Bayesian Treed Models*

HUGH A. CHIPMAN hachipman@uwaterloo.ca Department of Statistics and Actuarial Science, University of Waterloo, Waterloo, ON N2L 3G1, Canada

EDWARD I. GEORGE edgeorge@wharton.upenn.edu
Department of Statistics, The Wharton School, University of Pennsylvania, Philadelphia, PA 19104, USA

ROBERT E. McCULLOCH robert.mcculloch@gsb.uchicago.edu Graduate School of Business, University of Chicago, Chicago, IL 60637, USA

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Abstract. When simple parametric models such as linear regression fail to adequately approximate a relationship across an entire set of data, an alternative may be to consider a partition of the data, and then use a separate simple model within each subset of the partition. Such an alternative is provided by a treed model which uses a binary tree to identify such a partition. However, treed models go further than conventional trees (e.g. CART, C4.5) by fitting models rather than a simple mean or proportion within each subset. In this paper, we propose a Bayesian approach for finding and fitting parametric treed models, in particular focusing on Bayesian treed regression. The potential of this approach is illustrated by a cross-validation comparison of predictive performance with neural nets, MARS, and conventional trees on simulated and real data sets.

Keywords: binary trees, Markov chain Monte Carlo, model selection, stochastic search

1. Introduction

Parametric models such as linear regression can provide useful, interpretable descriptions of simple structure in data. However, sometimes such simple structure does not extend across an entire data set and may instead be confined more locally within subsets of the data. In such cases, the simple structure might be better described by a model that partitions the data into subsets and then uses separate submodels for the subsets of the partition. In this paper, we explore the use of "treed models" to describe such structure.

Basically, treed models are an elaboration of conventional tree models that use binary trees to partition the data into "homogeneous" subsets where the response can be described by a simple mean or proportion. Although such models can provide a useful approach to handling interactions and nonlinearities, they do not fully exploit partitions with more substantial statistical structure within the subsets. To overcome this limitation, treed models

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are constructed so that the *model structure*, as opposed to the data, is homogeneous within a terminal node. This allows richer submodels for each of the subsets of the tree-determined partition.

The main thrust of this paper is to propose and illustrate a Bayesian approach to finding and fitting treed models. In particular, we pay special attention to treed regressions, where linear regression models are used to describe the variation within each of these subsets. Alexander and Grimshaw (1996) coined the term "treed regression" to refer to a tree with simple linear regression in each terminal node. We adopt this terminology to include treed models with multiple regressions, and refer to our approach as Bayesian treed regression.

The idea of treed models is not new. Karalič (1992) considered multiple regression models in the terminal nodes, in conjunction with a greedy grow/prune algorithm. Quinlan's (1992) M5 algorithm also fits models of this form, but linear models were added to a conventional tree as part of the pruning stage. Torgo (1997) uses linear models, k-nearest neighbors, or kernel regressors in terminal nodes, but also adds these to a conventional tree after growing. Chaudhuri et al. (1994) consider trees with linear models (with possible polynomial terms) in the terminal nodes. Chaudhuri et al. (1995) extend this idea to non-normal response models. Loh (2002) also considers linear regressions in terminal nodes, and attempts to correct for possible bias in selection of splits. Waterhouse, MacKay, and Robinson (1996) consider a tree-structured mixture of experts model, in which the individual experts are linear models. Unlike the other treed models, the predictors are not used to specify what expert is used for a particular data point; rather mixing probabilities inferred from the data are used.

The Bayesian approach proposed here is substantially different from this past work. These earlier approaches are based on using some kind of greedy algorithm for tree construction, growing a large tree and then pruning it back. In contrast, the Bayesian approach induces a posterior distribution that can be used to guide a stochastic search towards 'more promising' treed models, thereby more fully exploring the model space. Such a Bayesian approach was successfully applied to conventional CART trees by Chipman, George, and McCulloch (1998) (hereafter denoted CGM) and Denison, Mallick, and Smith (1998). The Bayesian approach here elaborates this work—replacing the terminal node means or class probabilities in a conventional tree with more sophisticated models such as linear regression. The form of the terminal node models then becomes an integral part of the stochastic search, as the posterior is based on both the tree structure and terminal node models. The Bayesian approach also provides added flexibility for submodel specification. For example, the possibility of using different error variances for subset models, notably absent in previous methods, is easily accommodated.

A key aspect of any Bayesian approach is the introduction of prior distributions on all the unknowns, here the tree structures and terminal node models. In this paper we propose a class of priors and recommend specific "default" choices from within this class. We argue that, while no guarantees are made, these priors should lead to good results in a variety of problems and we illustrate this in the examples. Our prior plays two important roles in our methodology. First and foremost it provides a "penalty" to avoid overfitting in a natural, interpretable way. Secondly shrinkage of the bottom node parameters toward reasonable values stabilizes the estimation. Because treed model classes are so large, these aspects are critically important for obtaining realistic fits and predictions.

In Section 2, treed models are presented in a mathematical framework, and the important special case of treed regressions outlined and introduced via an example. Priors applicable to treed models of any form are discussed in Section 3.1. The search for interesting models is part of the Markov chain Monte Carlo method for estimating the posterior, discussed in Section 3.2. For treed regressions, the particular case considered here, prior parameters are selected in Section 4, in such a way that reasonable automatic choices of prior parameters can be made. In Section 5, two simulated and one real dataset are used to evaluate the predictive accuracy of treed regressions, and four competing models. We conclude in Section 6 with a discussion of directions for future investigations.

2. Treed models

2.1. The general model

Suppose we are interested in modeling the relationship between a variable of interest Y and a set of potential explanatory variables (a feature vector) x. A treed model is a specification of the conditional distribution of $Y \mid x$ consisting of two components—a binary tree T that partitions the x space, and a parametric model for Y that is associated with each subset of the partition.

