

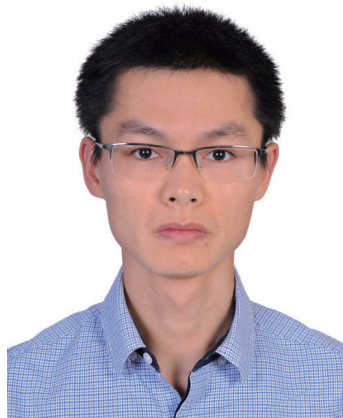
Decision tree modeling using R

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Author's introduction: Zhongheng Zhang, MMed. Department of Critical Care Medicine, Jinhua Municipal Central Hospital, Jinhua Hospital of Zhejiang University. Dr. Zhongheng Zhang is a fellow physician of the Jinhua Municipal Central Hospital. He graduated from School of Medicine, Zhejiang University in 2009, receiving Master Degree. He has published more than 35 academic papers (science citation indexed) that have been cited for over 200 times. He has been appointed as reviewer for 10 journals, including *Journal of Cardiovascular Medicine*, *Hemodialysis International*, *Journal of Translational Medicine*, *Critical Care*, *International Journal of Clinical Practice*, *Journal of Critical Care*. His major research interests include hemodynamic monitoring in sepsis and septic shock, delirium, and outcome study for critically ill patients. He is experienced in data management and statistical analysis by using R and STATA, big data exploration, systematic review and meta-analysis.



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Abstract: In machine learning field, decision tree learner is powerful and easy to interpret. It employs recursive binary partitioning algorithm that splits the sample in partitioning variable with the strongest association with the response variable. The process continues until some stopping criteria are met. In the example I focus on conditional inference tree, which incorporates tree-structured regression models into conditional inference procedures. While growing a single tree is subject to small changes in the training data, random forests procedure is introduced to address this problem. The sources of diversity for random forests come from the random sampling and restricted set of input variables to be selected. Finally, I introduce R functions to perform model based recursive partitioning. This method incorporates recursive partitioning into conventional parametric model building.

Keywords: Machine learning; R; decision trees; recursive partitioning; conditional inference; random forests

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Understanding decision tree

Decision tree learner is a technique of machine learning. As its name implies, the prediction or classification of outcomes is made going from root to leaves. The tree is made up of decision nodes, branches and leaf nodes. The tree is placed upside down, so the root is at the top and leaves indicating an outcome category is put at the bottom. At the root, all classifications are mixed, representing the original dataset. Then the tree grows to the first node where a certain feature variable is used to split the population into categories. Because the parent population can be split into in numerous patterns, we are interested in the one with the greatest purity. In technical terminology, purity can be described by entropy.

$$\text{entropy} = \sum_{i=1}^c -p_i \log_2(p_i)$$

where c is the number of different classifications. For the research of mortality outcome, c equals to two. p_i is the proportion of observations falling into class i . From the formula, we can see that entropy is zero when the population is completely homogenous; and 1 indicates the maximum degree of impurity. Other statistics such as Gini index, Chi-square statistics and gain ratio are also employed to describe the purity and to decide the best splitting (1).

The tree will stop growing by the following three criteria: (I) all leaf nodes are pure with entropy of zero; (II) a prespecified minimum change in purity cannot be made with any splitting methods; and (III) the number of observations in the leaf node reaches the prespecified minimum one (2). There are algorithms that allow the tree to grow and then the tree is pruned. Pruning technique aims to address the problem of overfitting as can occur in other parametric regression models (3).

Recursive partitioning within conditional inference framework, a type of decision tree learner, was developed by Hothorn and coworkers (2,4). It has many attractive features. Mathematical details of the technical are out of the scope of this paper. The word “recursive” refers to the fact that the decision nodes are repeatedly split. The algorithm works in the following steps:

Test the independence between any of the input variables and the response. Stop if there is no dependence between variables and response variable. Otherwise select the input variable with strongest association to the response. Permutation test is employed for the test of dependence (5). This step can prevent overgrowth of the tree (model overfitting) by statistical test;

Split observations in the decision node in the selected input variable;

Recursively repeat steps (I) and (II) until any of the stop criteria is reached.

