Exploiting tree-based variable importances to selectively identify relevant variables

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Abstract. This paper proposes a novel statistical procedure based on permutation tests for extracting a subset of truly relevant variables from multivariate importance rankings derived from tree-based supervised learning methods. It shows also that the direct extension of the classical approach based on permutation tests for estimating false discovery rates of univariate variable scoring procedures does not extend very well to the case of multivariate tree-based importance measures.

Key words: Relevant feature selection, tree-based importance measures, false discovery rate, random permutations

1 Introduction

In the context of supervised learning, feature selection may be decomposed into two complementary subproblems. The first subproblem aims at identifying among a given set of candidate input variables a maximal subset of so-called relevant variables, i.e. variables which convey information about the target output variable, in isolation or in conjunction with other relevant variables. The second subproblem aims at identifying among a given set of variables, maximal subsets of so-called redundant variables, i.e. maximal subsets of variables which conditionally to the other variables in that given set do not convey complementary information about the target output variable. Unfortunately, when the sole information available about the problem under consideration is limited to a training sample of input-output pairs, it is not possible to exactly identify maximal subsets of relevant or redundant inputs. Thus, any feature selection algorithm will be at risk of either missing some sought features (false negatives) or of erroneously selecting some truly non desired ones (false positives).

In this paper we consider the problem of identifying relevant features from a (typically very large) set of candidate features. Among the different possible selectivity/sensitivity compromises, we aim at high selectivity, i.e. at identifying subsets of relevant variables while maintaining the rate of false positives as small as possible. This type of compromise is typically sought in the context of ‘biomarker discovery’ in bioinformatics, where input variables correspond for example to RNA or protein expressions, or genetic polymorphisms, and where one
seeks to identify a maximum number of them which truly provide information about some biological condition (disease status, treatment response, etc.) for further analysis and biological insight, while aiming at a very low false positive rate, because of high costs of subsequent experiments (see e.g. [9]).

Univariate statistical hypothesis tests provide only a partial answer to this question, because they can only identify variables that provide a significant amount of information about the output variable in isolation from the other inputs. When one seeks for interacting effects, one could resort to importance measures provided by supervised learning methods, such as tree-based methods. However, while these importance measures indeed allow to rank input variables by decreasing order of relevance, they still lack of a statistically sound procedure for selecting from their ranking a maximal set of variables while keeping the rate of false positives below a specified level. In particular, subset selection based on estimating generalization error rates by cross-validation generally does not imply a low false positive rate.

One possible way to derive from tree-based variable importance measures a procedure for selectively identifying relevant variables would be to mimic the procedures that are used in the context of multiple hypothesis testing of univariate statistics (see [5] for a review): the approach there is to rank the features according to a relevance score derived from a (univariate) hypothesis test, and then for each score threshold to estimate the rate of false positives (called the false discovery rate, or FDR) among the variables that have a score greater than the threshold. In this context, the FDR is usually estimated in a non parametric way by assessing the average scores derived when randomly permuting the output variable values of the dataset.

In this paper, we first assess this FDR estimation approach when the relevance scores are derived by tree-based (multivariate) importance measures. We show empirically that this simple procedure typically overestimates in an unpredictable way the real FDR and thus can lead to unreliable selections of relevant subsets. We explain this by the fact that, contrary to the univariate case, the tree-based importance scores of different variables are not independent of each other. We then propose a novel alternative procedure for assessing the presence of irrelevant variables among a subset of variables top-ranked by tree-based variable importance measures. For a given importance threshold, this procedure first assumes that all and only those variables which have received an importance higher than the threshold are truly relevant, and then estimates the probability that any of the other variables would receive an importance higher than the given threshold. Experiments suggest that the latter quantity (estimated by an appropriately adapted permutation scheme) allows indeed to rather well identify an importance threshold below which the risk of having at least one false positive rapidly increases. The procedure may thus serve to identify in a more robust way than the FDR based approach a maximal subset of truly relevant variables among those proposed by the importance scoring method.

The rest of the paper is organized as follows. Section 2 describes the particular tree-based supervised learning method, the corresponding variable importance
measure, and the synthetic datasets that we will use in the paper for our empirical tests. Section 3 studies the classical permutation based FDR estimation scheme on these synthetic datasets, highlighting its main properties when applied to tree-based variable importances. Section 4 describes the proposed alternative approach, its permutation based estimation procedure and the empirical results obtained with the same synthetic datasets. Section 5 shows some results on a real biological dataset. Finally, section 6 concludes and gives a few directions for further research.