The binary tree T is the conventional tree associated with CART (Breiman et al., 1984). Each interior node of the tree splits into two children, using a single predictor variable in the splitting rule. For ordered predictors, the rule corresponds to a cutpoint. For categorical predictors, the set of possible categories is partitioned into two sets. The tree T has b terminal nodes with each terminal node corresponding to a region in the space of potential x vectors. The b regions corresponding to the b terminal nodes are disjoint so that the tree partitions the x space by assigning each value of x to one of the b terminal nodes.

The treed model then associates a parametric model for Y at each of the terminal nodes of T. More precisely, for values x that are assigned to the ith terminal node of T, the conditional distribution of $Y \mid x$ is given by a parametric model $Y \mid x \sim f(y \mid x, \theta_i)$ with parameter θ_i . Letting $\Theta = (\theta_1, \ldots, \theta_b)$, a treed model is fully specified by the pair (Θ, T) . It may be of interest to note that early tree formulations such as CART were essentially proposed as data analysis tools rather than models. By elaborating the formulation as a parametric model, we set the stage for a Bayesian analysis. By treating the training sample (y, x) as realizations from a treed model, we can then compute the posterior distribution over (Θ, T) .

CGM and Denison, Mallick, and Smith (1998) discuss Bayesian approaches to the CART model. In those papers, the terminal node distributions of Y, $f(y|x,\theta)$, do not depend on x. For example, a Bayesian version of a conventional regression tree is obtained by letting $\theta = (\mu, \sigma)$ and $f(y|x,\theta) = N(\mu, \sigma^2)$. The conditional distribution of Y given x is then $Y|x \sim N(\mu_i, \sigma_i^2)$ when x is assigned to the terminal node associated with $\theta_i = (\mu_i, \sigma_i)$. Such models correspond to step functions for the expected value of Y|x, and may require large trees to approximate an underlying distribution Y|x whose mean is continuously changing in x. By using a richer structure at the terminal nodes, the treed models in a sense transfer structure from the tree to the terminal nodes. As a result, we expect that smaller,

and hence more interpretable, trees can be used to describe a wider variety of distributions for Y|x.

Finally, we should point out that although we use the one symbol "x" for notational simplicity, we may decide, a priori, to use one subset of the components of x in the rules defining the binary tree T and another subset in the fitting the models $p(y \mid x, \theta)$. These subsets need not be disjoint.

2.2. Treed regression: An example

One of the most familiar and useful parametric models is the linear model with normal error: $Y \mid x \sim N(x'\beta,\sigma^2)$. A treed regression is readily obtained by incorporating this model into our treed model framework. Simply let $\theta_i = (\beta_i,\sigma_i)$ and associate $Y \mid x \sim N(x'\beta_i,\sigma_i^2)$ with the terminal node corresponding to x. We also assume that given the x values, the Y values are independent of each other. Note that this treed regression allows both β_i and σ_i to vary from node to node. This allows us to flexibly fit both global non-linearity and global heteroskedasticity.

In subsequent sections, we describe in detail a Bayesian approach for finding and fitting treed regressions to data. In particular, in Section 5.3, we illustrate the performance of our approach on the well-known Boston Housing data, Harrison and Rubinfeld (1978). To give the reader an immediate sense of the potential of our approach, we describe here some of what we found.

The data consist of 14 characteristics of 506 census tracts in the Boston area. The response is the logged median value of owner occupied homes in each tract, and is listed in Table 1, along with the 13 predictor variables. One of the treed regressions that was selected by

Table 1.	Variables	in the	Boston	dataset.

Name	Description	Min	Max
CRIM	per capita crime rate by town	0.006	88.976
ZN	proportion of residential land zoned for lots over 25,000 sq.ft.	0.000	100.000
INDUS	proportion of non-retail business acres per town	0.460	27.740
CHAS	Charles River dummy variable (=1 if tract bounds river; 0 otherwise)	0.000	1.000
NOX	nitric oxides concentration (parts per 10 million)	0.385	0.871
RM	average number of rooms per dwelling	3.561	8.780
AGE	proportion of owner-occupied units built prior to 1940	2.900	100.000
DIS	weighted distances to five Boston employment centres	1.130	12.126
RAD	index of accessibility to radial highways	1.000	24.000
TAX	full-value property-tax rate per \$10,000	187.000	711.000
PTRATIO	pupil-teacher ratio by town	12.600	22.000
В	$1000(Bk - 0.63)^2$ where Bk is the proportion of blacks by town	0.320	396.900
LSTAT	% lower status of the population	1.730	37.970
MEDV	Log Median value of owner-occupied homes in \$1000's	1.609	3.912

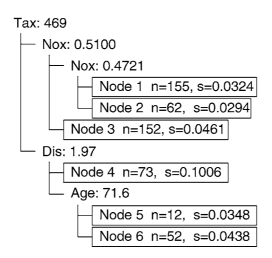


Figure 1. The tree structure of the treed regression model for the Boston data. Each node is represented by a line of text. Non-terminal nodes indicate the name of the splitting variable, and the cutpoint of the split. The first line after a nonterminal node is the child node with all values \leq the split value. In terminal nodes, the number of data points in the node (n), and the residual standard deviation (s) are given. Note that the units of split variables are on the original scale.

the Bayesian stochastic search is described by figures 1 and 2. Details of the Bayesian formulation such as prior choices that led to this model are discussed in Section 5.3.

Our fitted treed regression has 6 terminal nodes depicted in figure 1. One node that is immediately of interest is node 4, which has substantially larger residual standard deviation than the other nodes. This node consists of tracts with high tax rates (137 of the regions have taxes this high), and small distance to employment centres (105 of all tracts have this low a score, and 73 of the tracts with tax above 469). Among the high-tax, downtown areas, there appears to be more variation in the log median house price. Evidently, this is variation that is not readily explained by any predictors, and the treed regression search elects to leave the data alone, rather than overfit it.

