ITR Forest: Constructing a Treatment Decision Rule

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Introduction

The package ITR forest creates an individualized treatment decision rule using a recursive partitioning algorithm to grow a decision tree. The algorithm utilizes either an inverse probability weighted estimator (IPWE) or augmented inverse probability weighted estimator (AIPWE) as the reward function, maximizing the selected function at each split. The current version of this package can be applied to data from completed randomized controlled trials (RCTs) or an electronic medical record (EMR) database where there are two treatment options, a continuous outcome, and continuous or categorical predictors. Additionly, each patient needs to have an estimated propensity score, or probability of being assigned to treatment, $\hat{p}(t_i|\mathbf{x}_i)$. For an RCT with 1:1 treatment allocation, for example, $\hat{p}(t_i|\mathbf{x}_i) = 0.5$. For EMR data, $\hat{p}(t_i|\mathbf{x}_i)$ can be found using logistic regression. Without loss of generality, we assume that larger values of the outcome are better.

The IPWE is estimated from the data as

$$\hat{V}_{IPWE}(r) = \left(\sum_{i=1}^{N} \frac{\mathbf{I}_{t_i = r(\mathbf{x}_i)}}{\hat{p}(t_i | \mathbf{x}_i)} y_i\right) / \left(\sum_{i=1}^{N} \frac{\mathbf{I}_{t_i = r(\mathbf{x}_i)}}{\hat{p}(t_i | \mathbf{x}_i)}\right).$$

The AIPWE is estimated from the data as

$$\hat{V}_{AIPWE}(r) = \hat{V}_{IPWE}(r) - \left(\sum_{i=1}^{N} \frac{\mathbf{I}_{t_i = r(\mathbf{x}_i)} - \hat{p}(t_i|\mathbf{x}_i)}{\hat{p}(t_i|\mathbf{x}_i)} m(X_i)\right),$$

where $m(X_i) = \mu(t_i = 1, X_i) \cdot z_i + \mu(t_i = 0, X_i) \cdot (1 - z_i)$ is the estimated mean in a given node and is assigned based on the patient's original treatment assignment $t_i \in \{0, 1\}$ and the new treatment assignment under consideration $z_i \in \{0, 1\}$.

Further details can be found in "A Novel Algorithm for Generating Individualized Treatment Decision Trees and Random Forests" by Doubleday and Zhou (In Review).

The input dataset needs to have the following: (1) propensity score column labeled prtx, (2) binary treatment column labeled trt, (3) ID column labeled id, (4) continuous outcome column labeled y, and (5) columns of predictors.

Growing a Tree with Continuous Predictors

A single ITR tree can be constructed using the function grow.ITR which requires an input dataset and a set of splitting variable columns. By default the number of observations allowed in a terminal node is 20 (min.ndsz = 20), there must be at least 5 observations from each treatment group in a terminal node (n0 = 5), and the maximum tree depth is set at 15 (max.depth = 15). The initial value of the root node is the maximum of $\hat{V}(r)$ with all subjects given treatment or all patients given control. A initial split of the root node is only made if there is a split for which the value in the root node increases. The same is true for additional splits so that the tree cannot grow larger unless there is an increase in overall value of the tree given by the split.

Growing a Tree

We will use the following example dataset generated from the function gdataM() which simulates EMR data from the model

$$Y = 1 + 2X_2 + 4X_4 + \beta_1 * T * (\mathbf{X} \in A) + \beta_2 * (1 - T) * (\mathbf{X} \notin A) + \epsilon.$$

