An Introduction to Recursive Partitioning: Rationale, Application, and Characteristics of Classification and Regression Trees, Bagging, and Random Forests

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Recursive partitioning methods have become popular and widely used tools for nonparametric regression and classification in many scientific fields. Especially random forests, which can deal with large numbers of predictor variables even in the presence of complex interactions, have been applied successfully in genetics, clinical medicine, and bioinformatics within the past few years. High-dimensional problems are common not only in genetics, but also in some areas of psychological research, where only a few subjects can be measured because of time or cost constraints, yet a large amount of data is generated for each subject. Random forests have been shown to achieve a high prediction accuracy in such applications and to provide descriptive variable importance measures reflecting the impact of each variable in both main effects and interactions. The aim of this work is to introduce the principles of the standard recursive partitioning methods as well as recent methodological improvements, to illustrate their usage for low and high-dimensional data exploration, but also to point out limitations of the methods and potential pitfalls in their practical application. Application of the methods is illustrated with freely available implementations in the R system for statistical computing.

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Prediction, classification, and the assessment of variable importance are fundamental tasks in psychological research. A wide range of classical statistical methods—including linear and logistic regression as the most popular representatives of standard parametric models—is available to address these tasks. However, in certain situations, these classical methods can be subject to severe limitations.

One situation where parametric approaches are no longer applicable is the so-called small $n$ large $p$ case, where the number of predictor variables, $p$, is greater than the number of subjects, $n$. This case is common, for example, in genetics, where thousands of genes are considered as potential predictors of a disease. However, even in studies with much lower numbers of predictor variables, the combination of all main and interaction effects of interest—especially in the case of categorical predictor variables—may well lead to cell counts too sparse for reliable parameter estimation. Thus, interaction effects of high order usually cannot be included in standard parametric models.

Additional limitations of many standard approaches include the restricted functional form of the association pattern (with the linear model as the most common and most restrictive case), the fact that ordinally scaled variables, which are particularly common in psychological applications, are often treated as if they were measured on an interval or ratio scale, and the fact that measures of variable importance are available only for a small range of methods.
The aim of this article is to provide an instructive review of a set of statistical methods adopted from machine learning that overcome these limitations. The most important of these methods is the random forest approach of Breiman (2001a): A random forest is a so-called ensemble (or set) of classification or regression trees (CART; Breiman, Friedman, Olshen, & Stone, 1984). Each tree in the ensemble is built on the basis of the principle of recursive partitioning, where the feature space is recursively split into regions containing observations with similar response values. A detailed explanation of recursive partitioning is given in the next section.

In the past years, recursive partitioning methods have gained popularity as a means of multivariate data exploration in various scientific fields, including, for example, the analysis of microarray data, DNA sequencing, and many other applications in genetics, epidemiology, and medicine (cf., e.g., Bureau et al., 2005; Diaz-Uriarte & Alvarez de Andrés, 2006; Gunther, Stone, Gerwien, Bento, & Heyes, 2003; Huang et al., 2005; Lünetta, Hayward, Segal, & Edersewgh, 2004; Qi, Bar-Joseph, & Klein-Seetharaman, 2006; Segal, Barbour, & Grant, 2004; Shih, Seligson, Belldegrun, Palotie, & Horvath, 2005; Ward, Pajevic, Dreyfuss, & Malley, 2006).

A growing number of applications of random forests in psychology indicates a wide range of application areas in this field, as well: For example, Oh, Laubach, and Luczak (2003) and Shen, Ong, Li, Hui, and Wilder-Smit (2007) applied random forests to neuronal ensemble recordings and EEG data, which are too high dimensional for the application of standard regression methods. An alternative approach to cope with large numbers of predictor variables would be to first apply dimension reduction techniques, such as principle components or factor analysis, and then use standard regression methods on the reduced data set. However, this approach has the disadvantage that the original input variables are projected into a reduced set of components, so that their individual effect is no longer identifiable. As opposed to that, random forests can process large numbers of predictor variables simultaneously and provide individual measures of variable importance.

Interesting applications of random forests in data sets of lower dimensionality include the studies of Rossi, Amaddeo, Sandri, and Tansella (2005) on determinants of once-only contact in community mental health service and Baca-Garcia et al. (2007) on attempted suicide under consideration of the family history. For detecting relevant predictor variables, Rossi et al. (2005) pointed out that the random forest variable importance ranking proves to be more stable than stepwise variable selection approaches available for logistic regression, which are known to be affected by order effects (see, e.g., Austin & Tu, 2004; Derksen & Keselman, 1992; Freedman, 1983). Moreover, a high random forest variable importance of a variable that was not included in stepwise regression may indicate that the variable works in interactions that are too complex to be captured by the parametric regression model. As another advantage, Marinic et al. (2007) pointed out, in an application to the diagnosis of posttraumatic stress disorder, that random forests can be used to automatically generate realistic estimates of the prediction accuracy on test data by means of repeated random sampling from the learning data.

Luellen, Shadish, and Clark (2005) explored another field of application in comparing the effects in an experimental and a quasi-experimental study on mathematics and vocabulary performance: When the treatment choice in the quasi-experimental study is chosen as a working response, classification trees and ensemble methods can be used to estimate propensity scores (i.e., treatment probabilities). However, some of these seminal applications of recursive partitioning methods in psychology also reveal common misperceptions and pitfalls. For example, Luellen et al. (2005) suspected that ensemble methods could overfit (i.e., adapt too closely to random variations in the learning sample, as discussed in detail later) when too many trees are used to build the ensemble—even though this has been shown not to be the case—whereas recent results have indicated that other tuning parameters may be responsible for overfitting in random forests.

More common mistakes in the practical use and interpretation of recursive partitioning approaches are the confusion of main effects and interactions (see, e.g., Berk, 2006) as well as the application of biased variable selection criteria and a significance test for variable importance measures (see, e.g., Baca-Garcia et al., 2007) that has recently been shown to have extremely poor statistical properties. Some of these pitfalls are promoted by the fact that random forests were not developed in a stringent statistical framework, so their properties are less predictable than those of standard parametric methods, and some parts of random forests are still under construction (cf. also Polikar, 2006, for a brief history of ensemble methods, including fuzzy and Bayesian approaches). Therefore, the aim of this article is not only to point out the potential of random forests and related recursive partitioning methods to a broad scientific community in psychology and related fields, but also to provide a thorough understanding of how these methods function, how they can be applied practically, and when they should be handled with caution.

The next section describes the rationale of recursive partitioning methods, starting with single classification and regression trees and moving on to ensembles of trees. Examples are interspersed between the technical explanations and provided in an extra section to highlight potential areas of application. A synthesis of important features and advantages of recursive partitioning methods—as well as important pitfalls—with an emphasis on random forests is given in a later section. For all examples shown here, freely
available implementations in the R system for statistical computing (R Development Core Team, 2009) were used. The corresponding code is provided and documented in an online supplement as an aid for new users.

Methods

After the early seminal work on automated interaction detection by Morgan and Sonquist (1963), the two most popular algorithms for classification and regression trees (abbreviated as classification trees in most of the following), CART and C4.5, were introduced by Breiman et al. (1984) and independently by Quinlan (1986, 1993). Their nonparametric approach and the straightforward interpretability of the results have added much to the popularity of classification trees (cf., e.g., Hann¨over, Richard, Hansen, Martinovich, & Kordy, 2002; Kitsantas, Moore, & Sly, 2007, for applications on the treatment effect in patients with eating disorders and determinants of adolescent smoking habits). As an advancement of single classification trees, random forests (Breiman, 2001a), as well as its predecessor method, bagging (Breiman, 1996a, 1998), are termed ensemble methods, because an ensemble or committee of classification trees is aggregated for prediction. This section introduces the main concepts of classification trees, which are then used as so-called base learners in the ensemble methods bagging and random forests.

How Do Classification and Regression Trees Work?

Classification and regression trees are a simple nonparametric regression approach. Their main characteristic is that the feature space (i.e., the space spanned by all predictor variables) is recursively partitioned into a set of rectangular areas, as illustrated later. The partition is created such that observations with similar response values are grouped. After the partition is completed, a constant value of the response variable is predicted within each area.

The rationale of classification trees is explained in more detail by means of a simple psychological example: Inspired by the study of Kitsantas et al. (2007) on determinants of adolescent smoking habits, an artificial data set was generated for illustrating variable and split selection in recursive partitioning. Our aim is to predict the adolescents’ intention to smoke a cigarette within the next year (binary response variable intention_to_smoke) from four candidate risk factors (the binary predictor variables lied_to_parents, indicating whether the subject has ever lied to the parents about doing something they would not approve of, and friends_smoke, indicating peer smoking of one or more among the four best friends, as well as the numeric predictor variables age, indicating the age in years, and alcohol_per_month, indicating how many times the subject drank alcohol in the past month).