The data at each of the 6 terminal nodes of our fitted treed regression is described by a separate regression with 14 estimated coefficients (including an intercept). As shown in figure 2, the coefficient of each predictor varies from terminal node to terminal node, reflecting varying sensitivity of the response to the predictors. Note that in fitting this treed regression, we first standardized all the predictors (see Section 4), so that all β_i can be compared on the same scale in figure 2. The coefficients in node 4 for NOX and DIS are very different from other nodes. Recall that this node contains the close to downtown, high tax areas. The large negative coefficients mean that these tracts are more sensitive to NOX and DIS than other tracts. Specifically, high pollution and larger distance from city centers will decrease the median value of a house much more than in other tracts. A possible explanation is that people may walk or take public transport because of their proximity to work, and so a relatively small increase in distance could affect value. Other interesting patterns include the coefficients for crime rate (CRIM). In nodes 1–3, which are low property tax areas, the effect of crime is zero or small. In the remaining three nodes,

TRATIO Tercept Tercept AGE BM AGE BAD TRATIO BE STATE TERTIO TERTIO

Estimated coefficients by terminal node (1-6)

Figure 2. Regression coefficient estimates of the treed model for the Boston data. Estimates for a regression in a specific node are linked by lines and labeled by node number. Coefficients for different variables may be plotted on the same scale because in the regression models coefficients are scaled equally.

where taxes are higher, the coefficients are negative, implying that increasing crime rates decrease value. Again, some tracts are more sensitive to this variable than others.

3. A Bayesian approach for treed modeling

In a nutshell, the Bayesian approach to model uncertainty problems proceeds as follows. The uncertainty surrounding the space of possible models is described by a prior distribution p(model). Based on the observed data, the posterior distribution on the model space is $p(\text{model} \mid \text{data}) \propto p(\text{data} \mid \text{model}) \ p(\text{model})$. Under the chosen prior, the best candidates for selection are then the high posterior probability models. Although simple to state, the effective implementation of this Bayesian approach can be delicate matter. The two crucial steps are the meaningful choice of prior distributions, and a method for identification of the high posterior models. We now describe methods for accomplishing each of these steps for general treed model classes.

3.1. Prior formulations for treed models

Since a treed model is identified by (Θ, T) , a Bayesian analysis of the problem proceeds by specifying a prior probability distribution $p(\Theta, T)$. This is most easily accomplished by

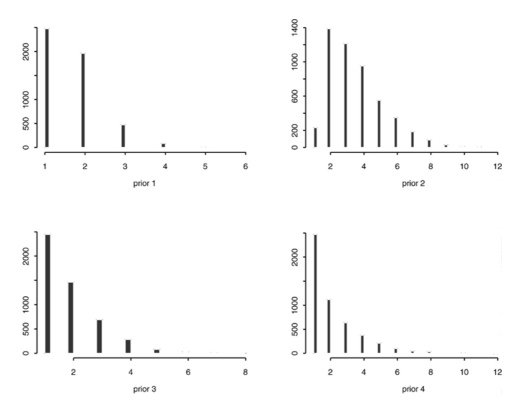


Figure 3. Marginal prior distribution for the number of terminal nodes, under various settings of α and β . Values for (α, β) are: upper left = (.5, 2), upper right = (.95, 1), lower left = (.5, 1), lower right = (.5, .5).

specifying a prior p(T) on the tree space and a conditional prior $p(\Theta \mid T)$ on the parameter space, and then combining them via $p(\Theta, T) = p(\Theta \mid T) p(T)$.

For p(T), we recommend the specification proposed in CGM for conventional trees. This prior is implicitly defined by a tree-generating stochastic process that "grows" trees from a single root tree by randomly "splitting" terminal nodes. A tree's propensity to grow under this process is controlled by a two-parameter node splitting probability $Pr(\text{node splits} \mid \text{depth} = d) = \alpha(1+d)^{-\beta}$, where the root node has depth 0. The parameter α is a "base" probability of growing a tree by splitting a current terminal node and β determines the rate at which the propensity to split diminishes as the tree gets larger. For details, see CGM. Figure 3 displays the marginal prior distribution on the number of terminal nodes (the tree size) for 4 different settings of (α, β) . (These figures are histograms of the tree sizes obtained by repeated simulation of trees from p(T)). Using such figures as a guide, values of (α, β) can be chosen to, for example, express the belief that reasonably small trees should yield adequate fits to the data. The tree prior p(T) is completed by specifying a prior on the splitting rules assigned to intermediate nodes. We use a prior that is uniform on available variables at a particular node, and within a given variable, uniform on all possible splits for that variable.

Turning to the specification of $p(\Theta \mid T)$, we note that while p(T) above is sufficiently general for all treed model problems, the specification of $p(\Theta \mid T)$ will necessarily be tailored to the particular form of the model $p(y|x,\theta)$ under consideration. This will be seen in Section 4 where we provide a detailed discussion of prior specification for the treed regression model. However, some aspects of $p(\Theta \mid T)$ specification should be generally considered. When reasonable, an assumption of independent and identically distributed (iid) components of Θ reduces the choice to that of a single density $p(\theta)$. However, even with this simplification, the specification of $p(\theta)$ can be both difficult and crucial. In particular, a key consideration is to avoid conflict between $p(\Theta \mid T)$ and the likelihood information from the data. On the one hand, if we make $p(\theta)$ too tight (i.e. with very small spread around the prior mean) the prior may be too informative and overwhelm the information in the data corresponding to a terminal node. On the other hand, if $p(\theta)$ is too diffuse (spread out), $p(\Theta \mid T)$ will be even more so, particularly for large values of b (large trees) given our iid model for the θ_i . Very spread out priors can "wash out" the likelihood in the sense of the Bartlett-Lindley paradox, Bartlett (1957). Thus, choice of an excessively diffuse prior will push the posterior of T towards concentrating on small trees. As will be seen in Section 4, both training data information and prior beliefs can be very useful for avoiding such pitfalls.