where $A = \{X_1 > 0.3 \cap X_3 > 0.1\}$ when depth=2 is specified. Covariates $X_1 - X_4 \sim \text{Unif}(0,1)$, errors follow N(0,1), treatment assignments are $T \in \{0,1\}$, prtx is the propensity score, and the signal to noise ratio θ is defined as β_1/β_2 .

```
set.seed(123)
dat \leftarrow gdataM(n = 2500, beta1 = 3, beta2 = 1, depth = 2)
head(dat)
       Х1
##
             Х2
                  ХЗ
                       Х4
                                  y trt prtx id
## 1 0.30 0.86 0.26 0.08 4.215369
                                      0 0.142
## 2 0.80 0.68 1.00 0.72 7.964857
                                       1 0.968
                                                2
## 3 0.42 0.74 0.72 0.86 4.827130
                                      0 0.703
                                                3
## 4 0.90 0.32 0.66 0.68 8.038266
                                      1 0.881
## 5 0.96 0.68 0.26 0.76 6.911391
                                       1 0.545
                                                5
## 6 0.06 0.48 0.10 0.78 8.211139
                                      0 0.035
set.seed(10)
covExtra<-matrix(sample(1:25, size = 10*nrow(dat), replace = T), nrow=nrow(dat))/25</pre>
colnames(covExtra)<-paste("E", seq(1,10), sep = "")</pre>
dat <- cbind(dat, covExtra)</pre>
```

This dataset has 2500 observations with $\theta = 3$. The argument depth=2 indicates that there are two subgroup defining variables, X_1 and X_3 . Changing this argument to depth=1 would change the subgroup definition to $X_1 < 0.5$, having only a single interacting covariate. Additionally, there are 10 noise variables added which are not shown above and which users can generate on their own if they would like. The tree is constructed using the grow.ITR() function as follows with the AIPWE since we are analyzing an EMR dataset.

```
tre <- grow.ITR(data = dat, split.var = c(1:4, 9:18), AIPWE = TRUE)
tre
##
        node size
                   n.1 n.0 trt.effect var
                                              vname cut.1 cut.2
## 1
           0 2500 1294 1206
                                                              0.3 0.7426
                               1.2540902
                                             1
                                                  X1
                                                          r
## 3
           01
               758
                    248
                         510 -0.9064623
                                           12
                                                  E4
                                                          1
                                                             0.04 0.8245
## 6
         011
                34
                       9
                           25 -0.9289875
                                           NA
                                                <NA>
                                                       <NA>
                                                             <NA>
                                                                       NA
## 5
         012
               724
                    239
                          485 -0.9053264
                                           11
                                                  ЕЗ
                                                          1
                                                             0.04 0.8260
        0121
                25
                                                <NA>
                                                             <NA>
## 8
                       9
                           16 -0.3540211
                                           NA
                                                       <NA>
                                                                       NA
        0122
               699
                    230
                          469 -0.9250415
                                                             0.96 0.8279
## 9
                                           18
                                                 E10
                                                          r
       01221
                                                             0.02 0.8292
## 11
               678
                    221
                          457 -0.9461866
                                            1
                                                  Х1
                                                          1
## 13 012211
                45
                       7
                           38 -0.4767850
                                                <NA>
                                           NA
                                                      <NA>
                                                             <NA>
                                                                       NA
## 12 012212
               633
                    214
                          419 -0.9734814
                                           NA
                                                <NA>
                                                      <NA>
                                                             <NA>
                                                                       NA
## 10
       01222
                21
                       9
                           12 -0.3644231
                                           NA
                                                <NA>
                                                       <NA>
                                                             <NA>
                                                                       NA
## 2
           02 1742 1046
                               2.1685733
                                            3
                                                  ХЗ
                          696
                                                          r
                                                              0.1 0.8225
## 4
         021
               168
                      28
                          140 -1.4782590
                                                <NA>
                                                      <NA>
                                                             <NA>
                                                                       NA
                                           NA
## 7
         022 1574 1018
                          556
                               2.5396112
                                           NA
                                                <NA>
                                                      < NA >
                                                             <NA>
                                                                       NA
```

The output contains a summary of the tree structure. The node column begins with the root node 0 and each subsequent number indicates the direction of the split, with 1 indicating the left (less than or equal to) node and 2 indicating the right (greater than) node. The first row indicates that the covariate X_1 is selected as the first splitting variable with a cut point of cut.2 = 0.3. The decision is to send treatment to the right node (cut.1 = "r"). size, n.1, and n.0 indicate there are 2500 observations in the root node, with 1294 treated and 1206 on control. The second row with node = 01 contains information from the left child node with interpretations similar to those described for the root node. The splitting information denoted by NA indicates a terminal node, 011 for instance.