The data were generated to resemble the key results of Kitsantas et al. (2007). However, the variables age and alcohol_per_month, which were used only in a discretized form by Kitsantas et al. (2007), were generated as numeric variables to illustrate the selection of optimal cutpoints in recursive partitioning. The generated data set, as well as the R code used for all examples, are available as online supplements.

The classification tree derived from the smoking data is illustrated in Figure 1A and shows the following: From the entire sample of 200 adolescents, a group of 89 adolescents is separated from the rest in the first split. This group (represented by Node 2, where the node numbers are more labels assigned sequentially from left to right starting from the top node) is characterized by the fact that none of their four best friends smoked and that within this group only a few subjects intended to smoke within the next year. The remaining 111 subjects are further split into two groups (Nodes 4 and 5) according to whether they drank alcohol (a) on fewer than one or one or (b) on more than one occasion in the past month. These two groups again vary in the percentage of subjects who intended to smoke.

The model can be displayed either as a tree, as in Figure 1A, or as a rectangular partition of the feature space, as in Figure 1B: The first split in the variable friends_smoke partitions the entire sample, whereas the second split in the variable alcohol_per_month further partitions only those subjects whose value for the variable friends_smoke is one or more. The partition representation in Figure 1B is even better suited than the tree representation to illustrating that recursive partitioning creates nested rectangular prediction areas corresponding to the terminal nodes of the classification tree. Details about the prediction rules derived from the partition are given later.

Note that the resulting partition is one of the main differences between classification trees and, for example, linear regression models: Whereas in linear regression the information from different predictor variables is combined linearly, here the range of possible combinations includes all rectangular partitions that can be derived by means of recursive splitting—including multiple splits in the same variable. In particular, this includes nonlinear and even nonmonotone association rules, which do not need to be specified in advance but are determined in a data driven way.

Of course, there is a strong parallel between tree building and stepwise regression, where predictors are also included one at a time in successive order. However, in stepwise linear regression, the predictors still have a linear effect on the dependent variable, whereas extensions of stepwise procedures, including interaction effects, are typically limited to the inclusion of twofold interactions, because the number of higher order interactions—that would have to be considered simultaneously when starting the selection proce-
dure—is too large. In contrast to this, in recursive partitioning, only those interactions that are actually used in the tree are generated during the fitting process. The issue of including main effects and interactions in recursive partitioning is discussed in more detail later.

Splitting and stopping. Both the CART algorithm of Breiman et al. (1984) and the C4.5 algorithm (and its predecessor ID3) of Quinlan (1986, 1993) conduct binary splits in numeric predictor variables, as depicted in Figure 1. In categorical predictor variables (of nominal or ordinal scale of measurement), C4.5 produces as many nodes as there are categories (often referred to as k-ary or multiway splitting), whereas CART again creates binary splits between the ordered or unordered categories. We concentrate on binary splitting trees in the following and refer to Quinlan (1993) for k-ary splitting.

For selecting the splitting variable and cutpoint, both CART and C4.5 follow the approach of impurity reduction, which we illustrate by means of our smoking data example: In Figure 2, the relative frequencies of both response classes are displayed not only for the terminal nodes, but also for the inner nodes of the tree previously presented in Figure 1. Starting from the root node at the top, we find that the relative frequency of yes answers in the entire sample of 200 adolescents is about 40%. By means of the first split, the group of 89 adolescents with the lowest frequency of yes answers (below 20%, Node 2) can be isolated from the rest, which have a higher frequency of yes answers (about 60%, Node 3). These 111 subjects are then further split to form two groups: one smaller group with a medium (below 40%, Node 4) and one larger group with a high (about 80%, Node 5) frequency of yes answers to the intention-to-smoke question.

From this example, we can see that, following the principle of impurity reduction, each split in the tree-building process results in daughter nodes that are more pure than the parent node in the sense that groups of subjects with a majority for either response class are isolated. The impurity reduction achieved by a split is measured by the difference between the impurity in the parent node and the average impurity in the two daughter nodes. Entropy measures, such as the Gini Index or the Shannon Entropy, are used to quantify the impurity in each node. These entropy measures have in common that they reach their minimum for perfectly pure nodes with the relative frequency of one response class being zero and their maximum for an equal mixture with the same relative frequencies for both response classes, as illustrated in Figure 3.

Although the principle of impurity reduction is intuitive and has added much to the popularity of classification trees, it can help our statistical understanding to think of impurity reduction as merely one out of many possible means of measuring the strength of the association between the splitting variable and the response. Most modern classification tree algorithms rely on this strategy and use the p values of association tests for variable and cutpoint selection. This
approach has additional advantages over the original impurity reduction approach, as outlined later.

Regardless of the split selection criterion, however, in each node the variable that is most strongly associated with the response variable (i.e., that produces the highest impurity reduction or the lowest $p$ value) is selected for the next split. In splitting variables with more than two categories, which offer more than one possible cutpoint, the optimal cutpoint is also selected with respect to this criterion. In our example, the optimal cutpoint identified within the range of the numeric predictor variable $\text{alcohol_per_month}$ is between the values 1 and 2, because subjects who drank alcohol on one or fewer occasions have a lower frequency of yes answers than those who drank alcohol in two or more occasions.

After a split is conducted, the observations in the learning sample are divided into the different nodes defined by the respective splitting variable and cutpoint, and in each node splitting continues recursively until some stop condition is reached. Common stop criteria are to split until (a) a given threshold for the minimum number of observations left in a node is reached or (b) a given threshold for the minimum change in the impurity measure is not met any more by any variable. Recent classification tree algorithms also provide statistical stopping criteria that incorporate the distribution of the splitting criterion (e.g., Hothorn, Hornik, & Zeileis, 2006), whereas early algorithms relied on pruning the complete tree to avoid overfitting.

The term overfitting refers to the fact that a classifier that adapts too closely to the learning sample not only discovers the systematic components of the structure that is present in the population, but also the random variation from this structure that is present in the learning data because of random sampling. When such an overfitted model is later

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**Figure 2.** Relative frequencies of both response classes in the inner nodes of the binary classification tree for the smoking data. The dark and light grey shaded areas again represent the relative frequencies of yes and no answers to the intention-to-smoke question in each group respectively.

**Figure 3.** Shannon entropy and Gini index as functions of the relative frequency of one response class. Pure nodes containing observations of only one class receive an impurity value of zero, whereas mixed nodes receive higher impurity values.
applied to a new test sample from the same population, its performance is poor because it does not generalize well. However, it should be noted that overfitting is an equally relevant issue in parametric models: With every variable, and thus every parameter, that is added to the regression model, its fit to the learning data improves, because the model becomes more flexible.

This is evident, for example, in the $R^2$ statistic reflecting the portion of variance explained by the model, which increases with every parameter added to the model. For example, in the extreme case where as many parameters as observations are available, any parametric model shows a perfect fit on the learning data, yielding a value of $R^2 = 1$, but performs poorly in future samples.

In parametric models, a common strategy to deal with this problem is to use significance tests for variable selection in regression models. However, one should be aware that in this case, significance tests do not work in the same way as in a designed study, where a limited number of hypotheses to be tested are specified in advance. In common forward and/or backward stepwise regression, it is not known beforehand how many significance tests will have to be conducted. Therefore, it is hard to control the overall significance level, which controls the probability of falsely declaring at least one of the coefficients as significant.

Advanced variable selection strategies, which have been developed for parametric models, use model selection criteria, such as Akaike’s information criterion and the Bayesian information criterion, which include a penalization term for the number of parameters in the model. For a detailed discussion of approaches that account for the complexity of parametric models, see Burnham and Anderson (2002) or Burnham and Anderson (2004).

Because information criteria, such as Akaike’s information criterion and the Bayesian information criterion are, however, not applicable to nonparametric models (see, e.g., Claeskens & Hjort 2008), in recursive partitioning the classic strategy to cope with overfitting is to prune the trees after growing them, which means that branches that do not add to the prediction accuracy in cross-validation are eliminated. Pruning is not discussed in detail here, because the unbiased classification tree algorithm of Hothorn et al. (2006), which is used here for illustration, uses $p$ values for variable selection and as a stopping criterion and therefore does not rely on pruning. In addition to this, ensemble methods, which are our main focus here, usually use unpruned trees.

**Prediction and interpretation of classification and regression trees.** Finally, a response class is predicted in each terminal node of the tree (or each rectangular section in the partition, respectively) by means of deriving from all observations in this node either the average response value in regression or the most frequent response class in classification trees. Note that this means that a regression tree creates a piecewise (or rectanglewise for two dimensions and cuboidwise in higher dimensions) constant prediction function.

Even though the idea of piecewise constant functions may appear very inflexible, such functions can be used to approximate any functional form, in particular nonlinear and nonmonotone functions. This is in strong contrast to classical linear or additive regression, where the effects of predictors are restricted to the additive form—the interpretation of which may appear easier, but which may also produce severe artifacts, because in many complex applications, the true data-generating mechanism is neither linear nor additive. We see later that ensemble methods, by combining the predictions of many single trees, can approximate functions more smoothly, too.