Motivating example

We use the *airquality* dataset for the illustration of how to work with decision tree using R. The dataset contains information on New York air quality measurement in 1973. We obtain the dataset and examine the data frame with the following codes.

```
> data(airquality)
> str(airquality)
'data.frame': 153 obs. of 6 variables:
 $ Ozone: int 41 36 12 18 NA 28 23 19 8 NA ...
 $ Solar.R: int 190 118 149 313 NA NA 299 99 19 194 ...
 $ Wind: num 7.4 8 12.6 11.5 14.3 14.9 8.6 13.8 20.1 8.6 ...
 $ Temp: int 67 72 74 62 56 66 65 59 61 69 ...
 $ Month: int 5 5 5 5 5 5 5 5 5 ...
 $ Day: int 1 2 3 4 5 6 7 8 9 10 ...
```

The data frame contains 153 observations and 6 variables. In the example, we will set the *Ozone* as response variable. However, there are missing values in the variable. We first have to impute these missing values, or simply exclude observations with missing values in *Ozone*. Because dropping observations result in information loss, imputation method is used to address this problem (6). We impute missing *Ozone* values with values randomly selected from the non-missing ones. The `sample()` function allows taking a sample of the specified size from given elements. Here the size of the sample is 37, and the given element is a vector containing non-missing *Ozone* values.

```
> set.seed(888)
> airquality[is.na(airquality$Ozone),1]<-
sample(airquality[!is.na(airquality$Ozone),1],37)
> summary(airquality$Ozone)
Min.    1st Qu.  Median    Mean    3rd Qu.    Max.
1.00    18.00    29.00    40.82    61.00    168.00
```

From the output of `summary()` function, it is apparent that *Ozone* contains no missing value any more.

Conditional inference tree

In this section we will use the imputed dataset to build a conditional inference tree. Owing to the `ctree()` function in *party* package, the code of building a tree is fairly simple (4).

```
> airct <- ctree(Ozone ~ ., data = airquality, controls =
ctree_control(maxsurrogate = 3))
```

The `ctree()` function created a conditional inference tree and returned an object of class *BinaryTree-class*. The first argument of the function is a formula defining the response and input variables. The format of the equation is similar to that used in building generalized linear model. “data” argument defines the data frame which the model is built upon. The `ctree_control()` function tunes various parameters to control aspects of the conditional inference tree fit. In the example, we adjusted the number of surrogate splits to evaluate to 3. We can take a look at the tree structure as follows.

```
> airct
```

Conditional inference tree with 5 terminal nodes

Response: Ozone

Inputs: Solar.R, Wind, Temp, Month, Day

Number of observations: 153

- 1) Temp <= 82; criterion = 1, statistic = 41.908
- 2) Wind <= 6.3; criterion = 1, statistic = 15.879
- 3)* weights = 8
- 2) Wind > 6.3
- 4) Solar.R <= 81; criterion = 0.977, statistic = 7.997
- 5)* weights = 26
- 4) Solar.R > 81
- 6)* weights = 71
- 1) Temp > 82
- 7) Wind <= 8; criterion = 0.977, statistic = 8.043
- 8)* weights = 29
- 7) Wind > 8
- 9)* weights = 19

The above output gives you a general glimpse of the tree structure. The response variable is *Ozone*, and input

variables include *Solar.R*, *Wind*, *Temp*, *Month* and *Day*. We can visualize the tree graphically using the generic function `plot()`, and it is more convenient to interpret the tree with graphics.

```
> plot(airct)
```

The decision nodes are represented by circles, and each circle is numbered. The input variable to split upon is shown in each circle, together with the P value of the dependence test. For example, the first decision node at the top shows that *Temp* is the variable that is most strongly associated with *Ozone* and thus is selected as the first node. The association is measured by a $P < 0.001$. The left and right branches show that a cutoff value equal to 82 is the best to reduce impurity. Thereafter, the decision nodes are further divided by variables *Wind* and *Solar.R* sequentially. The leaf nodes show the classifications of *Ozone* by the decision tree. Because variable *Ozone* is a continuous variable, the result is displayed by box plot (*Figure 1*).