3.2. Markov chain Monte Carlo posterior exploration

Given a set of training data, the Bayesian analysis would ideally proceed by computing the entire posterior distribution of (Θ, T) . Unfortunately, in problems such as this, the size of the model space is so large that complete calculation of the posterior is infeasible. However, posterior information can still be obtained by using a combination of limited analytical simplification together with indirect simulation sampling from the posterior via Markov chain Monte Carlo (MCMC) methods. In particular, the Markov chain simulation can be used as a stochastic search algorithm which can be more comprehensive than traditional greedy grow/prune algorithms.

Our algorithm has the potential to be more comprehensive in three important ways. First, the stochastic steps help move the search past local maxima. Second, in addition to grow/prune steps we use "swap" and "change" steps which help find better trees. Finally, repeated restarting of the chain identifies a more diverse set of trees. These ideas were first developed in CGM, in the context of conventional trees. There, simulation studies illustrated that stochastic search, change/swap steps and multiple restarts found better trees than a greedy search. Chipman, George, and McCulloch (2001) also showed that this MCMC algorithm can find a wider variety of trees than a bootstrapped greedy grow/prune algorithm. Lutsko and Kuijpers (1994) considered a MCMC-like approach using simulated annealing, and also found improvements.

In its essentials, the method we use is the same as that in CGM. Our first step is to simplify the problem by integrating Θ out of the posterior so only T remains. Given T, we index the observed data so that (y_{ij}, x_{ij}) is the jth observation amongst the data such that x_{ij} corresponds to the ith node. Let n_i denote the number of observations corresponding to the

*i*th node and let (Y, X) denote all of the training data. Then,

$$p(Y \mid X, T) = \int p(Y \mid X, \Theta, T) p(\Theta \mid T) d\Theta$$

$$= \prod_{i=1}^{b} \int \prod_{j=1}^{n_i} p(y_{ij} \mid x_{ij}, \theta_i) p(\theta_i) d\theta_i$$
(1)

where the second equality follows from the assumed conditional independence of the y's given the x's and the modeling of the θ_i as iid given T. Note that in the first line we assume that X and Θ are independent given T, so that the prior on each θ_i does not depend on the x_{ij} values.

We use MCMC to stochastically search for high posterior trees T by using the following Metropolis-Hastings algorithm which simulates a Markov chain T^0, T^1, T^2, \ldots with limiting distribution $p(T \mid Y, X)$. Starting with an initial tree T^0 , iteratively simulate the transitions from T^i to T^{i+1} by the two steps:

- 1. Generate a candidate value T^* with probability distribution $q(T^i, T^*)$.
- 2. Set $T^{i+1} = T^*$ with probability

$$\alpha(T^{i}, T^{*}) = \min \left\{ \frac{q(T^{*}, T^{i})}{q(T^{i}, T^{*})} \frac{p(Y \mid X, T^{*})p(T^{*})}{p(Y \mid X, T^{i})p(T^{i})}, 1 \right\}.$$
 (2)

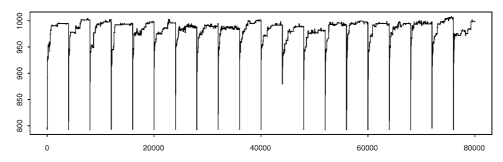
Otherwise, set $T^{i+1} = T^i$.

In (2), p(Y | X, T) is obtained from (1), and $q(T, T^*)$ is the kernel which generates T^* from T by randomly choosing among four steps:

- GROW: Randomly pick a terminal node. Split it into two new ones by randomly assigning it a splitting rule from the prior.
- PRUNE: Randomly pick a parent of two terminal nodes and turn it into a terminal node by collapsing the nodes below it.
- CHANGE: Randomly pick an internal node, and randomly reassign it a splitting rule from the prior.
- SWAP: Randomly pick a parent-child pair which are both internal nodes. Swap their
 splitting rules unless the other child has the identical rule. In that case, swap the splitting
 rule of the parent with that of both children.

CGM note that chains simulated by this algorithm tend to quickly gravitate towards a region where $P(T \mid Y, X)$ is large, and then stabilize, moving locally in that region for a long time. Evidently, this is a consequence of a proposal distribution which makes local moves over a sharply peaked multimodal posterior. Some of these modes will be better than others, and it is desirable to visit more than one. To avoid wasting time waiting for mode-to-mode moves, CGM recommend search with multiple restarts of the algorithm, saving the most promising trees from each run.

20 restarts of 4000 iterations



1 start of 80,000 iterations

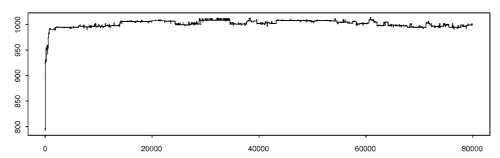


Figure 4. Comparison of log likelihood for 20 restarts of 4000 steps per start and 1 start chain with 80,000 steps.

By fitting parametric models relating y to x in each terminal node we hope to obtain good fit with much smaller trees than in the model considered by CGM. With smaller trees the MCMC method is better able to explore the space so that the approach is much more computationally stable. In fact, we have found that fewer restarts and iterations are typically needed here to find high posterior probability trees than in CGM. The decision of how many restarts and runs per start will vary according to sample size, signal to noise ratio, and complexity of the data.