The predicted response classes in our example are the majority class in each node in Figure 1A, as indicated by the shading in Figure 1B: Subjects who had no friends who smoked as well as those who had one or more friends who smoked but who drank alcohol on one or fewer occasions were not very likely to intend to smoke, whereas those who had one or more friends who smoked and who drank alcohol on two or more occasions were likely to intend to smoke within the next year.

For classification problems, it is also possible to predict an estimate of the class probabilities from the relative frequencies of each class in the terminal nodes. In our example, the predicted probabilities for answering *yes* to the intention-to-smoke question would thus be approximately 17%, 34%, and 79%, respectively, in the three groups—which may preserve more information than the majority vote that merely assigns the class with a relative frequency of $\geq 50\%$ as the prediction.

Reporting the predicted class probabilities more closely resembles the output of logistic regression models and can also be used (e.g., for estimating treatment probabilities or propensity scores). Note, however, that no confidence intervals are available for the estimates, unless, for example, bootstrapping is used in combination with refitting to assess the variability of the prediction.

The easy interpretability of the visual representation of classification trees, which we have illustrated in this example, has added much to the popularity of this method (e.g., in medical applications). However, the downside of this apparently straightforward interpretability is that the visual representation may be misleading, because the actual statistical interpretation of a tree model is not trivial. Especially the notions of main effects and interactions are often used rather incautiously in the literature, as seems to be the case in Berk (2006, p. 272), where it is stated that a branch that is not split any further indicated a main effect. However, when splitting continues in the other branch created by the same variable, as is the case in the example of Berk (2006), this statement is not correct.
The term *interaction* commonly describes the fact that the effect of one predictor variable, in our example `alcohol_per_month`, on the response depends on the value of another predictor variable, in our example `friends_smoke`. For classification trees, this means that, if in one branch created by `friends_smoke` it is not necessary to split in `alcohol_per_month`, whereas in the other branch created by `friends_smoke` it is necessary, as in Figure 1A, an interaction between `friends_smoke` and `alcohol_per_month` is present.

We further illustrate this important issue and source of misinterpretations by means of varying the effects in our artificial data set. The resulting classification trees are given in Figure 4. Only Figure 4A, where the effect of `alcohol_per_month` is the same in both branches created by `friends_smoke`, represents two main effects of `alcohol_per_month` and `friends_smoke` without an interaction: The main effect of `friends_smoke` shows in the higher relative frequencies of `yes` answers in Nodes 6 and 7 as compared to Nodes 3 and 4. The main effect of `alcohol_per_month` shows in the higher relative frequencies of `yes` answers in Nodes 4 and 7 as compared to Nodes 3 and 6, respectively.

As opposed to that, both Figure 4B and Figure 1A represent interactions, because the effect of `alcohol_per_month` is different in both branches created by `friends_smoke`. In Figure 4B, the same split in `alcohol_per_month` is conducted in every branch created by `friends_smoke`, but the effect on the relative frequencies of the response classes is different: For those subjects who have no friends who smoke, the relative frequency of a `yes` answer is higher if they drank alcohol in two or more occasions (Node 4 as compared to Node 3), whereas for those who have one or more friends that smoke, the frequency of a `yes` answer is lower if they drank alcohol on two or more occasions (Node 7 as compared to Node 6). This example represents a typical interaction effect as known from standard statistical models, where the effect of `alcohol_per_month` depends on the value of `friends_smoke`.

In Figure 1A, on the other hand, the effect of `alcohol_per_month` is also different in both branches created by `friends_smoke`, because `alcohol_per_month` has an effect only in the right branch, but not in the left branch. Although this kind of asymmetric interaction is very common in classification trees, one is unlikely to discover a symmetric interaction pattern like that in Figure 4B or even a main effect pattern like that in Figure 1A in real data. The reason for this is that, even if the true distribution of the data in both branches were very similar, because of random variations in the sample and the deterministic variable and cutpoint selection strategy of classification trees, it is extremely unlikely that the same splitting variable—and also the exact same cutpoint—would be selected in both branches. However, even a slightly different cutpoint in the same variable would, strictly speaking, represent an interaction. Thus, only if the two main effects and their respective cutpoints are very clear—and no other competitor variable is strong enough to outperform the two original variables in either node—the main effects pattern would be identified by a tree.

**Figure 4.** Classification trees based on variations of the smoking data with two main effects (Panel A) and interactions (Panel B). The tree depicted in Figure 1 that is based on the original data also represents an interaction.
Therefore, it is stated in the literature that classification trees cannot (or, rather, are extremely unlikely to) represent additive functions that consist only of main effects, although they are perfectly well suited for representing complex interactions. As opposed to that, standard regression models are—by definition—perfectly well suited for representing strictly additive functions but may not be able to identify complex interaction patterns and nonlinear effects.

In this sense, each statistical model imposes different limitations on the range of functions that can be represented by it—and may thus be more or less well suited to describing the (unknown) true structure of the data set at hand, which hardly ever follows a strict linear additive or strict stepwise recursive pattern. (Again, we find later that the ensemble methods bagging and random forests, which combine all patterns identified by a large set of single trees, can serve as a more flexible means for approximating different functional forms.)

Accordingly, what is easy for one class of statistical models may prove very hard for another class: Although it may seem surprising that classification trees cannot deal with such an easy problem as that of two main effects, one should note, for example, that logistic regression cannot deal with the—may be even easier—problem of perfectly separable response classes (in which case the coefficient estimates become infinite, i.e., there is no unique maximum-likelihood solution unless, e.g., a penalty term is used).

For exploratory data analysis, further means for illustrating the effects of particular variables in classification trees are provided by the partial dependence plots described in Hastie, Tibshirani, and Friedman (2001, 2009) and the CARTscans toolbox (Nason, Emerson, & Leblanc, 2004).

**Model-based recursive partitioning.** A variant of recursive partitioning, which can also be a useful aid for data exploration, is model-based recursive partitioning. Here, the idea is to partition the feature space not to identify groups of subjects with similar values of the response variable, but to identify groups of subjects with similar values of the parameters of a model of interest.

For example, linear regression could be used to model the dependence of a clinical response on the dose of medication. However, the slope and intercept parameters of this regression may be different for different groups of patients: older patients, for example, may show a stronger reaction to the medication; therefore, the slope of their regression line would need to be steeper than that of younger patients, or a group of nonresponders with a flat regression line may be identified by means of a combination of covariates. In this example, the model of interest is the regression between dose of medication and clinical response—however, the model parameters need to be chosen differently in the two or more groups defined by the covariates. Another example and visualization are given in the “Further Application Examples” section.

The model-based recursive partitioning approach of Zeileis, Hothorn, and Hornik (2009) offers a way to partition the feature space to detect parameter instabilities in the parametric model of interest by means of a structural change test framework. Similar to latent class or mixture models, the aim of model-based partitioning is to identify groups of subjects for which the parameters of the parametric model differ. However, in model-based partitioning, the groups are usually not defined by a latent factor but by combinations of observed covariates, which are searched heuristically. Thus, model-based partitioning can offer a heuristic but easy-to-interpret alternative to latent class—as well as random or mixed effects—models.

An extension of model-based partitioning for Bradley–Terry models is suggested by Strobl, Wickelmaier, and Zeileis (2009). An application to mixed models, including the Rasch model as a special case (as a generalized linear mixed model, see Doran, Bates, Bliese, & Dowling, 2007; Rijmen, Tuerlinckx, Boeck, & Kuppens, 2003), has been presented by Sanchez-Espigares and Marco (2008).

**What is wrong with trees?** The main flaw of simple tree models is their instability to small changes in the learning data: In recursive partitioning, the exact position of each cutpoint in the partition, as well as the decision about which variable to split in, determines how the observations are split up in new nodes, in which splitting continues recursively. However, the exact position of the cutpoint and the selection of the splitting variable strongly depend on the particular distribution of observations in the learning sample.

Thus, as an undesired side effect of the recursive partitioning approach, the entire tree structure could be altered if the first splitting variable, or only the first cutpoint, was chosen differently because of a small change in the learning data. Because of this instability, the predictions of single trees show a high variability.

The high variability of single trees can be illustrated, for example, by drawing bootstrap samples from the original data set and investigating whether the trees built on the different samples have a different structure. The rationale of bootstrap samples, where a sample of the same size as the original sample is drawn with replacement (so that some observations are left out, whereas others may appear more than once in the bootstrap sample) is to reflect the variability inherent in any sampling process: Random sampling preserves the systematic effects present in the original sample or population, but in addition to this, it induces random variability. Thus, if classification trees built on different bootstrap samples vary too strongly in their structure, this proves that their interpretability can be severely affected by the random variability present in any data set. Classification trees built on four bootstrap samples drawn from our original smoking data are displayed in Figure 5. Apparently, the effect of the variable friends_smoke is strong enough to
remain present in all four trees, whereas the further splits vary strongly with the sample.