Decision tree plot can be modified by setting graphical parameters. For example, if you want to display boxplot, instead of statistics, in inner nodes (decision nodes). The following code can be used:

```
> plot(airct, inner_panel = node_boxplot, edge_panel =
function(...) invisible(), tnex = 1)
```

As you can see from *Figure 2*, all inner nodes show boxplot of observations.

Sometimes investigators are interested in examining the split statistics based on which the split point is derived (4). Remember that the split input variable is continuous variable and there are numerous split points to choose from. For a given split point there is a statistic measuring how good the splitting is. A greater statistic value is associated with a better splitting. Split statistics can be derived from *splitstatistic* element of a split. The following syntax draws scatter plots of split statistics for inner nodes 1, 2, 4 and 7. The leaf nodes have no split statistics because they do not split any more (the tree stops growing at leaf node, *Figure 3*).

```
> inner <- nodes(airct, c(1, 2, 4, 7))
> layout(matrix(1:length(inner), ncol = length(inner)/2))
> out <- sapply(inner, function(i) {
splitstat <- i$psplit$splitstatistic
```

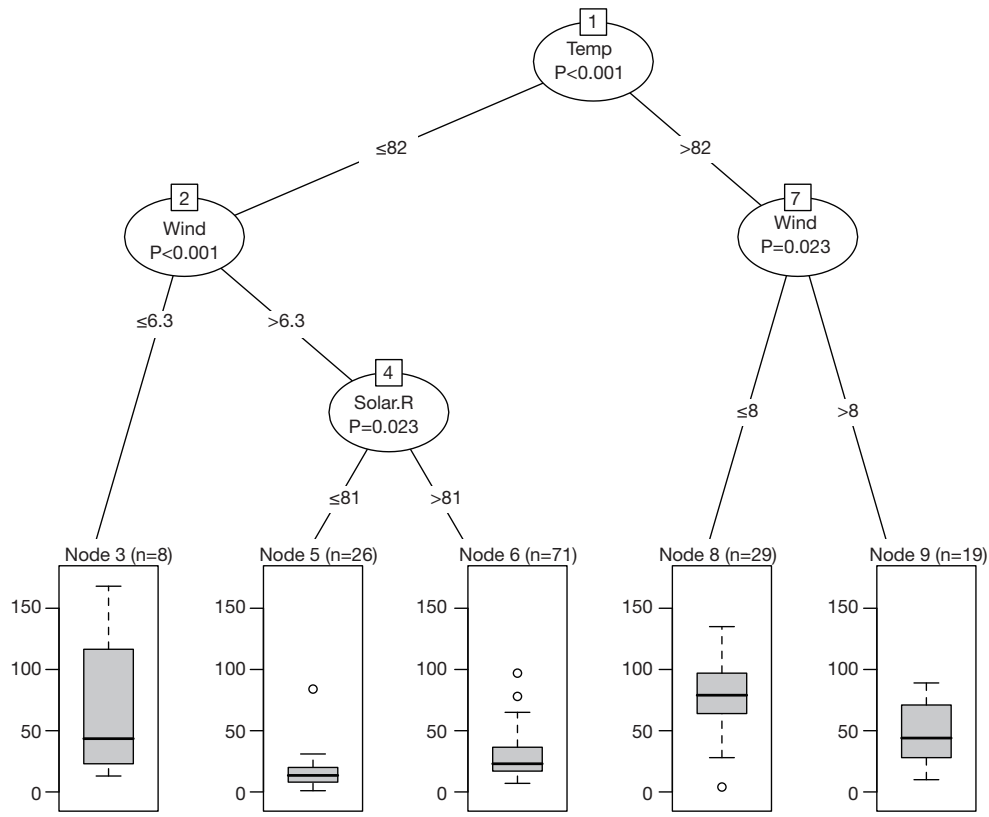


Figure 1 Conditional inference tree for the airquality data. For each inner node, input variable and P values are given, the boxplot of ozone value is displayed for each terminal node.

