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Stable variable selection for right censored data: comparison of methods

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Abstract

The instability in the selection of models is a major concern with data sets containing a large number of covariates. This paper deals with variable selection methodology in the case of high-dimensional problems where the response variable can be right censored. We focus on new stable variable selection methods based on bootstrap for two different methodologies commonly used in survival analysis: the Cox proportional hazard model and survival trees. As far as the Cox model is concerned, we investigate the bootstrapping applied to two variable selection techniques: the stepwise algorithm based on the AIC criterion and the $L_1$-penalization of Lasso. Regarding survival trees, we review two methodologies: the bootstrap node-level stabilization and random survival forests. We apply these different approaches to two real data sets, a classical breast cancer data set and an original infertility data set. We compare the methods on two criteria: the prediction error rate based on the Harrell concordance index and the relevance of the interpretation of the corresponding selected models, focusing on the original infertility data set. The aim is to find a compromise between a good prediction performance and ease to interpretation for clinicians. Results suggest that in the case of a small number of individuals, a bootstrapping adapted to $L_1$-penalization in the Cox model or a bootstrap node-level stabilization in survival trees give a good alternative to the random survival forest methodology, known to give the smallest prediction error rate but difficult to interpret for non-statisticians. In a clinical perspective, the complementarity between the methods based on the Cox model and those based on survival trees would permit to built reliable models easy to interpret for the clinician.

Key-words: censored data, variable selection, survival trees, survival random forests, Lasso, Cox model, bootstrap.
1 Introduction

Problems of variable selection arouse a growing interest in the processing of data sets containing more and more variables. In the last twenty years, many methods of variable selection have been proposed to handle these high-dimensional problems, especially when the number of covariates $p$ exceeds the number of observations $n$. To avoid a wrong estimation due to collinearity problems and to improve interpretation, the scientific community has developed tools to select the most relevant variables. A large literature concentrates on the case of the linear regression. A classical well-known method is the stepwise algorithm based on the Akaike Information Criterion (AIC). Recently, another field of research has focused on optimization problems, such as $L_1$-penalty approaches. On the other way, tree-based algorithms provide an interesting alternative to handle non-parametrically a large number of covariates.

We consider the special case where the response variable is right censored. In this context, the Cox proportional hazards model [1] has become the gold-standard tool for the statistical analysis, especially in the medical field. However, in the case of a large amount of covariates, it may be very unstable, even when stepwise selection or $L_1$-penalization of Lasso are added to the classical procedure. The instability of these selection approaches have also been encountered in the context of the linear regression [2, 3]. To remedy this problem, some authors have proposed to use bootstrapping to investigate the reliability of the choice of the variables in the final model. Bach [4] introduced stability in the selection by using bootstrapping in a Lasso algorithm, a method called Bolasso, and Meinshausen and Bühlmann [3] proposed to improve the $L_1$-penalization by randomizing the selection process of the covariates. These bootstrapped Lasso methods have only been considered in the framework of linear regression: we propose to extend them to the stabilization of the selection of covariates in a Cox model.

Alternatively to the Cox model, survival trees procedures permit to take into account non linear relationships between the censored variable and the covariates and yield easily interpretable classification rules to the clinicians, but they tend to overfit the data and suffer also from instability [5–8]. A stabilization technique known to improve the prediction performance of a single tree consists in aggregating a family of tree models using boostrapped samples and a random selection of the covariates at each node of the trees. This procedure, called “random forests” [9] has been adapted to the survival framework by Ishwaran et al. [10]. This Random Survival Forest (RSF) methodology is considered as the best modelization in terms of prediction performance but it is not easy to interprete as it does not provide a single tree. On the other way, Dannegger [6] introduced tools to control the stability of the selection of the covariates at each node of a tree and propose resampling techniques at node-level to stabilize the choice of the split. This methodology has the advantage to keep the simplicity of interpretation of a single final tree.

We review and compare these different stable variable selection methods derived on the Cox model or on survival trees on real data sets. We will base the comparison of the
proposed methods on statistical criteria as the prediction accuracy but also on the capacity
of the method to supply a final model whose clinical interpretation is easy and pertinent.
As a matter of fact, the objective of the statistician is to provide to clinicians a model
which is robust enough to permit a proper understanding of the relationships between the
variable of interest and the covariates in a predictive and explanatory purpose.

