpart)
rpartFit <- train(Zoo[, -17], Zoo[, 17], "rpart",
                  tuneLength = 10,
                  trControl = trainControl(method = "cv", indexOut = train))
rpartFit

## CART
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
## 'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 90, 90, 91, 92, 92, 90, ...
## Resampling results across tuning parameters:
##
##   cp          Accuracy    Kappa
##   0.00000000  0.8861616  0.8526187
##   0.03703704  0.8861616  0.8526187
##   0.07407407  0.8770707  0.8411604
##   0.11111111  0.8370707  0.7862417
##   0.14814815  0.8370707  0.7862417
##   0.18518519  0.7370960  0.6474605
##   0.22222222  0.6323485  0.4933646
##   0.25925926  0.6098485  0.4563303
##   0.29629630  0.6098485  0.4563303
##   0.33333333  0.4971212  0.2091997
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was cp = 0.03703704.

rpartFit$finalModel
```

```

## n= 101
##
## node), split, n, loss, yval, (yprob)
##      * denotes terminal node
##
## 1) root 101 60 mammal (0.41 0.2 0.05 0.13 0.04 0.079 0.099)
##    2) milk>=0.5 41 0 mammal (1 0 0 0 0 0 0) *
##    3) milk< 0.5 60 40 bird (0 0.33 0.083 0.22 0.067 0.13 0.17)
##      6) feathers>=0.5 20 0 bird (0 1 0 0 0 0 0) *
##      7) feathers< 0.5 40 27 fish (0 0 0.12 0.32 0.1 0.2 0.25)
##      14) fins>=0.5 13 0 fish (0 0 0 1 0 0 0) *
##      15) fins< 0.5 27 17 mollusc.et.al (0 0 0.19 0 0.15 0.3 0.37)
##      30) backbone>=0.5 9 4 reptile (0 0 0.56 0 0.44 0 0) *
##      31) backbone< 0.5 18 8 mollusc.et.al (0 0 0 0 0 0.44 0.56) *

accs <- accuracy(test_type, predict(rpartFit, test))
accs

## [1] 0.86

#####
# 2. Conditional Inference Tree (Decision Tree) (Ctree)
# Tuning parameters: mincriterion (1 - P-Value Threshold)

library(party)

## Loading required package: grid
## Loading required package: mvtnorm
## Loading required package: modeltools
## Loading required package: stats4
## Loading required package: strucchange
## Loading required package: zoo

##
## Attaching package: 'zoo'

## The following objects are masked from 'package:base':
##
##      as.Date, as.Date.numeric

## Loading required package: sandwich

ctreeFit <- train(Zoo[,-17], Zoo[,17], "ctree",
                  tuneLength = 10,
                  trControl = trainControl(method = "cv", indexOut = train))
ctreeFit

```

```

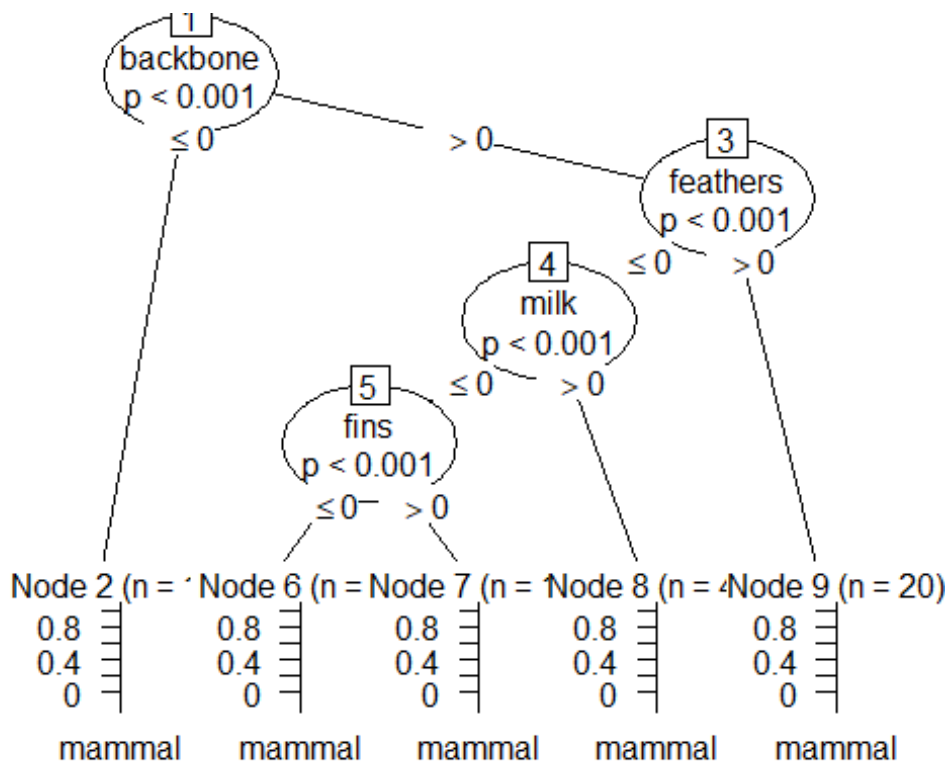
## Conditional Inference Tree
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 91, 90, 90, 92, 90, 91, ...
## Resampling results across tuning parameters:
##
## mincriterion Accuracy Kappa
## 0.0100000 0.8761616 0.8394608
## 0.1188889 0.8761616 0.8394608
## 0.2277778 0.8761616 0.8394608
## 0.3366667 0.8761616 0.8394608
## 0.4455556 0.8761616 0.8394608
## 0.5544444 0.8761616 0.8394608
## 0.6633333 0.8761616 0.8394608
## 0.7722222 0.8761616 0.8394608
## 0.8811111 0.8761616 0.8394608
## 0.9900000 0.8761616 0.8394608
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mincriterion = 0.99.