2 Tree-based ensemble learners and importance measure

In this paper, we focus on classification problems. We assume that we have at our disposal a learning sample of \( N \) input-output pairs drawn from some unknown probability distribution. The \( m \) input variables are denoted \( f_i, i = 1, \ldots, m \).

**Tree-based methods.** The basic idea of classification trees [3] is to recursively split the learning sample with binary tests based each on one input variable trying to reduce as much as possible the uncertainty about the output classification in the resulting subsets of examples. Single classification trees are usually very much improved by ensemble methods, which aggregate the predictions of several trees. In our experiments, we will consider two tree-based ensemble methods based on randomization, namely Random Forests [1] and Extra-Trees [6]. For these two methods, we use the default parameter setting (i.e., the number of variables randomly selected at each node fixed at the square root of \( m \) and no pruning) and grow ensembles of 100 trees, unless specified otherwise.

**Variable importance measure.** Several variable importance measures have been proposed in the literature for tree-based methods. In this paper, we consider a measure based on information theory [12], which at each test node \( n \) computes the total reduction of the class entropy due to the split, defined by:

\[
I(n) = \#S.H_C(S) - \#S_t.H_C(S_t) - \#S_f.H_C(S_f),
\]

where \( S \) denotes the set of samples that reach node \( n \), \( S_t \) (resp. \( S_f \)) denotes its subset for which the test is true (resp. false), \( H_C(\cdot) \) is the Shannon entropy of the class frequencies in a subset, and \( \# \) denotes the cardinality of a set of samples. For a single tree, the overall importance \( v_i \) of the \( i \)th variable is then computed by summing the \( I \) values of all tree nodes where this variable is used to split. For an ensemble of trees, the importances are averaged over all individual trees.

A property of this measure is that the sum of the importances of all variables for a tree is equal to the total mutual information brought by the tree about the classification variable, which in the case of unpruned trees is usually very close to the initial total entropy of the classification variable [12]. The sum of the importances, for a single tree as well as for an ensemble of trees, is thus usually almost constant for a given problem. For presentation purpose, the importances are often normalized for the different variables so that they sum up to 100%.
Each importance is thus the percentage of the total information brought by the ensemble of trees that is due to this variable.

Artificial problems. To validate our methods in a context where relevant variables are perfectly known, we generated two artificial problems by adapting the Matlab code originally used to produce the Madelon dataset for the NIPS 2003 feature selection challenge\footnote{http://www.clopinet.com/isabelle/Projects/NIPS2003/}. Both problems are binary classification problems with continuous input variables.

- **Dataset-3-20**: This dataset is composed of 200 objects and 20 variables. The first three are really relevant, while the others are pure Gaussian noise. The problem is such that the third variable is only relevant in combination with the first two.
- **Dataset-50-1000**: The second (larger) dataset is composed of 2000 objects and 1000 variables. 50 variables are relevant, among which 6 have been directly used to define the output and 44 are linear combinations of these 6 variables (these latter are thus redundant given the first 6 ones). The remaining 950 irrelevant variables are pure Gaussian noise.

### 3 False discovery rate

We assume now that from the learning sample, we have computed with a tree-based method the importance \( v_i \) of each input variable \( f_i \), \( \forall i = 1, \ldots, m \), and, without loss of generality, we further assume that the features are ordered according to their importances, i.e.

\[
v_1 \geq v_2 \geq \ldots \geq v_m.
\]

For a given importance threshold \( v \), we consider that all variables whose importance is greater than \( v \) are relevant and our concern is to estimate the expected rate of truly irrelevant features among these variables, the so-called false discovery rate (FDR) \cite{Benjamini1995}.

More formally, for a given importance threshold \( v \), the FDR is defined as:

\[
\text{FDR}(v) = E \left[ \frac{F(v)}{S(v)} \right],
\]

where \( S(v) \) is the number of variables considered relevant at threshold \( v \) and \( F(v) \) is the number of those variables that are truly irrelevant. The expectation is taken over different random learning samples drawn from the (usually unknown) joint distribution of the variables (assuming that the algorithm is deterministic).