We consider two real data sets. The first data set concerns the survival of breast
cancer patients in relation with their gene-expression signature [11] and is a very usual
case encountered in survival analysis. This classical example will validate the different
approaches discussed earlier. The second data set is much more original as it concerns
couples consulting for infertility and very few studies were led in this domain. The cen-
sored variable of interest is the duration between the first visit of the couple and the birth
of a alive child, which is difficult to model. The study takes into account several covariates
of the couple or man and woman, but also the associated medical treatment according
to the diagnosed causes of infertility. For example, a man with severe oligospermia will
receive ICSI (IntraCytoplasmic Sperm Injection), while a woman with a dysovulation or
with no tubal factor will enter an IVF (In Vitro Fertilization) program. In many couples,
causes of infertility are not identified and physicians can not clearly answer the question
of on-going pregnancy in couples.

In section 2, a presentation of the variable selection methods based on the Cox model
is given, followed, in section 3, by a review of the selection methods based on trees
algorithms. Section 4 compare the different approaches on the two data sets, giving
the prediction erro rate and the selected variables in the final model for each approach.
Concluding remarks and perspectives are presented in the last section.

2 Stable variable selection methods based on the Cox
model

Notations

We denote by $T$ the variable of interest, which is a time of failure. We suppose that $T$ may
be right censored at a non-informative censoring time $C$ such that $C$ is independent of $T$
conditionally on $Z$, a $p$-vector of covariates. The observations are therefore the possibly
censored time $X = \min(T, C)$ and the censoring indicator $\Delta = 1\{T \leq C\}$, where $1\{\cdot\}$ is
the indicator function. The observed sample is thus $(X_i, \Delta_i, Z_i)_{i=1,\ldots,n}$.

In survival analysis, the Cox model [1] has become the most popular method to model-
elize the relationship between a survival time and one or more predictors (i.e. covariates).
This model has the advantage to be semi-parametric in the sense that it does not re-
quire assumptions on the survival time distribution. Moreover, it is of easy interpretation
for clinicians in providing estimates of the effect of the covariates on survival time after
adjustment on the other covariates.
In the Cox regression model in its simplest form, the hazard function for the failure time of an individual takes the form
\[
\lambda(t|Z) = \lambda_0(t) \exp(\beta'Z),
\]
where $\beta$ is a $p$-vector of unknown regression parameters and $\lambda_0(t)$ is an unknown baseline hazard function. Denote $t_{(1)}, \ldots, t_{(k)}$ the $k$ ordered uncensored survival times. The parameter of interest $\beta$ is estimated by maximising the logarithm of the partial likelihood function
\[
L(\beta) = \sum_{i=1}^{k} \left( \beta'Z_i - \log \left( \sum_{j \in R_i} e^{\beta'Z_j} \right) \right) \tag{1}
\]
where $R_i$ is the set of indices of the individuals at risk at time $t_{(i)}$. Notice that the Cox model is a log-linear model based on the proportional hazard hypothesis, which stipulates that the ratio of the hazard functions of two individuals is constant over time.

In the following, we review three stable methods to select the relevant covariates using the Cox model: the bootstrap stepwise selection, the bootstrap Lasso selection and the bootstrap randomized Lasso selection.

### 2.1 Bootstrap stepwise selection (BSS)

In case of a large number of covariates, the selection of the predictors in the Cox model is usually made by a stepwise algorithm minimizing the Akaike Information Criteron (AIC). However, the stability of this method is questionable. Using a data splitting approach, Harrell et al. [2] showed variation in the selected predictors. This result was confirmed by Chen and George [12] who applied the stepwise procedure on 100 bootstrapped samples from a study of acute lymphocytic leukaemia. The original model which was built from the entire set by a Cox regression model coincide in only 2% of the cases with the models selected from the bootstrapped samples. Sauerbrei and Schumacher [13] developed a bootstrap selection procedure which combined the bootstrap method with stepwise selection in Cox regression. They examined the inclusion frequencies of the variables selected by the stepwise algorithm into the models derived from the bootstrapped samples and keep in the final model the variables for which the inclusion frequency exceeds a given cut-off value $\kappa$ in $(0, 1)$. The choice of $\kappa$ is arbitrary. The authors applied the methodology to a data set on brain tumours including 447 patients and 12 covariates, using two values of $\kappa$ (0.3 and 0.6). They showed that the choice of $\kappa = 0.6$ gives the same final model whatever the number of bootstrap samples (from 100 to 1000).
2.2 Bootstrap Lasso and bootstrap randomized Lasso selection