Because MCMC performance is apt to vary from problem to problem, our strategy has been to use the data to choose the number of restarts and iterations informally. We first run some initial exploratory chains to get a rough idea of how many iterations it takes the chain to get to the point where the likelihood has stabilized and only small local moves are being made. We then repeatedly run the chain, restarting it after this number of iterations. Figure 4 illustrates this process for the data considered in detail in Section 5.3. The top panel displays the log integrated likelihood (logged values of $p(Y \mid X, T)$) for the sequence of trees visited by 20 restarts of 4000 iterations. We see that within each restarted sequence, the values typically increase dramatically at first and then start to level off, suggesting small local moves. Notice that not all sequences plateau at the same level, indicating that the algorithm may not be mixing well, and may be getting stuck in local maxima. The lower panel shows

one long run of 80,000 iterations. While additional iterations do find larger values, the bulk of the gain occurs early in the run. In this example either approach (20 short runs or one long run) would likely yield comparable results. As discussed in CGM and Denison, Mallick, and Smith (1998), other strategies may be considered, and we are currently engaged in research directed at this issue.

4. Specification of $p(\Theta \mid T)$ for treed regression

In this section we discuss particular parameter prior $p(\Theta \mid T)$ specifications for the treed regression model described in Section 2.2. Coupled with the tree prior p(T) recommended in Section 3.1, such $p(\Theta \mid T)$ completes the prior specification for treed regression. Alternative approaches to $p(\Theta \mid T)$ specification that may also warrant consideration are briefly discussed in Section 6.

We begin with the simplifying assumption of independent components $\theta_i = (\beta_i, \sigma_i)$ of Θ , reducing the specification problem to a choice of $p(\beta, \sigma)$ for every component (i.e. terminal node). For this, we propose choosing the conjugate form $p(\beta, \sigma) = p(\beta \mid \sigma)p(\sigma)$ with:

$$\beta \mid \sigma \sim N(\bar{\beta}, \sigma^2 A^{-1}), \quad \sigma^2 \sim \frac{\nu \lambda}{\chi_{\nu}^2}$$
 (3)

so that the integral (1) of Section 3.2 can be performed analytically (see, for example, Zellner, 1971). This further reduces the specification problem to the choice of values for the hyperparameters ν , λ , $\bar{\beta}$ and A.

Before going into the specification details for these hyperparameters, we note in passing that methods which flexibly fit data typically have "parameters" that must be chosen. For example, in fitting a neural net one must choose a decay parameter. Often these parameters are related to penalties for model size to avoid overfitting. Models that are too large will have good training sample fit but poor test sample predictive performance. To some extent, both the choice of our prior on T (indexed by the two scalar parameters α and β , see Section 3.1) and the choice of the conjugate prior are related to the issue of penalizing overly large models. Our goal is to choose a single (or at least small) set of possible hyperparameter values such that the resulting choice of model works reasonably well in a variety of situations.

As noted in Section 3.1, the choice of this prior is a crucial part of the method. What is needed is a reasonably automatic way to specify a prior that is neither too tight nor too spread out. Furthermore, if the results are very sensitive to this choice then the method will not be of practical use. As will be seen below, our approach to hyperparameter selection makes use of a rough combination of training data information and prior beliefs, and is similar in spirit to the prior selection approach of Raftery, Madigan, and Hoeting (1997).

We begin by standardizing the training data by a linear transformation so that each x and y has an average value of 0 and a range of 1. The purpose of this standardization is to make it easier to intuitively gauge parameter values. For example, with the standardized data, 2.0 would be a very large regression coefficient because a half range increase in the corresponding predictor would result in a full range increase in y.

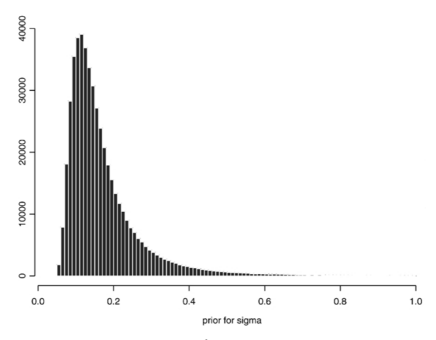


Figure 5. Prior distribution for residual variance σ^2 , Boston housing data.

4.1. Choosing v and λ

We choose ν to be 3. A common interpretation of this value is that this gives our prior information about σ the same weight as if it were based upon 3 observations, making this a reasonably uninformative prior. To choose λ we turn to the data. Let s be the classical unbiased estimate of σ based on a full linear regression fit to all of the standardized training data. We want to choose λ to express the idea that for terminal node regressions we expect the error standard deviation to be smaller than s, but maybe not too much smaller. Since we allow for a different σ value in each terminal node we also want to capture the prior belief that some σ values could be bigger than s. We thus choose a quantile q such that $Pr(\sigma < s) = q$, and then use the implied value of λ , $\lambda = s^2 \Phi_{\nu}^{-1} (1-q)/\nu$ where Φ_{ν} is the cumulative distribution function for the chi-squared distribution with ν degrees of freedom. In practice, we have tried the values q = .75 and q = .95 which correspond to $\lambda = .404s^2$ and $\lambda = .1173s^2$, respectively.

For the Boston housing data discussed in Sections 2.2 and 5.3, s = .2. Figure 5 displays draws from the prior distribution of σ^2 when q = .75, one of the priors considered in our analysis.

4.2. Choosing $\bar{\beta}$ and A

In the absence of prior information, we set $\bar{\beta} = 0$, the neutral choice indicating indifference between positive and negative values of the components of β . A prior mean of zero on the

intercept is especially plausible because of our standardization of predictors and response to have mean 0. Of course, given T, the average value of a variable using only those observations at a particular terminal node may differ from 0. To accommodate this possibility it is important to make the spread of the prior (defined by A) broad enough.

To choose A, we first make the simplifying restriction that A=aI, where I is the identity matrix. Making A diagonal is a simplifying assumption, though the use of common diagonal element a seems at least partly justified by the standardization. We choose a, the remaining unknown, by choosing c such that $Pr(-c < \beta < c) = .95$, and then using the relationship $a = \lambda 3.18^2/c^2$ (each component of β is marginally $\sim t_\nu \sqrt{\lambda/a}$, where t_ν is the t distribution with ν degrees of freedom). The constant 3.18 corresponds to the 95% cutoff for a t distribution with $\nu = 3$ degrees of freedom.