To select a subset of variables, one can then check the FDR for increasing values of the threshold \( v \) and choose the minimum value of \( v \) such that \( \text{FDR}(v) < \alpha \), where \( \alpha \) is typically small and reflects the risk one is ready to accept in terms of false positives when selecting the variables.
Table 1. Permutation algorithm for the estimation of the FDR

<table>
<thead>
<tr>
<th>Step</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>for (t = 1) to (T) (typically 1000) :</td>
</tr>
<tr>
<td></td>
<td>(a) Randomly permute the class labels.</td>
</tr>
<tr>
<td></td>
<td>(b) Compute variable importance values (v_j^t), (j = 1, \ldots, m), from the permuted data.</td>
</tr>
<tr>
<td></td>
<td>(c) Compute (R_i^t = #{k : v_k^t \geq v_i}), (\forall i = 1, \ldots, m).</td>
</tr>
<tr>
<td>2.</td>
<td>Then at (v_i), the FDR is estimated by (\hat{\text{FDR}}(v_i) = \frac{1}{T} \sum_{t=1}^T R_i^t) for (i = 1, \ldots, m).</td>
</tr>
</tbody>
</table>

3.1 Estimation by random permutations

To estimate this FDR, we adopt the same approximations as in [8]. When the number of features is large, one can approximate the expectation of the ratio by the ratio of the expectations:

\[
\text{FDR}(v) = E\left[\frac{F(v)}{S(v)}\right] \approx \frac{E[F(v)]}{E[S(v)]}. \tag{3}
\]

\(E[S(v)]\) can be simply approximated by the observed \(S(v)\), i.e. the number of variables with an importance greater than \(v\) in the original data. \(E[F(v)]\) is approximated by the expectation \(E[F(v)|H_0]\) over the null distribution \(H_0\) stating that all variables are truly irrelevant. In other words, \(E[F(v)]\) is taken as the expected number of variables that get an importance greater than \(v\) when none of them are truly relevant. We simulate this null hypothesis by applying the same tree-based method that was used to produce the original importances on datasets obtained from the original data by randomly permuting the class labels. This permutation decorrelates the classification variable from the inputs, making them all irrelevant, but keeps the dependencies that exist between the features in the original data.

The algorithm of Table 1 describes the procedure that we use to estimate the FDRs for all observed importance value thresholds in a learning sample.

3.2 Experiments on artificial data

We apply this procedure on the two artificial problems described in Section 2. Since we perfectly know the relevant variables for these two problems, we are also able to compute the observed FDR for a given subset of selected variables, i.e. the proportion of irrelevant variables among those selected. Figure 1 plots the
FDR estimated by the procedure of Table 1 and the observed FDR as a function of the rank of the variables on the two artificial datasets with Extra-trees and Random Forests. There is no important difference between the two methods. In all cases, we observe that the estimated FDR overestimates the observed FDR. On the small dataset, both methods are able to find the three relevant variables (as illustrated by the fact that the observed FDR is equal to 0 until rank 3). However, the curve of the estimated FDR that already starts increasing for the third variable wrongly suggests that this variable is a false positive. On the larger dataset, using a typical threshold of 0.05 on the estimated FDR leads to the selection of 26 variables with Extra-trees and 25 variables with Random Forests (all relevant in both cases), while the same threshold on the observed FDR would lead to 28 variables with Extra-trees and 33 variables with Random Forests, with 5% of irrelevant among them.

3.3 Discussion

The overestimation of the FDR by the procedure of Table 1 can be explained, at least partially, by the fact that this procedure does not take into account the dependence between the importance values for different variables. Indeed, as mentioned in Section 2, the sum of importances is roughly a constant for a given problem. In consequence, if a variable brings a lot of information about the classification, there is much less left to be explained by the remaining variables, whether they are relevant or not. Thus, if a relevant variable receives a high importance, it potentially hides a less important but still relevant variable that may consequently receive an importance which is small or even similar to that of an irrelevant variable in the permuted data. In this case, our estimation of

\begin{itemize}
  \item[Note] we are not talking here about the statistical dependence of the features that induces a dependence of their importances, however they are computed, but rather about the dependence between the importances that results from their joint computation by a multivariate approach.
  \item[Note] this applies whether or not the importances are normalized.
\end{itemize}
Fig. 2. Variable importance as a function of the rank with Extra-trees on Dataset-3-20, from the original and the permuted data. In the latter case, importances are averaged over the 1000 permutations.