As a solution to the problem of instability, the average over an ensemble of trees, rather than a single tree, is used for prediction in ensemble methods, as outlined in the following. Another problem of single trees, which is solved by the same model-averaging approach, is that the prediction of single trees is piecewise constant and thus may jump from one value to the next even for small changes of the predictor values. As described in the next section, ensemble methods have the additional advantage that their decision boundaries are more smooth than those of single trees.

How Do Ensemble Methods Work?

The rationale behind ensemble methods is to base the prediction on a whole set of classification or regression trees, rather than a single tree. The related methods bagging and random forests vary only in the way this diverse set of trees is constructed: In both bagging and random forests, a set of trees is built on random samples drawn from the learning sample. The only difference between bagging and random forests is that in bagging, variable selection follows the same principle as in single classification trees, whereas in random forests, variable selection is also randomized by means of random sampling from the set of all predictor variables to make the resulting set of trees even more diverse. Thus, first we explain the bagging procedure, which is based solely on random sampling from the learning data, and second we explain in more detail the random sampling from the predictor variables that distinguishes random forests from bagging.

Bagging. In each step of the algorithms for bagging and random forests, either a bootstrap sample (of the same size, drawn with replacement) or a subsample (of smaller size, drawn without replacement) of the learning sample is drawn randomly, and an individual tree is grown on each sample. As we saw earlier, each random sample reflects the same data-generating process but differs slightly from the original learning sample because of random variation. Keeping in mind that each individual classification tree depends highly on the learning sample as outlined earlier, the resulting trees can differ substantially. Another feature of the ensemble methods bagging and random forests is that usually trees are grown very large, without any stopping or pruning involved.
As illustrated again for four bootstrap samples from the smoking data in Figure 6, large trees can become even more diverse and include a large variety of combinations of predictor variables.

By combining the prediction of such a diverse set of trees, ensemble methods utilize the fact that classification trees are unstable but, on average, produce the right prediction (i.e., trees are unbiased predictors), which has been supported by several empirical as well as simulation studies (cf., e.g., Bauer & Kohavi, 1999; Breiman, 1996a, 1998; Dietterich, 2000) and especially by the theoretical results of Bühlmann and Yu (2002), which show the superiority in prediction accuracy of bagging over single classification or regression trees: Bühlmann and Yu (2002) were able to show by means of rigorous asymptotic methods that the improvement in the prediction is achieved by means of smoothing the hard cut decision boundaries created by splitting in single classification trees, which in return reduces the variance of the prediction (see also Biu, Devroye, & Lugosi 2008). The smoothing of hard decision boundaries also makes ensembles more flexible than single trees in approximating functional forms that are smooth rather than piecewise constant.

Grandvalet (2004) also pointed out that the key effect of bagging is that it equalizes the influence of particular observations—which proves beneficial when bad leverage points are downweighted but may be harmful when good leverage points that could improve the model fit are downweighted. The same effect can be achieved not only by means of bootstrap sampling as in standard bagging, but also by means of subsampling (Grandvalet, 2004), which is preferable in many applications because it guarantees unbiased variable selection (Strobl, Boulesteix, Zeileis, & Hothorn, 2007; see also section Bias in Variable Selection and Variable Importance). Ensemble construction can also

Figure 6. Classification trees (grown without stopping or pruning) based on four bootstrap samples of the smoking data, illustrating the principle of bagging.
be viewed in the context of Bayesian model averaging (cf., e.g., Domingos, 1997; Hoeting, Madigan, Raftery, & Volinsky, 1999, for an introduction). For random forests, which we consider in the next section, Breiman (2001a) stated that they may also be viewed as a Bayesian procedure but continued as follows: “Although I doubt that this is a fruitful line of exploration, if it could explain the bias reduction, I might become more of a Bayesian” (p. 25).

Random forests. In random forests, an extra source of diversity is introduced when the set of predictor variables to select from is randomly restricted in each split, producing even more diverse trees. The number of randomly preselected splitting variables, termed mtry in most algorithms, as well as the overall number of trees, usually termed ntree, are parameters of random forests that affect the stability of the results and are discussed further in the “Features and Pitfalls” section. Obviously, random forests include bagging as the special case where the number of randomly preselected splitting variables is equal to the overall number of variables.

Intuitively speaking, random forests can improve the predictive performance even further as compared to bagging, because the single trees involved in averaging are even more diverse. From a statistical point of view, this can be explained by Breiman (2001a)’s theoretical results showing that the upper bound for the generalization error of an ensemble depends on the correlation between the individual trees, such that a low correlation between the individual trees results in a low upper bound for the error.

In addition to the smoothing of hard decision boundaries, the random selection of splitting variables in random forests allows predictor variables that were otherwise outplayed by a stronger competitor to enter the ensemble: If the stronger competitor cannot be selected, a new variable has a chance to be included in the model—and may reveal interaction effects with other variables that otherwise would have been missed.

The effect of randomly restricting the splitting variables is again illustrated by means of four bootstrap samples drawn from the smoking data: In addition to growing a large tree on each bootstrap sample, as in bagging, now the variable selection is limited to mtry = 2 randomly preselected candidates in each split. The resulting trees are displayed in Figure 7: We find that, because of the random restriction, the trees have become even more diverse; for example, the strong predictor variable friends_smoke is no longer chosen for the first split in every single tree.

The reason that even suboptimal splits in weaker predictor variables can often improve the prediction accuracy of an ensemble is that the split selection process in regular classification trees is only locally optimal in each node: A variable and cutpoint are chosen with respect to the impurity reduction they can achieve in a given node defined by all previous splits, but regardless of all splits yet to come. Thus, variable selection in a single tree is affected by order effects similar to those present in stepwise variable selection approaches for parametric regression (which is also unstable against random variation of the learning data, as pointed out by Austin & Tu 2004). In both recursive partitioning and stepwise regression, the approach of adding one locally optimal variable at a time does not necessarily lead (or, rather, hardly ever leads) to the globally best model over all possible combinations of variables.

Because, however, searching for a single globally best tree is not computationally feasible (a first approach involving dynamic programming was introduced by van Os & Meulman 2005), the random restriction of the splitting variables provides an easy and efficient way to generate locally suboptimal splits that can improve the global performance of an ensemble of trees. Alternative approaches that follow this rationale by introducing even more sources of randomness are outlined later.

Besides intuitive explanations of how ensemble methods work, recent publications have contributed to a deeper understanding of the statistical background behind many machine learning methods: The work of Bühlmann and Yu (2002) provided the statistical framework for bagging; Friedman, Hastie, and Tibshirani (2000) and Bühlmann and Yu (2003) provided the framework for the related method of boosting; and, most recently, Lin and Jeon (2006) and Biau et al. (2008) provided the framework for random forests. In their work, Lin and Jeon explored the statistical properties of random forests by placing them in a k nearest neighbor (kNN) framework, where random forests can be viewed as adaptively weighted k nearest neighbors with the terminal node size determining the size of the neighborhood. However, to be able to mathematically grasp a computationally complex method like random forests, which involves several steps of random sampling, simplifying assumptions are often necessary. Therefore, well-planned simulation studies still offer valuable assistance for evaluating statistical aspects of the method in its original form.

Alternative ensemble methods. Alternative approaches for building ensembles of trees with a strong randomization component are the random split selection approach of Dietterich (2000), where cutpoints from a set of optimal candidates are randomly selected, and the perfect random trees approach of Cutler (1999) and Cutler (2000), where both the splitting variable and the cutpoint are chosen randomly for each split.

Another very intuitive approach, which resides somewhere in between single classification trees and the ensemble methods we have covered so far, is the trees with extra splits (TWIX) approach (Potapov, 2008; Potapov, Theus, & Urbanek, 2006). Here, the building of the tree ensemble starts in a single starting node but branches to a set of trees at each decision by means of splitting not only in the best cutpoint but also in reasonable extra cutpoints. A data-
driven approach for selecting extra cutpoints is suggested in Strobl and Augustin (2009).

However, although the approaches involving a strong randomization component manage to overcome local optimality as outlined earlier, the TWIX approach is still limited to a sequence of locally optimal splits. It has been shown to outperform single trees and even to reach the predictive performance of bagging, but, in general, it cannot compete because it becomes computationally infeasible for large sets of trees that are standard in today’s ensemble methods.

Predictions From Ensembles of Trees

In an ensemble of trees, the predictions of all individual trees need to be combined. This is usually accomplished by means of (weighted or unweighted) averaging in regression or voting in classification.