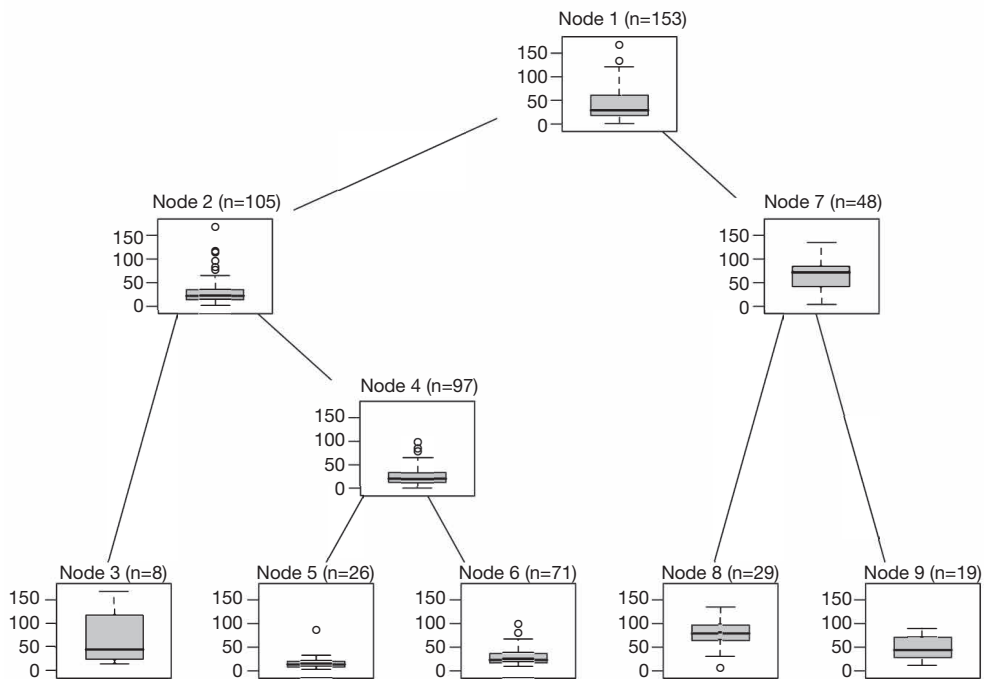


Figure 2 Conditional inference tree for the airquality data with the ozone value displayed for both inner and terminal nodes.