$E[F(v_i)]$ from random permutations will be positively biased and thus our estimate of the FDR will be too conservative. This phenomenon is clearly apparent on the small artificial dataset, where the relevant variable $f_3$ gets an importance in the original ranking that is lower than the average importance obtained by the most important variable in the permuted data (see Figure 2).

In [8], Listgarten and Heckerman observed a similar effect when trying to estimate the false discovery rate among the edges predicted by a Bayesian network learning algorithm.

4 An alternative measure

In order to overcome this limitation of the FDR, we propose an alternative measure to be associated with each importance threshold $v_i$ that takes into account the importances of the variables that are ranked above $f_i$. For a given importance threshold $v_i$, the procedure consists in computing the following conditional probability, which we call the conditional error rate (CER):

$$\text{CER}(v_i) = P(\max_{k=i, \ldots, m} V_k^H \geq v_i | H_{R}^{1-i-1}, H_{1}^{i-m}),$$

(4)

where $H_{R}^{1-i-1}$ denotes the hypothesis that all variables above $f_i$ in the original ranking are relevant, $H_{1}^{i-m}$ is the hypothesis that $f_i$ and all the variables below $f_i$ are irrelevant, and $V_k^H (k = 1, \ldots, m)$ is the random variable denoting the importance of $f_k$ under these two hypotheses. CER($v_i$) is thus the probability that at least one irrelevant variable among $m-i+1$ gets an importance greater or equal to $v_i$, when these importances are computed under the assumption that variables $f_1, \ldots, f_{i-1}$ are all relevant.

This value can be interpreted as a measure of evidence against the hypothesis that all variables above the threshold $v_i$ are relevant: a CER($v_i$) close to one
Table 2. Permutation algorithm for the estimation of the CER

<table>
<thead>
<tr>
<th>Compute variable importances from the original data and assume, without loss of generality, that the variables are ordered according to their importance $v_i$, i.e. $v_1 \geq v_2 \geq \ldots \geq v_m$.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. for $i = 1$ to $m$ :</td>
</tr>
<tr>
<td>(a) for $t = 1$ to $T$ (typically 1000) :</td>
</tr>
<tr>
<td>$-$ Keep the class labels and the values of the $i - 1$ first variables of the original ranking fixed and randomly permute the values of the remaining variables using the same permutation vector for all variables</td>
</tr>
<tr>
<td>$-$ Compute variable importance values $v_{ij}^t$, $j = 1, \ldots, m$, from the permuted data.</td>
</tr>
<tr>
<td>(b) Then at $v_i$, the conditional probability (4) is estimated by:</td>
</tr>
<tr>
<td>$\hat{\text{CER}}(v_i) = \frac{# { t : \max_{j=i,\ldots,m} v_{ij}^t \geq v_i } }{T}$</td>
</tr>
</tbody>
</table>

means that it is very likely to observe an irrelevant variable with an importance above $v_i$ while a $\text{CER}(v_i)$ close to zero means that it is very unlikely that an irrelevant variable could reach the threshold $v_i$. The limit between relevant and irrelevant variables in the ranking can then be determined by looking for the minimal threshold $v_i$ such that $\text{CER}(v_i) < \alpha$, for some small value of $\alpha$.

Since the CER tries to detect at least one false positive above the threshold, we expect it to evolve much more abruptly than the FDR and thus that it will indicate more clearly the risk of selecting some irrelevant variables in the ranking.

### 4.1 Estimation by random permutations

We propose to estimate the probabilities (4) by random permutations as well. $H_{k_{i-1}}^{i-1}$ is approximated by keeping the class labels and the first $i-1$ variables unchanged (which amounts at considering that variables 1 to $i$ are truly relevant), while hypothesis $H_{k_{i-m}}^{i}$ is simulated by randomly permuting the values of the variables $i$ to $m$, that have an importance equal or smaller than $v_i$ in the original ranking. To adhere as much as possible to the original joint distribution of the variables, they are furthermore permuted jointly, i.e. using the same permutation vector. The resulting procedure is described in Table 2.

Because the importance of the variables $i$ to $m$ in the random permutations are computed with the values of the variables 1 to $i$ being unchanged, these importances should not suffer from the same bias as in the estimation of the FDR. We thus expect that the algorithm of Table 2 will produce unbiased estimates of the CER and thus be more adapted to highlight the true frontier between relevant and irrelevant features than the procedure of Table 1 based on the FDR.
4.2 Experiments on artificial data

Figure 3 compares the CER as estimated by the procedure of Table 2 with the FDR estimated by the procedure of Table 1 on both datasets and with the two ensemble methods.