ctreeFit$finalModel

##
## Conditional inference tree with 5 terminal nodes
##
## Response: .outcome
## Inputs: hair, feathers, eggs, milk, airborne, aquatic, predator, toothed,
backbone, breathes, venomous, fins, legs, tail, domestic, catsize
## Number of observations: 101
##
## 1) backbone <= 0; criterion = 1, statistic = 100
## 2)* weights = 18
## 1) backbone > 0
## 3) feathers <= 0; criterion = 1, statistic = 82
## 4) milk <= 0; criterion = 1, statistic = 62
## 5) fins <= 0; criterion = 1, statistic = 21
## 6)* weights = 9
## 5) fins > 0
## 7)* weights = 13
## 4) milk > 0
## 8)* weights = 41
## 3) feathers > 0
## 9)* weights = 20

```

```
plot(ctreeFit$finalModel)
```



```

accs <- accuracy(test_type, predict(ctreeFit, test))
accs

## [1] 0.86

#####
# 3. Support Vector Machines with Linear Kernel
# Tuning parameters: C (Cost)

library(e1071)
svmFit <- train(Zoo[, -17], Zoo[, 17], "svmLinear2",
               tuneLength = 10,
               trControl = trainControl(method = "cv", indexOut = train))
svmFit

## Support Vector Machines with Linear Kernel
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
## 'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 91, 93, 91, 92, 91, 90, ...

```

```

## Resampling results across tuning parameters:
##
##   cost      Accuracy  Kappa
##   0.25      1          1
##   0.50      1          1
##   1.00      1          1
##   2.00      1          1
##   4.00      1          1
##   8.00      1          1
##   16.00     1          1
##   32.00     1          1
##   64.00     1          1
##   128.00    1          1
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was cost = 0.25.

svmFit$finalModel

##
## Call:
## svm.default(x = as.matrix(x), y = y, kernel = "linear", cost = param$cost,
##   probability = classProbs)
##
##
## Parameters:
##   SVM-Type:  C-classification
##   SVM-Kernel: linear
##   cost:      0.25
##   gamma:     0.0625
##
## Number of Support Vectors:  48

accs <- accuracy(test_type, predict(svmFit, test))
accs

## [1] 1

#####
# 4. k-Nearest Neighbors
# Tuning parameters: k (neighbors)

knnFit <- train(Zoo[, -17], Zoo[, 17], "knn",
  tuneLength = 10,
  trControl = trainControl(method = "cv", indexOut = train))
knnFit

## k-Nearest Neighbors
##
## 101 samples
## 16 predictor

```



```

## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 90, 90, 91, 92, 90, 91, ...
## Resampling results across tuning parameters:
##
## k Accuracy Kappa
## 5 0.9020455 0.8676959
## 7 0.8456818 0.7942121
## 9 0.8065909 0.7393416
## 11 0.7963889 0.7236157
## 13 0.7863889 0.7095492
## 15 0.7863889 0.7095492
## 17 0.7863889 0.7095492
## 19 0.7763889 0.6935863
## 21 0.7380051 0.6346827
## 23 0.7280051 0.6155792
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was k = 5.

knnFit$finalModel

## 5-nearest neighbor model
## Training set outcome distribution:
##
## mammal bird reptile fish amphibian
## 41 20 5 13 4
## insect mollusc.et.al
## 8 10

accs <- accuracy(test_type, predict(knnFit, test))
accs

## [1] 0.96

#####
# 5. Neural Network
# Tuning parameters: 1) size (#Hidden Units)
# 2) decay (Weight Decay)

library(nnet)
nnetFit <- train(Zoo[,-17], Zoo[,17], "nnet",
  tuneLength = 5,
  trControl = trainControl(method = "cv", indexOut = train))