The term voting can be taken literally here: Each subject with given values of the predictor variables is dropped through every tree in the ensemble, so that each single tree returns a predicted class for the subject. The class that most trees vote for is returned as the prediction of the ensemble. This democratic voting process is the reason why ensemble methods are also called committee methods. Note, however, that there is no diagnostic for the unanimity of the vote. For regression and for predicting probabilities (i.e., relative class frequencies), the results of the single trees are averaged; some algorithms also use weighted averages. A summary over several aggregation schemes is given in Gattin (2008). However, even with the simple aggregation schemes used in the standard algorithms, ensembles methods reliably outperform single trees and many other advanced methods (examples of benchmark studies are given in the “Discussion and Conclusions” section).
Aside from the issue of aggregation, for bagging and random forests there are two different prediction modes: ordinary prediction and the so-called out-of-bag prediction. Whereas in ordinary prediction each observation of the original data set—or a new test data set—is predicted by the entire ensemble, out-of-bag prediction follows a different rationale: Remember that each tree is built on a bootstrap sample that serves as a learning sample for this particular tree. However, some observations, namely the out-of-bag observations, were not included in the learning sample for this tree. Therefore, they can serve as a built-in test sample for computing the prediction accuracy of that tree.

The advantage of the out-of-bag error is that it is a more realistic estimate of the error rate that is to be expected in a new test sample than the naive and overoptimistic estimate of the error rate resulting from the prediction of the entire learning sample (Breiman, 1996b) (see also Boulesteix, Strobl, Augustin, & Daumer, 2008, for a review on resampling-based error estimation). For example, the standard accuracy and the out-of-bag prediction accuracy for bagging in our smoking data example are 78% and 76.5%, respectively, where the out-of-bag prediction accuracy is more conservative.

However, in this very simple artificial example, random forests and even a single tree would perform as well as bagging, because the interaction of friends_smoke and alcohol_per_month, which was already correctly identified by the single tree, is the only effect that was induced in the data, whereas in most real data applications—especially in cases where many predictor variables work in complex interactions—the prediction accuracy of random forests is found to be higher than for bagging, and both ensemble methods usually highly outperform single trees.

### Variable Importance

As described in the previous sections, single classification trees are easily interpretable, both intuitively at first glance and descriptively when looking in detail at the tree structure. In particular, variables that are not included in the tree did not contribute to the model—at least not in the context of the previously chosen splitting variables. As opposed to that, ensembles of trees are not easy to interpret at all, because the individual trees in them are not nested in any way: Each variable may appear at different positions, if at all, in different trees, as depicted in Figures 6 and 7, so that there is no such thing as an average tree with a simple structure, that could be visualized for interpretation.

On the other hand, an ensemble of trees has the advantage of giving each variable the chance to appear in different contexts with different covariates; thus, the ensemble can better reflect that variable’s potentially complex effect on the response. Moreover, order effects induced by the recursive variable selection scheme used in constructing the single trees are eliminated by averaging over the entire ensemble. Therefore, in bagging and random forests variable importance measures are computed to assess the relevance of each variable over all trees of the ensemble.

In principle, a possible naive variable importance measure would be to merely count the number of times each variable is selected by all individual trees in the ensemble. More elaborate variable importance measures incorporate a (weighted) mean of the improvements of the individual trees in the splitting criterion produced by each variable (Friedman, 2001). An example for such a measure in classification is the Gini importance available in random forest implementations. It describes the average improvement in the Gini gain splitting criterion that a variable has achieved in all of its positions in the forest. However, in many applications involving predictor variables of different types, this measure is biased, as outlined in the Bias in Variable Selection and Variable Importance section.

The most advanced variable importance measure available in random forests is the permutation accuracy importance measure (termed permutation importance in the following). Its rationale is the following: By randomly permuting the values of a predictor variable, its original association with the response is broken. For example, in the original smoking data, adolescents who drank alcohol on more occasions were more likely to intend to smoke. Randomly permuting the values of alcohol_per_month over all subjects, however, destroys this association. Accordingly, when the permuted variable, together with the remaining unpermuted predictor variables, is then used to predict the response, the prediction accuracy decreases substantially. Thus, a reasonable measure for variable importance is the difference in prediction accuracy (i.e., the number of observations classified correctly; usually the out-of-bag prediction accuracy is used to compute the permutation importance) before and after permuting a variable, averaged over all trees.

If, on the other hand, the original variable was not associated with the response, either it is not included in the tree (and its importance for this tree is zero by definition), or it is included in the tree by chance. In the latter case, permuting the variable results in only a small random decrease in prediction accuracy, or the permutation of an irrelevant variable can even lead to a small increase in the prediction accuracy (if, by chance, the permuted variable happens to be slightly better suited than the original one). Thus, the permutation importance can even show (small) negative values for irrelevant predictor variables, as illustrated for the irrelevant predictor variable age in Figure 8B.

Note that in our simple example, the two relevant predictor variables friends_smoke and alcohol_per_
month are correctly identified by the permutation variable importance of both bagging and random forests, even though the positions of the variables vary more strongly in random forests (cf. again Figures 6 and 7). In real data applications, however, the random forest variable importance may reveal higher importance scores for variables working in complex interactions, which may have gone unnoticed in single trees and bagging (as well as in parametric regression models, where modeling high-order interactions is usually not possible at all).

Another important thing to note in the permutation importance scores for bagging and random forests displayed in Figure 8 is that, even though the two relevant predictor variables of the smoking data from bagging and random forests.

![Figure 8: Permutation variable importance scores for the predictor variables of the smoking data from bagging and random forests.](image)

Formally, the permutation importance for classification can be defined as follows: Let $B(t)$ be the out-of-bag sample for a tree $t$, with $t \in \{1, \ldots, n_{\text{tree}}\}$. Then, the importance of variable $X_j$ in tree $t$ is

$$VI_{t}(X_j) = \frac{\sum_{i \in \mathcal{B}(t)} I(y_i = \hat{y}^{(t)}_i) - \sum_{i \in \mathcal{M}} I(y_i = \hat{y}^{(t)}_{i,p})}{|\mathcal{B}(t)|}$$

(1)

where $\hat{y}^{(t)}_i = f(t(x_i))$ is the predicted class for observation $i$ before and where $\hat{y}^{(t)}_{i,p} = f(t(x_{i,p}))$ is the predicted class for observation $i$ after permuting its value of variable $X_j$, that is, with $x_{i,p} = (x_{i,1}, \ldots, x_{i,j-1}, x_{i,j-1,p}, x_{i,j+1}, \ldots, x_{i,p})$. Note that $VI_{t}(X_j) = 0$ by definition, if variable $X_j$ is not in tree $t$.

From this raw importance score, a standardized importance score can be computed with the following rationale: The individual importance scores $VI_{t}(X_j)$ are computed from $n_{\text{tree}}$ bootstrap samples, that are independent given the original sample, and are identically distributed. Thus, if each individual variable importance $VI_{t}(X_j)$ has standard deviation $\sigma$, the average importance from $n_{\text{tree}}$ replications has standard error $\sigma/\sqrt{n_{\text{tree}}}$. The standardized or scaled importance, also called $z$ score, is then computed as

$$z(x_j) = \frac{VI(x_j)}{\sigma \sqrt{n_{\text{tree}}}}$$

(3)

When the central limit theorem is applied to the mean importance $VI(x_j)$, Breiman and Cutler (n.d.) argued that the $z$ score is asymptotically standard normal. This property is
often used for a statistical test; however, it shows very poor statistical properties as outlined in the “Features and Pitfalls” section.

As already mentioned, the main advantage of the random forest permutation accuracy importance, as compared to univariate screening methods, is that it covers the impact of each predictor variable individually as well as in multivariate interactions with other predictor variables. For example, Lunetta et al. (2004) found that genetic markers relevant in interactions with other markers or environmental variables can be detected more efficiently by means of random forests than by means of univariate screening methods like Fisher’s exact test.

This, together with its applicability to problems with many predictor values, also distinguishes the random forest variable importance from the otherwise appealing approach of Azen, Budescu, and Reiser (2001) and advanced in Azen and Budescu (2003) for assessing the criticality of a predictor variable, termed 

dominance analysis: These authors suggested using bootstrap sampling and selecting the best regression model from all possible models for each bootstrap sample to estimate the empirical probability distribution of all possible models. From this empirical distribution for each variable the unweighted or weighted sum of probabilities associated with all models containing the predictor is computed and suggested as an intuitive measure of variable importance. This approach, where for \( p \) predictor variables \( 2^p - 1 \) models are fitted in each bootstrap iteration, has the great advantage of providing sound statistical inference. However, it is computationally prohibitive for problems with many predictor variables of interest, because all possible models have to be fitted on all bootstrap samples.

In random forests, on the other hand, a tree model is fit to every bootstrap sample only once. Then, the predictor variables are permuted in an attempt to mimic their absence in the prediction. This approach can be considered in the framework of classical permutation test procedures (Strobl, Boulesteix, Kneib, Augustin, & Zeileis, 2008) and is feasible for large problems, but it lacks the sound statistical background available for the approach of Azen et al. (2001). Another difference is that random forest variable importances reflect the effect of a variable in complex interactions as outlined earlier, whereas the approach of Azen et al. reflects the main effects—at least as long as interactions are not explicitly included in the candidate models. A conditional version of the random forest permutation importance that resembles the properties of partial correlations rather than those of dominance analysis was suggested by Strobl et al. (2008).