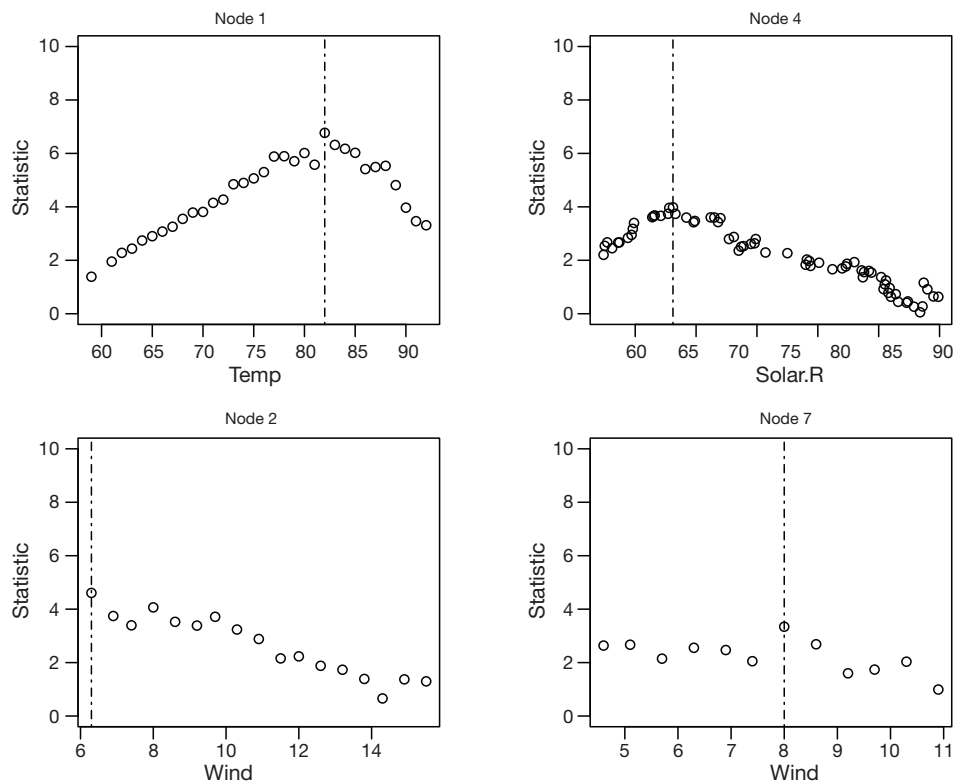


Figure 3 Split point estimation based on split statistics in inner nodes 1, 2, 4 and 7. Standardized two-sample test statistics are plotted against each possible split point in the selected input variable. The estimated split point is denoted by a vertical dotted line.

```
x <- airquality[[i$psplit$variableName]][splitstat > 0]
plot(x, splitstat[splitstat > 0], main =
paste("Node", i$nodeID), xlab = i$psplit$variableName,
ylab = "Statistic", ylim = c(0, 10), cex.axis = 1.2, cex.lab =
1.2, cex.main = 1.2)
abline(v = i$psplit$splitpoint, lty = 4)
})
```

Prediction with decision tree

The purpose of building a decision tree model is to predict responses in future observations. A subset of the *airquality* data frame is employed as a new cohort of observations.

```
> ind <- sample(2, nrow(airquality), replace=TRUE, prob =
c(0.7, 0.3))
> newData <- airquality[ind==2,]
> newpred <- predict(airct, newdata= newData)
```

```
> plot(newpred, newData$Ozone, xlab="Ozone value
predicted by decision tree", ylab="Observed ozone value")
```

The `sample()` function returns indicator vector of the same length to the row number of the *airquality* data frame. There are two levels of “1” and “2” in this vector. Observations in the *airquality* data frame denoted by “2” are selected to constitute the *newData*. Then the `predict()` function is used to make a prediction based on input variables. Because the decision tree results in five classifications (5 leaf nodes), the `predict()` function returns a vector with five values. Each value is the mean value of observations in each leaf nodes (*Figure 4*). We plot predicted value against observed value in a scatter plot. Not surprisingly, the predicted values take only five values.

Random forests

In the above sections, I grow a single tree for prediction. This method has been criticized for its instability to small changes

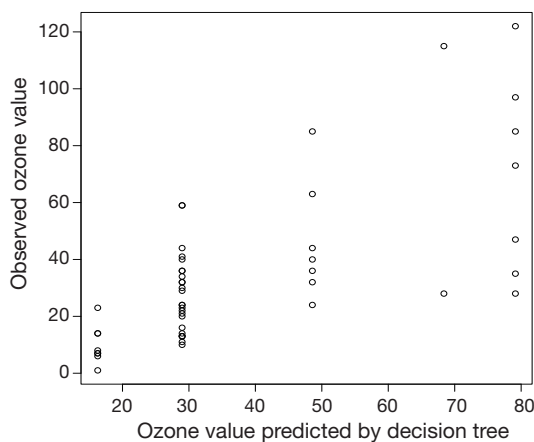


Figure 4 Scatter plot showing predicted value based on conditional inference tree against observed value. The predicted values are only allowed to take five values that are the mean values of each classification.

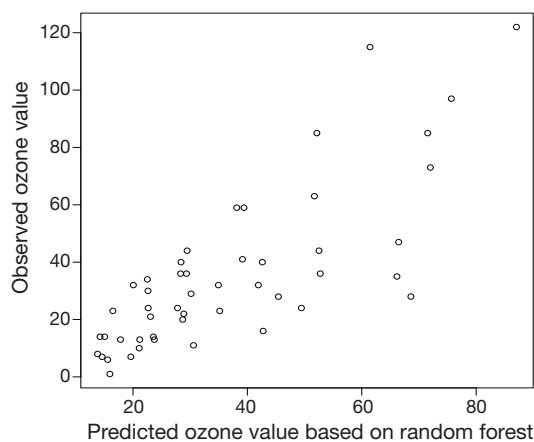


Figure 5 Scatter plot showing the observed ozone values against predicted values by random forests model. This time the predicted value is not restricted to five levels. Instead, they are evenly distributed across the ozone range, which is more likely to occur in the real world.