## # weights: 79
## initial value 225.600631
## iter 10 value 106.668955

```

```
## iter 20 value 59.575738
## iter 30 value 46.440020
## iter 40 value 32.127355
## iter 50 value 24.372662
## iter 60 value 18.818845
## iter 70 value 16.661347
## iter 80 value 16.206064
## iter 90 value 15.091604
## iter 100 value 13.797728
## final value 13.797728
## stopped after 100 iterations
```

nnetFit

```
## Neural Network
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 92, 91, 92, 88, 91, 91, ...
## Resampling results across tuning parameters:
##
##   size  decay  Accuracy  Kappa
##   1     0e+00  0.7400253  0.6344099
##   1     1e-04  0.7270960  0.6192481
##   1     1e-03  0.8521212  0.7946360
##   1     1e-02  0.8279798  0.7731977
##   1     1e-01  0.7370960  0.6451797
##   3     0e+00  0.9154545  0.8898625
##   3     1e-04  0.9436364  0.9269775
##   3     1e-03  0.9818182  0.9763331
##   3     1e-02  0.9818182  0.9767189
##   3     1e-01  0.9527273  0.9391183
##   5     0e+00  0.9818182  0.9768421
##   5     1e-04  0.9727273  0.9651400
##   5     1e-03  0.9818182  0.9767189
##   5     1e-02  0.9818182  0.9767189
##   5     1e-01  0.9818182  0.9767189
##   7     0e+00  0.9727273  0.9651374
##   7     1e-04  0.9818182  0.9768395
##   7     1e-03  0.9818182  0.9768395
##   7     1e-02  0.9818182  0.9767189
##   7     1e-01  0.9818182  0.9767189
##   9     0e+00  0.9818182  0.9767189
##   9     1e-04  0.9818182  0.9767189
##   9     1e-03  0.9818182  0.9767189
```

```

##      9      1e-02  0.9818182  0.9767189
##      9      1e-01  0.9818182  0.9767189
##
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were size = 3 and decay = 0.01.

nnetFit$finalModel

## a 16-3-7 network with 79 weights
## inputs: hairTRUE feathersTRUE eggsTRUE milkTRUE airborneTRUE aquaticTRUE
predatorTRUE toothedTRUE backboneTRUE breathesTRUE venomousTRUE finTRUE legs
tailTRUE domesticTRUE catsizeTRUE
## output(s): .outcome
## options were - softmax modelling  decay=0.01

accs <- accuracy(test_type, predict(nnetFit, test))
accs

## [1] 1

#####
# 6. Random Forest
# Tuning parameters: mtry (#Randomly Selected Predictors)

library(randomForest)

## randomForest 4.6-14

## Type rfNews() to see new features/changes/bug fixes.

##
## Attaching package: 'randomForest'

## The following object is masked from 'package:ggplot2':
##
##      margin

randomForestFit <- train(Zoo[, -17], Zoo[, 17], "rf",
                        tuneLength = 10,
                        trControl = trainControl(method = "cv", indexOut =
train))
randomForestFit

## Random Forest
##
## 101 samples
## 16 predictor
## 7 classes: 'mammal', 'bird', 'reptile', 'fish', 'amphibian', 'insect',
'mollusc.et.al'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)

```

```

## Summary of sample sizes: 92, 91, 92, 91, 92, 91, ...
## Resampling results across tuning parameters:
##
##   mtry Accuracy  Kappa
##    2      1      1
##    3      1      1
##    5      1      1
##    6      1      1
##    8      1      1
##    9      1      1
##   11      1      1
##   12      1      1
##   14      1      1
##   16      1      1
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.

randomForestFit$finalModel

##
## Call:
## randomForest(x = x, y = y, mtry = param$mtry)
##           Type of random forest: classification
##           Number of trees: 500
## No. of variables tried at each split: 2
##
##           OOB estimate of  error rate: 2.97%
## Confusion matrix:
##           mammal bird reptile fish amphibian insect mollusc.et.al
## mammal          41    0      0    0      0      0      0
## bird             0   20      0    0      0      0      0
## reptile          0    1      3    1      0      0      0
## fish             0    0      0   13      0      0      0
## amphibian        0    0      1    0      3      0      0
## insect           0    0      0    0      0      8      0
## mollusc.et.al    0    0      0    0      0      0     10
##
##           class.error
## mammal          0.00
## bird             0.00
## reptile          0.40
## fish             0.00
## amphibian        0.25
## insect           0.00
## mollusc.et.al    0.00

accs <- accuracy(test_type, predict(randomForestFit, test))
accs

## [1] 1

```

#####