**Literature and Software**

Random forests have only recently been included in standard textbooks on statistical learning, such as Hastie et al. (2009; the previous edition, Hastie et al. 2001, did not yet cover this topic). In addition to a short introduction of random forests, this reference gives a thorough background on classification trees and related concepts of resampling and model validation; it is therefore highly recommended for further reading. For the social sciences audience, a first instructive review on ensemble methods, including random forests and the related method bagging, was given by Berk (2006). We suggest this reference for the treatment of unbalanced data (e.g., in the case of a rare disease or mental condition), which can be treated either by means of asymmetric misclassification costs or equivalently by means of weighting with different prior probabilities in classification trees and related methods (see also Chen, Liaw, & Breiman, 2004, for the alternative strategy of 
down sampling, i.e., sampling from the majority class as few observations as there are of the minority class), even though the interpretation of interaction effects in Berk (2006) is not coherent, as demonstrated earlier. The original works of Breiman (1996a, 1996b, 1998, 2001a, 2001b), to name a few, are also well suited and not too technical for further reading.

For practical applications of the methods introduced here, several up-to-date tools for data analysis are freely available in the R system for statistical computing (R Development Core Team, 2009). Regarding this choice of software, we believe that the supposed disadvantage of command line data analysis criticized by Berk (2006) is easily outweighed by the advanced functionality of the R language and its add-on packages at the state of the art of statistical research. However, in statistical computing, the textbooks also lag behind the latest scientific developments: The standard reference, Venables and Ripley (2002), does not (yet) cover random forests either, whereas the handbook of Everitt and Hothorn (2006) gives a short introduction to the use of both classification trees and random forests. This handbook, together with the instructive examples in the following section and the R code provided in an online supplement to this work, can offer a good starting point for applying random forests to data. Interactive means of visual data exploration in R that can support further interpretation are described in Cook and Swayne (2007).

**Further Application Examples**

For further illustration, two additional application examples of model-based partitioning and random forests are outlined. The data and source code for reproducing all steps of the following analyses as well as the examples in the previous sections in the R system for statistical computing (R Development Core Team, 2009) are provided as an online supplement.
Model-Based Recursive Partitioning

From a study on attitudes toward statistics among university students, several covariates (gender, age, major subject of study, and whether the person achieved his or her high school diploma through continuing education) are available for a sample of 430 first-year students together with their scores on the Cognitive Competence scale of the Survey of Attitudes Towards Statistics (Schau, Stevens, Dauphinee, & Vecchio, 1995) and their statistics grade in the final exam (see also Strobl, Dittrich, Seiler, Hackensperger, & Leisch, 2009, for details on the study).

As an example for model-based partitioning, we consider a linear regression model for predicting the statistics grade in the final exam from the Cognitive Competence scale score obtained in the first week of the semester. (Although the linear regression model may not be perfectly suited for describing these data, it is very well suited for illustrating the principle of model-based partitioning because it has only two model parameters: intercept and slope.)

In the resulting partition, the parameters of the linear regression model vary with respect to the students’ major subject and age, whereas the remaining covariates show no effect on the model parameters. As illustrated in Figure 9, although the intercept is similar, the slope parameter varies notably among all three groups.

The results imply a different impact of the cognitive aspect of the attitude toward statistics in the different groups of students: For students over the age of 20 with a business science major (Node 4), an increase in their negative attitude implies the strongest aggravation of the expected performance in the final exam, whereas for students with a social science major (Node 5), it implies the least, and for students up to the age of 20 with a business science major (Node 3), it implies a medium aggravation.

Of course, these differences among the groups of students could also be modeled by means of, for example, random effects or latent class models—but again the visual inspection of the model-based partition, which requires no further assumptions, can provide a helpful first glance impression of different association patterns present in the sample.

Random Forests

When the number of variables is very high, as, for example, in gene expression studies, parametric regression models are no longer applicable, and ensemble methods are often applied for prediction and the assessment of variable

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**Figure 9.** Model-based partition for the attitude toward statistics data. The model of interest relates the final statistics grade to the Cognitive Competence score obtained in the first week. (For the interpretation, note that the Cognitive Competence score was recoded such that high values correspond to a negative attitude, i.e., agreement with items such as “I will find it difficult to understand statistical concepts,” and numerically high grades indicate poor performance.)
importance. For an exemplary analysis of gene data, we adopted a data set originally presented by Ryan et al. (2006): The data were collected in a case-control study on bipolar disorder including 61 samples (from 30 cases and 31 controls) from the dorsolateral prefrontal cortex cohort. In the original study of Ryan et al., no genes were clearly found to be differentially expressed (i.e., to have an effect on the disease) in this sample. Therefore, two genes were artificially modified to have an effect, so that we can later ascertain whether these genes are correctly identified.

To be able to illustrate the variable importances in a plot, in addition to the two simulated genes and the three covariates age, gender, and brain pH level, a subset of 100 genes was randomly selected from the 22,283 genes originally presented by Ryan et al. (2006) for the example. Note, however, that the application to larger data sets is only a question of computation time. The permutation importances for all 105 variables are displayed in Figure 10. The effects of the two artificially modified genes can be clearly identified. With respect to the remaining variables, a conservative strategy for exploratory screening would be to include all genes whose importance scores exceed the amplitude of the largest negative scores (that can only be due to random variation) in future studies.

A prediction from the random forest can be given either in terms of the predicted response class or the predicted class probabilities, as illustrated in Table 1 for some exemplary subjects, with a mismatch between the true and predicted class for Subject 29. For the entire learning sample, the prediction accuracy estimate is overly optimistic (90.16%), whereas the estimate based on the out-of-bag sample is more conservative (67.21%). The confusion matrices in Table 2 display misclassifications separately for each response class.

Note that in this example, the application of logistic regression would be problematic even for the reduced data set with only 102 genes and even if only main effects are considered: Obviously, estimating the full model including all variables at a time is not possible when the number of predictors exceeds the sample size. However, even in forward stepwise selection, there are several occurrences of perfectly separable classes (inducing unidentifiable coefficient estimates), so that—aside from the issue of order effects—the model selection path is questionable.

In general, however, the complex random forest model, involving high-order interactions and nonlinearity, should be compared to a simpler model (e.g., a linear or logistic regression model including only low-order interactions) whenever possible to decide whether the simpler, interpretable model would be equally adequate. To further explore and interpret the effects and interactions of the predictor variables that were found relevant in a random forest, multivariate data visualization tools, such as those described in Cook and Swayne (2007), are strongly suggested.

Features and Pitfalls

The way recursive partitioning methods—in particular the ensemble methods bagging and random forests—work induces some special characteristics that distinguish them from other (even other nonparametric) approaches. Some of these special features are mostly technical, whereas others can prove very beneficial in applications, and yet others may pose severe practical problems, which we want to address here.

Small n Large p Applicability

The fact that variable selection can be limited to random subsets in each step of random forests makes them particularly well applicable in small $n$ large $p$ problems with many more variables than observations and has added much to the popularity of random forests. However, even if the set of candidate predictor variables is not restricted as in random forests, but covers all predictor variables as in bagging, the search is only a question of computational effort: Unlike logistic regression models, for example, where parameter estimation is not possible (e.g., because of linear constraints in the predictors or perfect separation of response classes in some predictor combinations as in the previous example) when there are too many predictor variables and too few observations, tree-based methods like bagging and random forests consider only one predictor variable at a time and can thus deal with high numbers of variables sequentially. Therefore, Bureau et al. (2005) and Heidema et al. (2006) pointed out that the recursive partitioning strategy is a clear advantage of random forests as opposed to more common methods like logistic regression in high-dimensional settings.

Nonlinear Function Approximation

Classification and regression trees are provably Bayes consistent, that is, in principle they can approximate any decision boundary, whether linear or highly nonlinear, given a sufficiently large data set and if allowed to grow at a proper rate (see, e.g., Devroye, Györfi, & Lugosi, 1996). For linear functions, the problem from a practical point of view is that a single tree’s step-function approximation is rather poor. Ensembles of trees, however, can approximate functions more smoothly by averaging over the step-functions of the single trees.

Therefore, bagging and random forests can be used to approximate any unknown function, even if it is nonlinear and involves complex interactions. An advantage of ensemble methods in this context is that, as compared to other nonlinear regression approaches, such as smoothing splines, neither the shape of the function nor the position or number of knots needs to be prespecified (see, e.g., Wood, 2006, for knot selection approaches in generalized
Figure 10. Variable importances for the original and modified gene data.
additive models). On the other hand, the resulting functional shape cannot be interpreted or grasped analytically and (aside from measures of overall variable importance) can only serve as a black box for prediction. This characteristic of many machine learning approaches has fueled discussions about the legitimacy and usefulness of such complex, nonlinear models (see, e.g., Hand, 2006, and the corresponding discussion).