On the small dataset, the CER correctly starts increasing at the fourth variable. It thus gives more chance than the FDR to the third variable to be selected.

On the larger dataset, the transition region between low and high CER is quite well centered at the point where irrelevant variables start appearing in the ranking (indicated by the vertical line). Setting a small threshold $\alpha = 0.05$ on the CER and looking for the last variable $f_i$ in the ranking such that CER($v_i$) < $\alpha$ leads to the selection of 22 variables with Extra-trees and 21 variables with Random Forests. In both cases, all the selected variables are truly relevant but the selection remains however quite conservative (with Extra-trees, the first 27 variables are relevant and with Random forests, the first 30). This is because the transition region between low and high CER is quite large (especially for Random Forests).

4.3 Link with FWER based univariate procedures

The CER has a nice interpretation when the importances are actually derived from univariate statistics and a variable is, by definition, irrelevant when it satisfies the null hypothesis $H_0$ of the corresponding statistical test (e.g. $v_i$ is the $t$-statistic associated to variable $i$). Indeed, in this case, importances $v_i$ are computed independently of each other and probability (4) can thus be rewritten as follows:

$$P(\max_{k=i,\ldots,m} V_k^H \geq v_i|H_1^{i-1}, H_i^{i-m}) = P(\max_{k=i,\ldots,m} V_k^H \geq v_i|H_1^{i-m})$$

$$= P(\max_{k=i,\ldots,m} V_k^H \geq v_i|H_i^{1-m}), \quad (5)$$
where $H^{x}_{J|m}$ is the hypothesis that all variables satisfy the null hypothesis $H_0$. Expression (5) corresponds precisely to the definition of Westfall and Young’s *stepdown maxT adjusted p-values* (see [13, 5]). The direct application of the procedure of Table 2 in this case thus produces estimates of these adjusted $p$-values by random permutations. Under some conditions about the distribution of the statistic, Westfall and Young showed that selecting all variables such that their adjusted $p$-values is lower than some threshold $\alpha$ guarantees that the family-wise error rate, or FWER (i.e. the probability to include at least one false positive among the selected variables) is lower than $\alpha$. In our context, however, given the strong dependency between the importances that are computed by tree-based methods, it is not clear whether this guarantee still applies.

### 5 Experiments on a real dataset

To highlight the behaviour of both measures on a real problem, we run experiments on a biological dataset. The goal of the study [4] is the identification of the genes with altered expression in the livers of knock-out mice compared to control mice. The dataset contains 5548 gene expression measurements for 16 mice divided into two classes: 8 wild-type mice and 8 mice whose Apo AI gene was knocked out. This dataset was also used in [5] to compare several statistical procedures for controlling multiple testing issues for univariate statistical tests.

Although the truly relevant variables are unknown, in [4], eight variables were identified as differentially expressed using univariate statistical tests and their relevance was experimentally confirmed. We expect that multivariate approaches will at least highlight these eight variables and we will thus check their presence in the rankings below. Note however that this does not mean that only those eight variables are relevant. All additional variables found by our multivariate procedures certainly deserve to be checked experimentally.

On this dataset, we apply the Extra-trees algorithm with four different settings of its parameters, i.e. the number $K$ of variables that are randomly selected at each node and the number $T$ of trees in the ensemble: $(K = \sqrt{m}, T = 100)$, $(K = \sqrt{m}, T = 1000)$, $(K = m, T = 100)$, and $(K = m, T = 1000)$. The estimated FDR and CER as a function of the ranking are plotted in Figure 4 in all four cases.

Several conclusions can be drawn from these plots. First, using $K = \sqrt{m}$ and $T = 100$ does not bring interesting results on this dataset. The method is unable to distinguish truly relevant variables from randomly permuted ones, which translates into a high value of both the FDR and the CER. Simply increasing the number of ensemble terms already gives much better results in terms of the number of variables that appear as relevant. A threshold of 0.05 on the CER selects a subset of 7 variables and a threshold of 0.05 on the FDR gives 10 variables that actually contains the 8 variables identified in [4]. Only 3 of them were present in the top 10 variables with $T = 100$, confirming that the ranking