Note that in the case of this simulated data (depth 2) the correct tree structure splits the root node at $X_1 = 0.3$ and sends treatment to node 02. Next, node 02 should be split at $X_3 = 0.1$ and sends treatment to the right. We see that this tree structure represents this well, but has some additional splits which are not necessary.

Pruning a Tree

To avoid overfitting, a pruning procedure is introduced to penalize additional splits in the tree growing process. The function prune(tre, a, train, test) accomplishes this with penalty $\lambda = a$ applied using the weakest link criteria. The penalty is applied to subtree Γ .

$$V_{\lambda}(\Gamma) = V(\Gamma) - \lambda \cdot |\Gamma - \tilde{\Gamma}|$$

where $|\Gamma - \tilde{\Gamma}|$ is the number of internal nodes of subtree Γ , $V(\Gamma)$ is the value of the entire subtree, and $V_{\lambda}(\Gamma)$ is the penalized value. We trim the weakest branches first which are those with (1) the greatest number of parent nodes, and (2) contributes the smallest additional value to the tree. We can prune a tree as follows. The following example uses the tree above (tre) with a penalty of 0.05.

```
pruned <- prune(tre, a = 0.05, train = dat)
pruned</pre>
```

```
##
     subtree node.rm size.tree size.tmnl
                                             alpha
                                                          V
                                                               V.a V.test Va.test
## 1
                   01
                              13
                                          7 6.2855 6.3001 6.0001
                                                                      <NA>
            1
                                                                               <NA>
## 2
            2
                   02
                                          3 6.1903 6.2855 6.1855
                                                                               <NA>
                               5
                                                                      <NA>
## 3
            3
                    0
                               3
                                          2 5.9411 6.1903 6.1403
                                                                      <NA>
                                                                               <NA>
## 4
                                1
                                               9999 5.9411 5.9411
                   NA
                                                                      <NA>
                                                                               <NA>
```

The first row represents the entire tree (subtree 1) and summarizes the value, penalized value, and weakest node (node to be removed next) by V, V.a, and node.rm. The size.tree and size.tmnl columns give the total number of nodes and terminal nodes in a given subtree. We want to select the tree with the highest penalized value, which would correspond to subtree 2 with a penalized value of V.a = 6.1855. This subtree has 3 terminal nodes.

Cross Validation for Model Selection

One issue with the approach above for model selection is the risk of overfitting through using the training data alone for model selection. The function treeCV() will perform n-fold cross validation to select the optimal tuning parameter (λ). The function returns the optimal model, selected penalty, and several summary measures.

[1] 0.04

```
n.0 trt.effect var vname cut.1 cut.2
##
     node size
                n.1
## 1
        0 2500 1294 1206
                                               X1
                                                            0.3 0.7426
                            1.2540902
                                          1
                                                       r
##
                 248
                       510
                           -0.9064623
                                        NA
                                             <NA>
                                                    <NA>
                                                           <NA>
                                                                    NA
##
                                          3
                                               ХЗ
                                                           0.1
                                                                0.8225
       02
          1742
                1046
                       696
                            2.1685733
                                                       r
##
   4
      021
            168
                   28
                       140
                           -1.4782590
                                        NA
                                             <NA>
                                                    <NA>
                                                           <NA>
                                                                    NA
      022 1574 1018
                       556
                            2.5396112
                                        NA
                                             <NA>
                                                           <NA>
                                                    <NA>
                                                                    NA
```

The optimal lambda selected is 0.04 and the optimal tree has 3 terminal nodes. This tree structure corresponds to the correct tree structure for this particular simulation. Figure 1 shows the results of the cross validation as λ increases.