In practice, for a given data set, where nonlinear associations or high-order interactions are suspected, complex approaches like random forests can at least serve as a benchmark predictor: If a linear or other parametric model with a limited number and degree of interaction terms can reach the (cross-validated or test sample) prediction accuracy of the more complex model, the extra complexity may be uncalled for and the simpler, interpretable model should be given preference. If, however, the prediction accuracy cannot be reached with the simpler model, and, for example, the high importance of a variable in a random forest is not reflected by its respective parameters in the simpler model, relevant nonlinear or interaction effects may be missing in the simpler model, and it may not be suited to grasping the complexity of the underlying process.

In addition to this, a black box method like random forests can be used to identify a small number of potentially relevant predictors from the full feature list, which can then be processed (e.g., by means of a familiar parametric method). This two-stage approach has been successfully applied in a variety of applications (see, e.g., Ward et al., 2006). Note, however, that variable selection should not be conducted before applying another statistical method on the same learning data (Ambroise & McLachlan, 2002; Boulesteix et al., 2008; Leeb & Pötscher, 2006).

The XOR Problem and Order Effects

In the literature on recursive partitioning, you may come across the so-called XOR problem (where XOR stands for the logical exclusive or concatenation), which describes a situation where two variables show no main effect but a perfect interaction. In this case, because of the lack of a marginally detectable main effect, none of the variables may be selected in the first split of a classification tree, and the interaction may never be discovered.

In such a perfectly symmetric, artificial XOR problem, a tree would indeed not find a cutpoint to start with. However, a logistic regression model would not be able to identify an effect in any of the variables either, if the interaction was not explicitly included in the logistic regression model—and in that case a tree model, where an interaction effect of two variables can also be explicitly added as a potential predictor variable, would do equally well.

In addition to this, a tree, and even better an ensemble of trees, is able to approximate the XOR problem by means of a sequence of cutpoints driven by random fluctuations that are present in any real data set. In this case, the random preselection of splitting variables in random forests again increases the chance that a variable with a weak marginal effect is still selected, at least in some trees, because some of its competitors are not available.

A similar argument applies to order effects when comparing stepwise variable selection in regression models with the variable selection that can be conducted on the basis of random forest variable importance measures: In both stepwise variable selection and single trees, order effects are present, because only one variable at a time is considered—in the context of the variables that were already selected but regardless of all variables yet to come. However, the advantage of ensemble methods, which use several parallel tree models, is that the order effects of all individual trees counterbalance, so that the overall importance ranking of a variable is much more reliable than its position in stepwise selection (see also Rossi et al., 2005).

Out-of-Bag Error Estimation

A feature that was already mentioned and used in the application example is that bagging and random forests come with their own built-in test sample: the out-of-bag observations, which provide a fair means of error estimation (Breiman, 1996b). Of course, similar validation strategies, based either on sample splitting or resampling techniques (see, e.g., Boulesteix et al., 2008; Hothorn, Leisch, Zeileis, & Hornik, 2005;) or ideally even on external validation

### Table 1
Predicted Response Class or Class Probability for Part of the Gene Data

<table>
<thead>
<tr>
<th>Subject</th>
<th>y</th>
<th>ŷ</th>
<th>( \hat{p}(y = 1) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>28</td>
<td>1</td>
<td>1</td>
<td>.80</td>
</tr>
<tr>
<td>29</td>
<td>1</td>
<td>2</td>
<td>.46</td>
</tr>
<tr>
<td>30</td>
<td>1</td>
<td>1</td>
<td>.64</td>
</tr>
<tr>
<td>31</td>
<td>2</td>
<td>2</td>
<td>.48</td>
</tr>
<tr>
<td>32</td>
<td>2</td>
<td>2</td>
<td>.43</td>
</tr>
</tbody>
</table>

### Table 2
Confusion Matrices With Prediction From Learning and Out-of-Bag Sample for the Gene Data

<table>
<thead>
<tr>
<th>ŷ</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Learning sample</td>
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<td>2</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>27</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ŷ</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Out-of-bag sample</td>
<td>20</td>
<td>10</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>21</td>
</tr>
</tbody>
</table>
samples (König, Malley, Weimar, Diener, & Ziegler, 2007), can and should be applied to any statistical method. However, in many disciplines, intensive model validation is not common practice. Therefore, a method that comes with a built-in test sample, like random forests, may help sensitize the issue and relieve the user of the decision for an appropriate validation scheme.

**Missing Value Handling by Means of Surrogate Splits**

Besides imputation approaches offered by some random forests algorithms, all tree-based methods provide another intuitive strategy for missing-value handling: This strategy is that, at first, observations that have missing values in the variable that is currently evaluated are ignored in the computation of the impurity reduction for this variable. However, the same observations are included in all other computations, so that the method does not involve cancelation of observations with missing values (which can result in heavy data loss).

After a splitting variable is selected, it would be unclear to what daughter node the observations that have missing values in this variable should be assigned. Therefore, a so-called **surrogate variable** is selected that best predicts the values of the splitting variable. By means of this surrogate variable, the observations can then be assigned to the left or right daughter node (see, e.g., Hastie et al., 2001). A flaw of this strategy is, however, that currently the permutation variable importance measure is not defined for variables with missing values.

**Bias in Variable Selection and Variable Importance**

In the classical classification and regression tree algorithms CART and C4.5, variable selection is biased in favor of variables with certain characteristics, even if these variables are no more informative than their competitors. For example, variables with many categories and numeric variables or, even more unintuitively, variables with many missing values are artificially preferred (see, e.g., Kim & Loh, 2001; Strobl, Boulesteix, & Augustin, 2007; White & Liu, 1994).

This bias is carried forward to ensembles of trees: Especially the variable importance can be biased when a data set contains predictor variables of different types (Strobl, Boulesteix, Zeileis, & Hothorn, 2007). The bias is particularly pronounced for the Gini importance, which is based on the biased Gini gain split selection criterion (Strobl, Boulesteix, & Augustin, et al., 2007), but can also affect the permutation importance. Only when subsamples drawn without replacement, instead of bootstrap samples, in combination with unbiased split selection criteria, are used in constructing the forest, can the resulting permutation importance be interpreted reliably (Strobl, Boulesteix, Zeileis, & Hothorn, 2007).

For applications in R, the functions `ctree` for classification and regression trees and `cforest` for bagging and random forests (both freely available in the add-on package, party; Hothorn et al., 2006; Hothorn, Hornik, & Zeileis, 2009) guarantee unbiased variable selection when used with the default parameter settings, as documented in the online supplement to this work.

The functions `tree` (Ripley, 2007) and `rpart` (Therneau & Atkinson, 2006) for trees and `randomForest` (Breiman, Cutler, Liaw, & Wiener, 2006; Liaw & Wiener, 2002) for bagging and random forests, on the other hand, which resemble the original CART and random forests algorithms more closely, induce variable selection bias and are not suggested when the data set contains predictor variables of different types.

**Scaled and Unscaled Importance Measures**

For the permutation importance, a scaled version, the $z$ score, is available or even the default in many implementations of random forests. The term **scaled** here is somewhat misleading, however, for two reasons: First, the variable importance does not depend on the scaling or variance of the predictor variables in the first place (in fact, the whole method is invariant against the scaling of numeric variables). Therefore, it is not necessary to account for the scaling of predictors in the variable importance. Second, for a scaled measure, one may assume that its values are comparable over different studies—which is not the case for the $z$ score in random forests that heavily depends on the choice of tuning parameters, as outlined in the next section. Therefore, we suggest not interpreting or comparing the absolute values of the importance measures, not even the $z$ scores, but relying only on a descriptive ranking of the predictor variables.

**Tests for Variable Importance and Variable Selection**

In addition to using variable importance measures as a merely descriptive means of data exploration, different significance tests and schemes for variable selection have been suggested: On the official random forests Web site, a simple statistical test based on the supposed normality of the $z$ score is proposed by Breiman and Cutler (n.d.); this test has been applied in a variety of studies—ranging from the investigation of predictors of attempted suicide (Baca-Garcia et al., 2007) to the monitoring of a large area space telescope on board a satellite (Paneque et al., 2007). This approach may appear more statistically advanced than a merely descriptive use of the random forest variable importance. However, it shows such alarming statistical properties that any statement of significance made with this
test is nullified (Strobl & Zeileis, 2008): Among other things, the power of this test depends on the arbitrarily chosen number of trees in the ensemble ntree, over which the importance is averaged (cf. Equations 2 and 3 in the Variable Importance section; see also Lunetta et al., 2004). Thus, reporting the significance of variable importance scores (like, e.g., Baca-Garcia et al., 2007, who did not even report the parameter settings they used for fitting the random forest) can be highly misleading, because the number of variables whose scores exceed a given threshold for significance depends on the arbitrary choice of a tuning parameter.