in the training data. In recursive partitioning framework, the decision on which input variable to split in and the split point determine how observations are split into leaf nodes. The exact split position and input variable to be selected are highly dependent on the training data. A small change to the training data may alter the first variable to split in, and the appearance of tree can be completely altered. One approach to this problem is to grow a forest instead of one tree (7). Diversity is introduced from two sources: (I) a set of trees is built on random samples of the training sample; and (II) the set of variables to be selected from in each decision node is restricted. Trees of the forest can be built on bootstrap

samples or a subsample of the training dataset. If there are 10 variables to be selected from, we can restrict it to 5 at given split node. The five variables are randomly selected and the pools of candidate variables can be different across forest trees. The final prediction model is the combination of all trees (8).

```
> aircf<-cforest(Ozone ~ ., data = airquality)
> aircf
      Random Forest using Conditional Inference Trees
Number of trees: 500
Response: Ozone
Inputs: Solar.R, Wind, Temp, Month, Day
Number of observations: 153
```

The syntax for fitting a random forests model is similar to a single conditional inference tree. The output of *aircf* shows that we have grown 500 trees. The response variable is *Ozone* and input variables are *Solar.R*, *Wind*, *Temp*, *Month* and *Day*. Tree plot cannot be drawn because these trees are different in structure. However, we can make prediction with the random forests model. Again, we use the *predict()* function, but this time the model is *aircf*.

```
> predforest<-predict(aircf, newdata= newData)
> plot(predforest,newData$Ozone,ylab="Observed
ozone value",xlab="Predicted ozone value based on
random forest")
```

In the graphical output (*Figure 5*), observed ozone values are plotted against predicted values by random forests model. This time the predicted value is not restricted to five levels. Instead, they are evenly distributed across the ozone range, which is more likely to occur in the real world.

Model based recursive partitioning

Recursive partitioning can be incorporated into conventional parametric model building. Model based recursive partitioning algorithm consists of the following steps: (I) a parametric model (such as linear model and logistic regression model) is fit to current dataset; (II) parameter instability is tested over partitioning variables; (III) tree node is split by partitioning variable associated with highest parameter instability; (IV) repeat above steps in each of the daughter nodes (9). Model based recursive partitioning algorithm can be easily performed using R.