How do we choose c? Given the standardization we can argue that ± 1 is a reasonable range for a coefficient since a value of 1 indicates that all the variation in y could be explained by a single x. However, such reasoning is clearly flawed in at least two ways. First of all, given a tree T the observations corresponding to that node are no longer standardized (although they would have a range less than 1, and a mean between -1 and 1). Indeed, the whole point of the model is that the coefficients may vary substantially from node to node. Secondly, the presence of multicollinearity throws all such reasoning out the window. Even with standardized data, the MLE of a coefficient could be arbitrarily large. However, if severe multicollinearity is present we can usually shrink coefficients substantially (in this case towards 0) without appreciable damage to the fit. In fact, such shrinkage often stabilizes calculations and even improves predictions. Given these considerations and the realistic goal of hoping to find something that is "in the right ballpark" for a variety of problems, we have tried values such as c=1 and c=3. Generally, smaller c values will result in estimated coefficients that are shrunk towards 0, and trees with fewer terminal nodes.

Finally, the intercept perhaps warrants special consideration. If all the slopes are 0, then clearly a range of ± 1 is appropriate for the intercept. If the slopes are not 0, then the intercept could be anything (just as the slopes could be anything in the presence of multicollinearity). In the examples in the next section, we have used the same c values for the intercept as for the slopes, but it may be that larger c values would have been more appropriate.

5. Three examples

In this section, we illustrate Bayesian treed regression and evaluate its performance on two simulated datasets, and a single real dataset, the Boston housing data. The first simulated example was chosen because we expected to do very well on it and the second simulated example was chosen because we expected to do poorly.

Comparisons are made with the following other methods:

- 1. The linear regression model, using all available predictors.
- 2. Conventional tree models in which the prediction is a constant in each terminal node. Clark and Pregibon's (1992) implementation in S (Becker, Chambers, & Wilks, 1988), with greedy grow/prune steps similar to CART was used.

- 3. Multivariate Adaptive Regression Splines (MARS) (Friedman, 1991). The S implementation described in Kooperberg, Bose, and Stone (1997) was used.
- 4. Neural networks, with a single layer of hidden units, and a skip layer. The S implementation (Venables & Ripley, 1999) was used. The skip layer allows a directly connection from inputs to output without an activation function. For the hidden units, a logistic activation function was used, while the output unit was linear. Training was via a quasi-Newton algorithm.

Linear models and trees were chosen since they are the building blocks of our method. The piecewise linear and locally adaptive functional form of MARS makes it a good competitor. Neural nets are included as a black-box nonparametric method. Methods for combing models, such as bagging, boosting, or Bayesian model averaging, were not considered. This decision is discussed in Section 6.

For each of the three data sets we used the same set of 6 different prior settings. For the tree prior, we used $\alpha = .5$ and $\beta = 2$ in all examples. This is a very conservative choice in that it puts a lot of weight on small trees (see figure 3). For λ we tried both of the choices discussed in Section 4.1: $\lambda = .404s^2$ for q = .75, and $\lambda = .1173s^2$ for q = .95. For the coefficients we tried both of the choices discussed in Section 4.2: c = 1 and c = 3 and we also tried c = 10. This gives $2 \times 3 = 6$ possible priors. Based on our heuristics in Section 4.2, only the choice c = 10 is considered potentially unreasonable. The other four priors are chosen in an attempt to cover a wide range of reasonable beliefs. Our hope is that the results are not overly sensitive to the prior choice.

Other choices in addition to the prior must be made to use Bayesian treed models. As discussed in Section 3.2, each Markov chain was repeatedly restarted after a predetermined number of iterations. This number differed from example to example as discussed in detail therein. Note that where we restarted the chain many times, good trees were often found in the first few restarts of the chain. Longer runs were simply used to see just how good a tree could be found.

To actually make a prediction we must choose a specific tree from all of those visited by the chain. The single tree with the highest integrated likelihood P(T | Y, X) was chosen. Given the tree T, the posterior expected value of the parameters in each node were used as estimates of β and σ .

In every example, all the components of x were used in both the tree and the bottom node regressions.

5.1. First simulated example

Data with two predictors and 200 observations were generated according to the model

$$\mu = \begin{cases} 1 + 2x_1 & \text{if } x_2 \ge 0.5\\ 0 & \text{if } x_2 < 0.5 \end{cases} \tag{4}$$

where the response has expected value μ , x_1 , x_2 are independent uniforms on (0,1), and independent normal errors with mean 0 and standard deviation 0.10 were added to each

observation. We expect treed regression to do well with this data, since the functional form of the model matches the data generation mechanism. 100 simulated datasets were generated according to this model, and each model trained on the same datasets.

For the other methods, several algorithm parameters have to be set. To explore the effect of the parameters, we identify a range of good values, and then run the procedure with several values in this range. For example, with trees, cross validation suggests that trees with 5 or more nodes will fit about equally well. Consequently, we considered 10, 20, 30, 40, and 50 terminal nodes. For MARS, the generalized cross-validation penalty term (gcv) has a recommended value of 4, and values 2–6 were tried. For neural nets, 3, 5, 8, 10, and 15 hidden units plus a skip layer were used. Weight decay values of 10^{-2} , 10^{-3} , 10^{-4} , 10^{-5} , as suggested by Venables and Ripley (1999) were considered. This gives $5 \times 4 = 20$ different combinations of neural networks.

For the Bayesian methods, we found that after 2000 to 3000 iterations, the chain stabilized yielding subsequent trees with similar integrated likelihoods. We thus ran the chain with 10 restarts of 5000 iterations for each of the 6 prior settings (although far fewer iterations would have given similar results). Each run of 50,000 total iterations took about 20 seconds (Pentium III, 500 MHz).