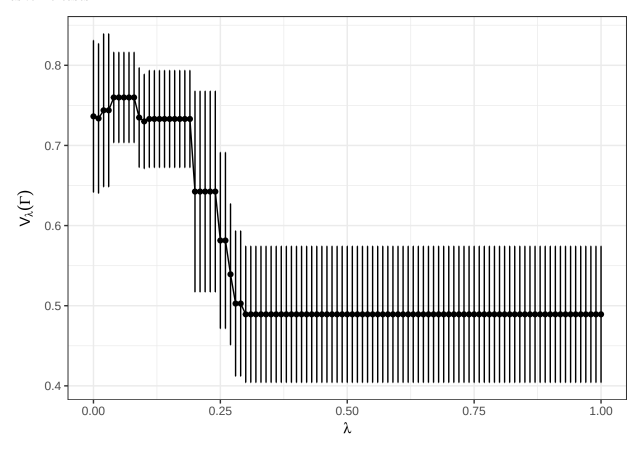


Figure 1. Cross validated value versus λ with 95% error bars.

Constructing an ITR Forest to Give Decision Rule

A single tree which is trained using all available data may be overfitted and not extendable to subsequent observations. Hence, we make a decision rule using a forest of ITR trees in which each tree is more variable, but the aggregation of the trees in the forest mitigates this variance. The ITR forest is contructed using the function Build.RF.ITR() and requires the entry of a dataset, columns for the outcome, treatment, propensity score, and splitting variables. To randomized the growth of trees in the forest a subset of predictors, mtry, is selected as potential splitting variables at each split which defaults to the maximum of 1/3 the number of splitting variables and 1. The number of observations and the number of treated and control subjects allowed in a terminal node is given by NO and nO. By default the number of trees contructed, ntree, is 500. Each tree is grown using a bootstrap sample taken from the input dataset. The function returns the bootstrap samples used in tree construction, the trees, and the model parameters. The forest is contructed as follows and the first two trees are displayed as examples.

```
set.seed(2)
forest <- Build.RF.ITR(dat, split.var = c(1:4, 9:18), col.y="y", col.trt="trt",</pre>
                        col.prtx="prtx", ntree = 500)
forest$TREES[1:2]
## [[1]]
##
     node size n.1 n.0 trt.effect var vname cut.1 cut.2
## 1
        0 1583 813 770 1.2594453
                                      1
                                            Х1
                                                   r
                                                       0.3 0.7173
## 2
       01
           483 160 323 -0.9498811
                                     NA
                                         <NA>
                                                <NA>
                                                      <NA>
                                                                NA
## 3
       02 1100 653 447
                         2.1869441
                                      3
                                                       0.1 0.8139
                                           ХЗ
                                                   r
## 5
      021
           105
                17
                     88 -1.8317697
                                     NA
                                         <NA>
                                                <NA>
                                                      <NA>
                                                                NA
      022
           995 636 359
## 4
                         2.5747458
                                     NA
                                         <NA>
                                                <NA>
                                                      <NA>
                                                                NA
##
## [[2]]
     node size n.1 n.0 trt.effect var vname cut.1 cut.2 score
##
        0 1576 831 745
## 1
                          1.309622
                                     NA
                                            NA
                                                  NA
                                                        NA
                                                               NA
```

We see that the first tree corresponds to the expected result and the second tree is a null tree. If the user wants, the generation of null trees can be avoided using the additional argument avoid.nul.tree = TRUE. We can now run a new observation down each of the trees and obtain a vote from each tree as to what the treatment assignment should be. The majority vote from the forest will be the ITR forest decision rule. This can be obtained using the predict.ITR() function. First we generate a new observation and second we make a treatment predition for this new observation. Shown is the treatment summary as the proportion of trees voting for treatment and the voting record for the first 10 trees.

```
set.seed(10)
new.obs \leftarrow gdataM(n = 1, depth = 2, beta1 = 3, beta2 = 1)
##
       Х1
             Х2
                  X3 X4
                                 y trt prtx id
## 1 0.52 0.32 0.44 0.7 3.834144
                                     0 0.44 1
new.obs <- cbind(new.obs, matrix(sample(1:25, size = 10, replace = TRUE)/25, ncol = 10))
colnames(new.obs) <- colnames(dat)</pre>
preds <- predict.ITR(forest, new.obs)</pre>
preds$SummaryTreat
## [1] 1
preds$tree.votes[1:20]
```

```
[1]
   1 NA 1 1 NA
               1 NA 1 1 1 NA 1 1 1 1 NA 1 NA NA
```