Lasso selection

An alternative to the stepwise selection procedure is the Lasso selection. Adapted by Tibshirani [14] to the Cox model, the method estimates the $\beta$ parameter via maximising the log partial likelihood function (1) with the constraint $\sum_{j=1}^{p} |\beta_j| \leq \lambda$ where $\lambda$ is a regularization parameter. The Lasso constraint selects variables by shrinking estimated coefficients towards 0. This leads to coefficients exactly equal to zero and allows a parsimonious and interpretable model. The choice of a proper $\lambda$ is sensitive and leads to variation in model selection as demonstrated by Meinshausen and Bühlmann [3]. In a linear regression framework, they studied the stability of the selection in a gene expression data set. They introduced noise in variables in permuting all but 6 of the 4088 gene expressions. They showed that, when $\lambda$ increases, the final model retained as well the 6 unpermuted variables as the irrelevant noised variables. Although cross-validation seems a natural solution to choose $\lambda$, it is not a good alternative for high-dimensional problems. As a matter of fact, the authors showed in their example that 14 permuted variables are still retained in the final model obtained in choosing $\lambda$ by cross-validation. Moreover, Meinshausen and Bühlmann [15] proved that if the number of covariates tends to infinity then the probability of selecting a wrong variable converges to 1.

Bootstrap Lasso selection (BLS)

In order to obtain a consistent model selection, Bach [4] proposed for the linear model a bootstrapped version of the Lasso, referred as the Bolasso. He defined the Bolasso model estimate as the final model composed by only the variables which are selected in all bootstrapped samples; in other words, using the terminology of the BSS, it corresponds to a cut-off value $\kappa$ equal to 1. However, Meinshausen and Bühlmann [3] showed with the permuted gene data set that even with random subsampling (which is a procedure close to the bootstrap [16]), the Lasso algorithm could select irrelevant variables when $\lambda$ is too large.

Bootstrap randomized Lasso selection (BRLS)

To deal with the choice of $\lambda$, Meinshausen and Bühlmann [3] proposed a generalisation of the bootstrap Lasso procedure called bootstrap randomized Lasso where the covariates are penalized by different values randomly chosen in the range $[\lambda, \lambda/\alpha]$ with $\alpha$ in $(0, 1)$. This turns out to estimate the $\beta$ parameter with the constraint $\sum_{j=1}^{p} \left| \frac{\beta_j}{W_j} \right| \leq \lambda$. In practice, the set of covariates $\{Z_j : j = 1, \ldots, p\}$ are weighted by the set $\{W_j : j = 1, \ldots, p\}$ randomly generated where $P(W_j = \alpha) = p_w$ and $P(W_j = 1) = 1 - p_w$ with $p_w$ in $(0, 1)$. The authors give no indication on the choice of $p_w$ and in the following we take $p_w = 0.5$. 

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This procedure is very simple to implement and the authors showed that choosing $\alpha$ in the range of $(0.2, 0.8)$ gives a consistent variable selection whatever the choice of the penalty $\lambda$. The authors explained that a low value of $\alpha$ will decrease the selection probabilities of irrelevant variables even if the penalty $\lambda$ is large. As the BLS and BRLS methods are based on a penalization of the parameter $\beta$, it is necessary to first normalize the covariates such as $||Z_j||_2 = (\sum_{i=1}^{n} Z_{ij}^2)^{1/2} = 1$ for all $j$ in $\{1, \ldots, p\}$.

3 Stable variable selection methods based on survival trees

Although they are not so popular than the Cox model, tree-based methods in survival analysis (the so-called survival trees) have known a great development in the last decades. They provide a good alternative to the Cox regression model in identifying covariates which play a role on the survival outcome and in predicting the individual risk of failure. In addition to be easy to interpret in a large frame of applications, survival trees methods can incorporate non linear effects, and also take into account interactions between covariates.

First developed for basic classification trees, the Classification and Regression Trees (CART) algorithm of Breiman [17] is based on binary recursive partitioning. This is an iterative process which splits the data into two subgroups (daughter nodes) according to the value of one of the predictors. The splitting rule maximises the difference between nodes. Let denote $\{Z_j : j = 1, \ldots, p\}$ the set of covariates. Formally, a split is induced by a question of the form “Is $Z_j \leq b$ ?” for a continuous covariate where $b$ is a cut-off value in the set of realizations of $Z_j$. For nominal covariates with possible values in the set $B = \{b_1, \ldots, b_r\}$, the question is of the form “Is $Z_j$ in $S$?” where $S \subset B$. To determine the best split $s$ among all covariates in the current node $h$, a measure of improvement $G(s, h)$ is evaluated, producing the most homogeneous daughter nodes. The best split $s_{best}$ is the optimal split among all possible splits in the set $S_h$ such as $G(s_{best}, h) = \max_{s \in S_h} G(s, h)$. The process goes on until each node reaches a user-specified minimum node size and becomes a terminal node, or is homogeneous. To control the size of the tree, a stopping rule is used to prune the “large” tree containing “pure” nodes. The method employed by CART is called cost-complexity pruning. The complexity of a tree $\Theta$ is :

$$R_{cp}(\Theta) = R(\Theta) + cp |\tilde{\Theta}|,$$

where $R(\Theta)$ is the raw error of measure (sum of the error of measure over the terminal nodes), $|\tilde{\Theta}|$ is the number of terminal nodes, and $cp$ is an arbitrary penalty weight between 0 and $\infty$ called the complexity parameter. A sequence of nested trees is built. The final tree is the smallest tree for which $R_{cp}(\Theta)$ is minimized. To deal with the choice of $cp$, cross-validation techniques can be used to determine the optimal size of the tree.