In addition to this, all statistical tests and variable selection schemes based on the original permutation importance, such as those suggested by Diaz-Uriarte and Alvarez de Andrés (2006) and Rodenburg et al. (2008), show another—potentially unwanted—peculiarity that is induced by the way the permutation importance is constructed: Correlated predictor variables are systematically preferred over uncorrelated ones. This issue is addressed in a permutation test framework by Strobl et al. (2008), who suggested a conditional importance measure for random forests. The interpretation of this conditional measure more closely resembles that of partial correlations and parametric regression model coefficients.

For selecting variables for further investigation in an exploratory study, we suggest a conservative decision aid for variable selection that was already hinted at in the application example: All variables with importance that is negative, zero, or positive but with a value that lies in the same range as the negative values can be excluded from further exploration. The rationale for this rule of thumb is that the importance of irrelevant variables varies randomly around zero. Therefore, positive variation of an amplitude comparable to that of negative variation does not indicate an informative predictor variable, whereas positive values that exceed this range may indicate that a predictor variable is informative.

**Randomness and Stability**

One special characteristic of random forests and bagging that new users are often not entirely aware of is that they are truly random models in the sense that, for the same data set, the results may differ between two computation runs. The two sources of randomness that are responsible for these possible differences are (a) the bootstrap samples (or subsamples) that are randomly drawn in bagging and random forests and (b) the random preselection of predictor variables in random forests. When the permutation importance is computed, another source of variability is the random permutation of the predictor vectors.

Because of these random processes, a random forest is only exactly reproducible when the random seed, a number that can be set by the user and determines the internal random number generation of the computer, is fixed. Otherwise, the results vary between two runs of the same code. To illustrate this point, random seeds are set in the online supplement code for the random forest application example whenever random sampling is involved.

The differences induced by random variations are, however, negligible—as long as the parameters of a random forest have been chosen to guarantee stable results:

- The number of trees ntree highly affects the stability of the model. In general, the higher the number of trees, the more reliable the prediction and the interpretability of the variable importance.
- The number of randomly preselected predictor variables mtry may also affect the stability of the model and the reliability of the variable importance. In general, random forests with random preselection perform better than bagging with no random preselection at all, but small values of mtry do not always prove beneficial: When predictor variables are highly correlated, the results of Strobl et al. (2008) indicate that a higher number of randomly preselected predictor variables is better suited to reflect conditional importance. In addition to that, if the number of randomly preselected predictor variables is very low, interactions of high order may be missed in the tree-building process. In situations with few relevant variables, “small mtry results in many trees being built that do not incorporate any of the relevant [variables]” (Diaz-Uriarte & Alvarez de Andrés, 2006), which would lead to a decrease in prediction accuracy.

The number of randomly preselected predictor variables can also be chosen to optimize prediction accuracy by means of cross validation in some algorithms. Note, however, that the choice of tuning parameters in random forests is not as critical as in other computer-intensive approaches, such as support vector machines (Svetnik, Liaw, Tong, & Wang, 2004), and random forests often produce good results even off the shelf without tuning.

- Note that the two tuning parameters, ntree and mtry, also interact: To assess a high number of predictor variables in a data set, a high number of trees or a high number of preselected variables for each split, or ideally both, are necessary so that each variable has a chance to occur in enough trees. Only then is its average variable importance measure based on enough trials to actually reflect the importance of the variable and not just a random fluctuation.

In summary, this means that if you observe that, for a different random seed, your prediction results and variable importance rankings (for the top-scoring variables) differ notably, you should not interpret the results but adjust the number of trees and preselected predictor variables.
Do Random Forests Overfit?

The study referred to in Breiman (2001b), where it is stated (and has been extensively cited ever since) that random forests do not overfit, may be a prominent example for a premature conclusion drawn from an unrepresentative sample. A variety of studies exploring the characteristics of machine learning tools, such as random forests, are based on only a few, real data sets that happen to be freely available in some data repository. The particular data sets investigated by Breiman (2001b) seem to enhance the impression that random forests would not overfit, but this notion was heavily criticized by Segal (2004).

The theoretical results of Breiman (1996a) do support the fact that ensemble methods do not overfit with an increasing number of trees. However, the real data “case studies” referred to in Breiman (2001b) do not exclude the possibility that they overfit for other reasons. For further methodological investigations of machine learning algorithms, we therefore strongly suggest the use of well-designed and controlled simulation experiments, rather than case studies with an unrepresentative selection of real data sets with unknown distributional properties, where analytical results are not feasible. With regard to the theoretical foundations and practical applications of random forests, Segal (2004) implied that the depth of the trees in random forests, rather than the number of trees (as suspected, e.g., by Luellen et al., 2005) may regulate overfitting.

Although most previous publications have argued that in an ensemble, each individual tree should be grown as large as possible and that trees should not be pruned, the recent results of Lin and Jeon (2006) also showed that creating large trees is not necessarily the optimal strategy. In problems with a high number of observations and few variables, a better convergence rate (of the mean squared error as a measure of prediction accuracy) can be achieved when the terminal node size increases with the sample size (i.e., when smaller trees are grown for larger samples). On the other hand, for problems with small sample sizes or even small n large p problems, growing large trees usually does lead to the best performance.

Discussion and Conclusion

Recursive partitioning methods have become popular and widely used tools in many scientific fields. Random forests especially have been widely applied in genetics and related disciplines within the past few years. First applications in psychology show that random forests can be of use in a wide variety of applications in this field as well. With this review, we hope to have given the necessary background for a successful—yet sensible—use of recursive partitioning methods, in particular of random forests, which have drawn much attention because of their applicability to even high-dimensional problems.

Besides the applications to regression and classification problems covered here, the function cforest (Hothorn et al., 2006, 2009) used in the application example can even be applied to survival data with a censored response and thus can also serve as a means of data exploration in a broad range of longitudinal studies. Of course, other recent statistical learning methods, such as boosting (Freund & Schapire, 1997) and support vector machines (cf. Vapnik, 1995, for an introduction), can also be applied to the scope of problems we suggested for the application of random forests. The performance of these methods is within close range of random forests; therefore, in some comparison studies, random forests clearly outperform their competitors (cf., e.g., Wu et al., 2003), whereas in others they are slightly outperformed (cf., e.g., König et al., 2008, for a comparison of several statistical learning methods in a medical example of moderate size, where logistic regression was also applicable).

In summary, one can conclude, in accordance with Heidema et al. (2006), that high-dimensional data should be approached by several different methods because each single method has its strengths and weaknesses: Boosting, for example, can be used for variable selection in linear and other additive models (see Bühlmann, 2006; Bühlmann & Hothorn, 2007, for an implementation in R). Similarly, shrinkage approaches like the LASSO (least absolute shrinkage and selection operator; cf., e.g., Hastie et al., 2001), the elastic net (Zou & Hastie, 2005), and the recent approach of Candes and Tao (2007) perform variable selection in linear models by means of penalization of the model coefficients. However, in contrast to random forests, for these methods it has to be assumed that the model is linear or additive and that the problem is sparse (meaning that only few predictor variables have an effect). For extremely small sample sizes, on the other hand, exact methods like the multivariate permutation tests described in Mielke and Berry (2001) or Good (2005) may be more suited.

With respect to ease of application, the results of the empirical comparisons between different supervised learning methods conducted by Caruana and Niculescu-Mizil (2006) and Svetnik et al. (2004) indicate that random forests are among the best performing methods even without extra tuning. Therefore, random forests can be considered as a valuable off-the-shelf tool for exploring complex data sets, which may in a few years become as popular in psychology as it is now in the fields of genetics and bioinformatics.

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INTRODUCTION TO RECURSIVE PARTITIONING


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Call for Nominations: Journal of Neuroscience, Psychology, and Economics

The Publications and Communications (P&C) Board of the American Psychological Association has opened nominations for the editorship of the Journal of Neuroscience, Psychology, and Economics, for the years 2011–2016. The editor search committee is chaired by Peter Ornstein, PhD.

The Journal of Neuroscience, Psychology, and Economics (JNPE), first published by Educational Publishing Foundation of the APA in 2009, publishes original research dealing with the application of psychological theories and/or neuroscientific methods to business and economics. Therefore, it is the first peer-reviewed scholarly journal that publishes research on neuroeconomics, decision neuroscience, consumer neuroscience, and neurofinance, besides more classical topics from economics and business research.

As an interdisciplinary journal, JNPE serves academicians in the fields of neuroscience, psychology, business, and economics and is an appropriate outlet for articles designed to be of interest, concern, and value to its audience of scholars and professionals.

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Candidates should be nominated by accessing APA’s EditorQuest site on the Web. Using your Web browser, go to http://editorquest.apa.org. On the Home menu on the left, find “Guests.” Next, click on the link “Submit a Nomination,” enter your nominee’s information, and click “Submit.”

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The deadline for accepting nominations is January 31, 2010, when reviews will begin.