Model accuracy was evaluated via the root mean squared error,

$$RMSE = \sqrt{\sum_{i=1}^{n} \frac{(\mu_i - \hat{Y}_i)^2}{n}},$$

where μ_i is the expected value of observation Y_i , given by (4). An RMSE value for each method was obtained for each of the 100 repetitions. These are plotted in figure 6, using boxplots to represent the 100 values for each method. Some of the 20 neural nets are omitted from the plot, to improve clarity. None of the neural nets omitted are among the best models found. Our method does best, with an average RMSE ranging from 0.016 to 0.018. A 10 node tree and a neural net with 15 nodes and decay of 10^{-5} are the closest competitors, with respective mean RMSE values of 0.057 and 0.065. Although the boxplots ignore the matched nature of this experiment (i.e. each method is applied to the same dataset), it is clear that our procedure wins hands down, in part due to the discontinuous nature of the data. Note that the performance of the Bayesian procedure is insensitive to the choice of prior.

5.2. Second simulated example

Friedman (1991) used this simulated example to illustrate MARS. n = 100 observations are generated with 10 predictors, each of which is uniform on (0,1). The expected value of Y is

$$\mu = 10\sin(\pi x_1 x_2) + 20(x_3 - 0.5)^2 + 10x_4 + 5x_5 \tag{5}$$

with random noise added from a standard normal distribution. The remaining five predictors have no relationship with the response.

This example is considerably more difficult, and the sine term should prove especially challenging to our method, as there is an abrupt changepoint with a saddlepoint at 45 degrees

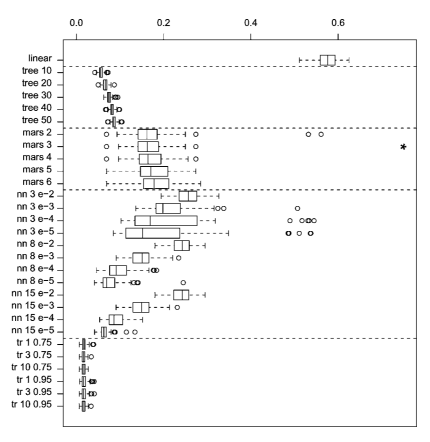


Figure 6. Boxplots representing RMSE for various models and parameter settings, first simulation example. Trees are identified as having 10–50 nodes, MARS by the GCV penalty parameter (2–6), neural nets by the number of hidden units (3, 8, 15) and weight decay (1e-2, 1e-3,1e-4,1e-50), and Bayesian regression tree by the prior on λ (75% or 90% tail) and the prior variance on c (1, 3, 10). The * represents a single MARS model with an outlying RSS of 5.91 (truncated to fit in the plotting region).

to either axis. Since the splits for the tree must be aligned to the axes, this will be difficult to identify. The signal to noise ratio is high, however.

In this example, preliminary runs indicated that even after a large number of iterations the chain would still make substantial moves in the tree space. Given reasonable time limitations, we thus opted for one long run of 500,000 iterations for each of the 6 prior settings. Each run of 500,000 took about 12.2 minutes.

The RMSE values for 50 simulations are given in figure 7.

In this case neural networks are the most consistent performer, although MARS occasionally gives better fits. Our tree procedure fares surprisingly well, considering the small number of terminal nodes (in the range 2–6 for various different priors). The fact that greedy trees improve their fit as the number of terminal nodes is increased suggests that choosing priors which force further growing of the treed regression might further help prediction. This was not explored because of our stated goal of evaluating default choices of priors.

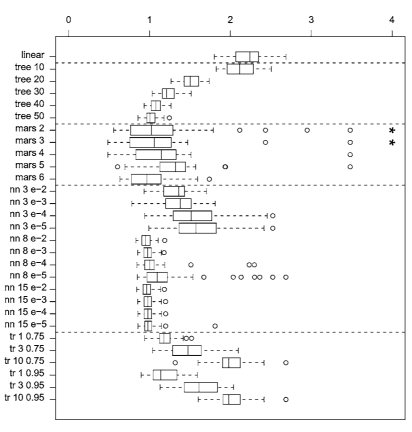


Figure 7. Boxplots representing RMSE for various models and parameter settings, second simulation example. Labeling is the same as in figure 6.

In this case, the choice of prior does seem to affect the performance of treed regression. However, the choice c = 10, which was a priori deemed to be extreme, here fares the worst.

Several methods appear to perform comparably, from the boxplot. A more accurate comparison can be made by exploiting the fact that the models were all trained on the same datasets. By calculating the differences in RMSE for each dataset, a more accurate comparison can be made. The mean values of RMSE over the 50 simulated datasets are 1.192, 1.018, 0.983, 0.987 for treed regression, trees, neural networks and MARS, respectively. Based on the standard errors of paired differences, all differences seem significant except those between MARS and neural networks or trees (smallest t-statistic for any paired comparison is 5.17).

5.3. Boston housing results

In this section we report RMSE values for our procedure and the four other methods, as in the simulation cases. The data consists of 506 observations, and RMSE performance is

now assessed using 10-fold cross validation. That is, the data are randomly divided into 10 roughly equal parts, each model is trained on 9/10 of the data, and tested on the remaining 1/10. This is repeated 10 times, testing on a different tenth each time. Unlike the simulated examples, the true function is unknown, meaning that μ_i in RMSE is replaced by observed y_i in the 1/10 test portion of the data. To reduce variation due to any particular division of the data, the entire experiment is repeated 20 times. The 20 repetitions play a similar role to the 50 or 100 simulations in earlier examples. Analogously, we will look at boxplots of RMSE to get a sense of performance and variation in the performance over different "datasets". Note that cross-validation is used only to assess predictive error. The methods for hyperparameter selection described in Section 4 are still used.