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4 http://www.stat.berkeley.edu/users/terry/zarray/Html/apodata.html
Fig. 4. CER and FDR for increasing variable importance rank (Extra-Trees). Top left with \((K = \sqrt{m}, T = 100)\), top right with \((K = \sqrt{m}, T = 1000)\). Bottom left with \((K = m, T = 100)\), bottom right with \((K = m, T = 1000)\).

is indeed improved by increasing \(T\). It is interesting to note that, on the other hand, increasing \(T\) from 100 to 1000 does not affect the error rate (which is equal to 25% in both cases, as estimated by leave-one-out), meaning that here accuracy would not be a relevant criterion to assess the quality of the ranking. The high improvement of the FDR and CER values when \(T\) is increased is here a consequence of the very high ratio between the number of variables and the number of examples that makes the random trees, and thus the corresponding rankings, highly unstable and thus requires to average a very large number of trees for stabilization.

When \(K\) is increased to its maximum value (i.e. randomization is reduced), the CER is also very much improved, even with \(T = 100\). It shows an abrupt change between the 11th and the 12th variables, suggesting that about 11 variables are relevant. As a confirmation, the 8 variables identified in [4] are again among these 11 ones. On the other hand, the estimated FDR seems to be highly overestimated in this case. We explain this by the fact that increasing \(K\) in Extra-trees makes the model less random and thus increases the importance of the top ranked variables relatively to the low ranked ones, thus emphasizing the phenomenon highlighted in Section 3.3 and responsible for the overestimation of the FDR.

6 Conclusions

In this paper, we have proposed and evaluated two statistical procedures to extract a subset of truly relevant variables from multivariate importance rankings obtained from tree-based supervised learning methods. The first one is a direct application of FDR based procedures used in the context of univariate statistics. Unfortunately, this procedure can rather strongly overestimate the real FDR and can thus potentially lead to overly conservative selections of relevant subsets. We then have proposed an alternative procedure based on the estimation
of a conditional error rate, whose estimation by random permutations takes into account the dependencies between the importances and thus leads to more robust estimates. Although further experiments are needed to better validate this procedure, we believe that our results on artificial and real problems are already quite encouraging.

One drawback of the CER procedure is that it is very computationally demanding: $T \times m$ models are needed for the estimation of the CER for all possible importance thresholds, where $m$ is the number of variables and $T$ is the number of random permutations. The computing times can however be decreased, e.g. by stopping the procedure when the estimated probability is greater than some threshold, as we are typically only interested in small values of the CER.

In terms of future works, it would be interesting to investigate better procedures to estimate the FDR. As noted in Section 3.3, one of the reasons of the overestimation of the FDR is the overestimation of $E[F(v)]$ that results from the simultaneous permutation of all features. We have tried to adopt the same sequential approach for estimating this term as for the CER, i.e. only permuting the $m-i+1$ last variables and estimate $E[F(v_i)]$ as the expected number of permuted variables whose importance exceeds threshold $v_i$. This yields better estimates of the FDR for small $i$ but when $i$ grows to $m$, the number of permuted variables, and thus the FDR, decreases to 0. Further investigations are thus necessary to really assess the conditions of validity of this procedure. Several improved permutation procedures have also been proposed to better estimate the FDR in the context of univariate tests (e.g. [14]). The extension of these approaches to our multivariate context is also an interesting future work direction.

Along a different line, Stoppiglia et al. [10] have proposed a related approach for feature selection that introduces a random variable among the input variables and exploits its rank among the other variables to determine the relevance threshold. Although they developed this idea in the context of linear models only (where the distribution of the rank of the random feature can be computed analytically), this approach could be straightforwardly extended to tree based ranking. Its comparison with our approach will be the subject of future work.

Another important direction of investigation is of course the application of these procedures in the context of other multivariate importance measures derived from machine learning algorithms. In the context of tree-based methods, our analysis should carry over straightforwardly to regression measures based on variance reduction [3]. It would be interesting also to consider other importance measures such as, for example, Breiman’s permutation based importance measure [2].

On the other hand, we have already carried out some preliminary experiments with variable importances derived from the weights of linear SVM models [7]. One problem however with these importances is that their scale of measurement is dependent on one hand of the range of variation of the variables (although this can be circumvented by a prior normalization of the features) and on the other hand on the specific margin that can be achieved given the features (since the
margin is proportional to the norm of the weight vector). This scaling instability suggests that some further caution should be taken when comparing the original importances with importances derived from randomly permuted data as required in our procedure.

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