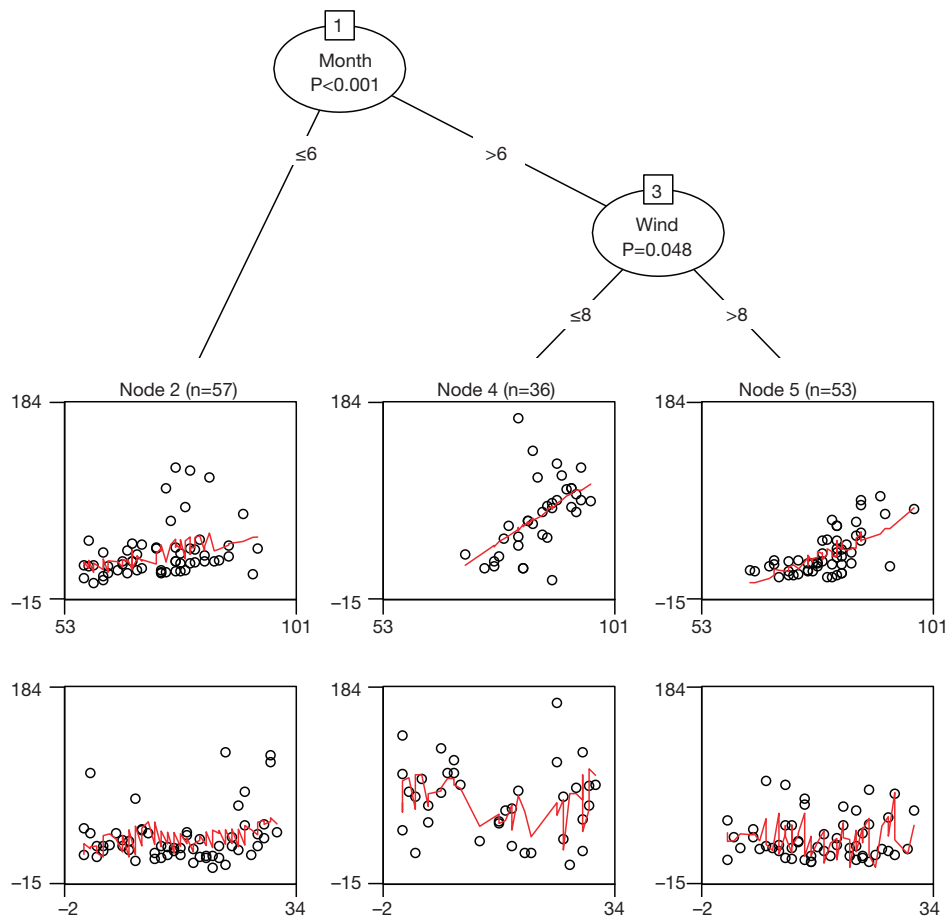


Figure 6 Linear-regression-based tree for the airquality data. The leaf nodes give partial scatter plots for *Temp* (upper panel) and *Day* (lower panel).

```
> airmod <- mob(Ozone ~ Temp + Day | Solar.
R + Wind + Month, data = airquality)
> plot(airmod)
```

In the above code, we used variables *Temp* and *Day* to construct a linear regression model with the *Ozone* as the response variable. Variables after symbol “|” are partitioning variables. The result is displayed with decision tree plot (Figure 6). Because *Temp* is used as a predictor in regression model, it no longer participates in partitioning. Thus the first partition is made on the variable *Month*. There seems no strong association between *Day* and *Ozone* as shown in the Figure. Therefore, we need to take a close look at statistics of the model.

```
> airmod
1) Month <= 6; criterion = 1, statistic = 32.402
```

```
2)* weights = 57
Terminal node model
Gaussian GLM with coefficients:
(Intercept) Temp Day
-52.3217 0.9880 0.7325
```

```
1) Month > 6
3) Wind <= 8; criterion = 0.952, statistic = 15.979
4)* weights = 36
```

```
Terminal node model
Gaussian GLM with coefficients:
(Intercept) Temp Day
-200.2079 3.1813 -0.1323
```

```
3) Wind > 8
5)* weights = 53
```


Terminal node model**Gaussian GLM with coefficients:**

```
(Intercept) Temp Day
-149.8501 2.1784 0.5631
```

The above output displays coefficient of fitted linear regression model. The coefficient of *Temp* varied remarkably across leaf nodes, indicating a good partitioning. The coefficient of each leaf nodes can be extracted using `coef()` function.

```
> coef(airmod)
      (Intercept)      Temp      Day
2      -52.32172    0.9880186  0.7325495
4     -200.20787    3.1812577 -0.1323226
5     -149.85014    2.1783656  0.5630807
```

The output of `coef()` function displays model coefficient for each leaf node. Test for parameter stability can be examined with the `sctest()` function by passing a model based tree and node number to the function.

```
> sctest(airmod, node = 1)
      Solar.R      Wind      Month
statistic 7.3785883 3.016852e+01 3.240226e+01
p.value 0.8682267 1.012705e-04 3.405009e-05
```

In the above output, we can see that the P value for *Month* is the smallest and thus split of the first decision node is made on *Wind* (Figure 6).

Summary

The article introduces some basic knowledge on decision tree learning. It employs recursive binary partitioning algorithm that splits the sample in partitioning variable with the strongest association with the response variable. The process continues until some certain criteria are met. In the example I focused on conditional inference tree, which incorporated tree-structured regression models into conditional inference procedures. While growing a single tree is subject to small changes in the training data,

random forests procedure is introduced. The sources of diversity for random forest come from the random sampling and restricted set of input variables to be selected. Finally, I introduce R functions to perform model based recursive partitioning. This method incorporates recursive partitioning into conventional parametric model building.

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Footnote

Conflicts of Interest: The author has no conflicts of interest to declare.

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