Figure 4 at the end of Section 3.2 displays runs of our chain based on all 506 observations. For each of the six prior setting we ran 20 restarts of 4000 iterations. As discussed in Section 3.2 one long run might have done just as well. Each run of 80,000 iterations took about 13.5 minutes. The tree reported in Section 2.2 used one of these six settings, namely the 75th percentile of σ , and c = 3.

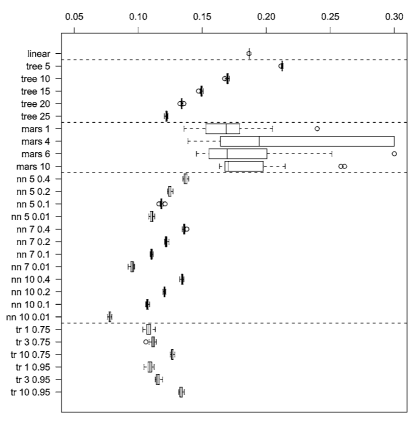


Figure 8. RMSE for training data, Boston housing example.

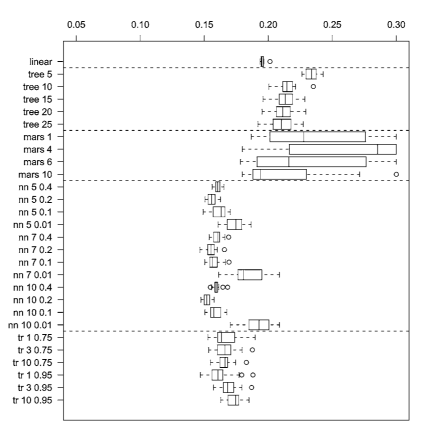


Figure 9. RMSE for test data, Boston housing example.

To reflect the higher signal to noise ratio, conventional trees of size 5, 10, 15, 20, and 25 were used. We were unsure of how to choose hyperparameters for MARS and neural nets. We tried a variety of choices and report results for a subset of choices which perform well in terms of test RMSE. For MARS, we report results for GCV values of 1, 4, 6, and 10. For neural nets, we report results for 5, 7, and 10 hidden units and decay parameter values of 0.4, 0.2, 0.1, and 0.01.

Figures 8 and 9 give boxplots of training and test error of the various methods. Only neural nets and treed regression outperform a linear regression. Their performance is comparable, although neural nets have a slight edge.

The success of our procedure in this example may be because treed regressions allow the variances to be different across terminal nodes. This may seem counterintuitive, since one might expect that if the variance in a terminal node is larger, the predictions will suffer. This may not be the case. If there really is greater noise in some regions of the predictor space, a method that assumes constant variance may overfit. In the test sample, such methods will underperform, while a procedure that leaves the unexplainable variation alone may do better.

6. Discussion

The main thrust of this paper has been to propose a Bayesian approach to treed modeling and to illustrate some of its features. However, we have only explored part of its potential here and there are several important avenues which warrant further investigation in the future.

Because of the increasing prevalence of large data sets with many variables, it will be very important to consider the performance of our approach on such data sets. Two key issues in this area will be the algorithm's speed and the extent to which it mixes (i.e. avoids getting stuck in local maxima). A promising feature of our approach is its ability to find good models quickly, as is typical of many heuristic procedures. Nonetheless, it will be of interest to develop fast alternatives to stochastic search for exploring the posterior. As suggested by one of the referees, a Bayesian greedy algorithm guided by the posterior may be very effective, either as a starting point for stochastic search, or as an end in itself. We also plan to explore methods for improving the mixing of the chain.

Our proposed approach to hyperparameter specifications in Section 4 entails explicit user specified choices based on basic data features such as the scale of the predictors and noise level. Hopefully such specifications can be reasonably automatic with only weak subjective input. The robustness of our procedures as the prior inputs varied in the examples of Section 5, provides some evidence that our recommendations are reasonable. However, this is another crucial issue requiring further study. Although much more time consuming, it may be better to use hyperparameter selection based on the data using methods such as cross validation, MCMC sampling (Neal, 1996) or empirical Bayes (George & Foster, 2000). At the very least, a comparison with such methods is needed.

Although we have focused on Bayesian model selection, our Bayesian platform can easily be used for Bayesian model averaging. Such averaging could be obtained by posterior or likelihood weighting of the MCMC output. One might also consider averaging only a subset of best models. Although such treed model averages forego the interpretability features of single treed models, they will probably provide improved predictions similar to those in Raftery, Madigan, and Hoeting (1997). It will be of interest to investigate the extent to which such averaging offers improved prediction. We suspect this will hinge on the stability of treed models (Breiman, 1996). The inclusion of more complicated models in terminal nodes may mean less instability in treed regressions, due to smaller tree sizes. However, treed regressions may also benefit from model averaging in situations like our Section 5.2 example, where the functional form is not readily represented by a small to moderate number of partitioned regression models.

Following CGM, we based model selection here on the integrated log likelihood. In fact, the posterior distribution of trees may contain a variety of interesting and distinct trees. Methods for making some sense of this "forest" of trees may be advantageous, offering the possibility of selecting several interesting trees for either interpretation or model averaging. The extension of the ideas in Chipman, George, and McCulloch (2001) should prove fruitful in this regard.

The Bayesian approach to treed modeling provides a coherent method for combining prior information with the training data to obtain a posterior distribution that guides a stochastic search towards promising models. The internal consistency of the Bayesian framework

seems to naturally guard against overfitting with the very rich class of treed models. The within model shrinkage mentioned in Section 4.2 provides some stability in this regard. However, it should be possible to obtain further improvements using shrinkage across models. This would allow parameter estimates that do not change substantially across terminal nodes to be shrunk towards a common value, or set to 0, if variable selection were of interest. Leblanc and Tibshirani (1998), Hastie and Pregibon (1990), and Chipman, George, and McCulloch (2000) illustrate the promise of shrinkage in conventional trees models. We are hopeful that these methods will extend naturally to the current framework.

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