Note that the observation should receive treatment since $X_1 > 0.3$ and $X_3 > 0.1$. This forest has 100% of the trees voting that the patient should receive treatment which would be a good decision for this patient since they are in the subgroup which benefits from treatment. Note that several of the trees return votes of NA. This means that the tree did not make an initial split, or was a null tree.

Predictions for outside data

Use the predict.ITR() function to make treatment predictions for a forest object created using Build.RF.ITR() or a single tree object created using grow.ITR(). For a forest, the output includes the proportion of trees voting for the treatment (trt = 1), votes from all the trees, the number of null trees in the forest, and a summary of the data and trees. An example for a single tree is shown below using the tree tree from above and 4 newly generated observations.

```
set.seed(1)
new.dat <- gdataM(4, 2, 1, 1)
new.dat <- cbind(new.dat, matrix(sample(1:25, size = 10, replace = TRUE)/25, ncol = 10))
colnames(new.dat) <- colnames(dat)</pre>
preds <- predict.ITR(tre, new.dat)</pre>
new.dat
##
       X1
            X2
                 ХЗ
                                 y trt prtx id E1
                                                       F.2
                                                            E3
                                                                  E4
                                                                       E5
                                                                            E6
## 1 0.28 0.22 0.64 0.70 6.364931
                                               1 0.8 0.12 0.76 0.44 0.84 0.68
                                     0 0.510
## 2 0.38 0.90 0.08 0.40 4.789843
                                     1 0.079
                                               2 0.8 0.12 0.76 0.44 0.84 0.68
                                              3 0.8 0.12 0.76 0.44 0.84 0.68
## 3 0.58 0.96 0.22 0.78 6.023810
                                     0 0.239
## 4 0.92 0.68 0.18 0.50 3.145300
                                     1 0.416 4 0.8 0.12 0.76 0.44 0.84 0.68
##
      E7
           E8
                E9 E10
## 1 0.8 0.56 0.56 0.8
## 2 0.8 0.56 0.56 0.8
## 3 0.8 0.56 0.56 0.8
## 4 0.8 0.56 0.56 0.8
preds$SummaryTreat
## [1] 0 0 1 1
```

Each element of SummaryTreat is the treatment decision for one of the new observations.

Variable Importance

Last, we include a function to calculate the importance of a predictor in making the treatment assignment for an ITR forest. The variable importance is calculated by determining the out of bag (OOB) value $V_{OOB}(r)$ for the sample not used in tree construction (OOB sample), permuting the variable values for any predictor used in the tree construction, and re-running the OOB sample down the tree to obtain $V_{OOBpermuted}(r)$. The larger the difference between $V_{OOB}(r)$ and $V_{OOBpermuted}(r)$ the more important the predictor. This is done for each tree in the forest and importance measures for each variable are summed. We scale the measure to be out of 1 for easy interpretibility. This is done using the function Variable.Importance.ITR().

```
VI <- Variable.Importance.ITR(forest)</pre>
VI
                                                       E5
             X1
                            ХЗ
                                         X4
## 0.7064653941 0.2561928760 0.0209976418 0.0044585622 0.0027144093
             X2
                            E3
                                         E9
                                                       E7
## 0.0022568726 0.0016150340 0.0015710675 0.0012761554 0.0010778375
##
             E2
                          E10
                                         E4
## 0.0005656072 0.0003416131 0.0002766305 0.0001902988
```

We see that the variable forming the interaction subgroup, X_1 and X_3 , are returned as the most important predictors.

References

[In Review] Doubleday, K., Zhou, J., Fu, H. (2017), "A Novel Algorithm for Generating Individualized Treatment Decision Trees and Random Forests," *Journal of Computational and Graphical Statistics*.