Various splitting and pruning approaches have been proposed to adapt regression trees to survival data. Gordon and Olshen [18] used the Wasserstein metrics to measure the
distance between two Kaplan-Meier estimates of the survival distribution. The split criterion chooses the predictor (and if necessary the cut-off) which maximises this distance for the left and right daughter nodes. To prune the survival tree, they generalized the cost-complexity measure for censored data. A more usual way to measure the difference between survival curves is the logrank test statistic. This was done by Ciampi et al. [19], Segal [20] and LeBlanc and Crowley [21] who suggested to use the logrank test statistic as a between-node heterogeneity measure. As an alternative to the standard cost-complexity pruning approach using proportional hazards martingale residuals as error measures, LeBlanc and Crowley [21] proposed a "goodness-of-split" complexity based on the sum of the standardized splitting logrank test statistics of the internal nodes. Others authors suggested to use split criteria based on the likelihood function. Davis and Anderson [22] assume that the survival function within each node is an exponential function with a constant hazard, while LeBlanc and Crowley [23] only assume proportionality for the hazard functions of two daughter nodes. These latter used for estimation the full or partial likelihood function in the Cox proportional hazards model.

Instability in the selection of covariates by regression trees has been observed and demonstrated by many authors [5 – 8] This instability may be due to an overfitting of data. The variance observed may also come from arbitrary cutpoints defined by the dichotomization of continuous covariates. We review in the following two remedies to instability of survival trees: the bootstrap node-level stabilization proposed by Dannegger [6] and the random survival forests of Ishwaran et al. [10].

### 3.1 Bootstrap node-level stabilization (BNLS)

Dannegger [6] proposed a bootstrap node-level stabilization procedure for survival trees. The algorithm consists, at node \( h \), in drawing bootstrapped samples from the original set, and for each of them, in finding the best split. The split which appears the most of the time at the node \( h \) is selected. For a continuous variable, the cut-off value \( b \) chosen in the set of realizations of the split variable is the median of all the \( b \)-values proposed at each bootstrap. As Dannegger [6] did not propose a choice of the cut-off for a categorical variable, we decide to affect to the daughter node the level of the categorical variable which was mainly chosen by the bootstrapped samples. To find the optimal complexity parameter \( cp \) to prune the tree, we use a ten-fold cross-validation procedure as suggested by Dannegger [6].

Using simulated data, Dannegger [6] compared his methodology (for 100 bootstraps at each node) to the CART algorithm and to the bagging method and found that the BNLS reduces the predictor error rate from 26% (the value obtained with CART) to 6%. The bagging method has the best performance with a prediction error of 3%. However the model obtained with this latter method is not easy to use and interpret by non-statisticians, unlike BNLS.
3.2 Random Survival Forests (RSF)

In order to stabilize the trees obtained by the CART procedure, Breiman [5] proposed the bagging method (so called for bootstrap aggregating) based on a family of random trees: multiple versions of predictor models obtained from bootstrapped samples are aggregated in order to construct a robust model estimator. Bagging was then adapted to the survival framework by Efron [24] and Akritas [25]. Another way to improve stability of trees is the boosting, developed by Freund and Schapire [26]. As for the bagging, the boosting consists in aggregating a family of models. Each tree is built iteratively from a weighted sample (an individual who is misclassified gains weight and an individual who is classified correctly loses weight) and then evaluated according to its ability to classify data. Considered as better than the bagging, the boosting is yet limited when data are too noised: the algorithm gives a heavy weight to noised data which leads to a bad overfit [27].

Breiman [5] proposed a random selection approach which combines the bagging method with a random selection of the covariates at each node of the tree. This methodology, called random forests, is more stable than the two previous ones [5, 26]. The method was adapted to the survival framework in an approach called “random survival forests” by Ishwaran [10]. Bootstrapped samples were drawn from the original data set. Notice that each bootstrapped sample excludes on average 37% of the data, a set called out-of-bag (OOB) data. For each bootstrapped sample \( b \) in \( (1, \ldots, n_{\text{tree}}) \), a survival tree is built: at each node, a subset of covariates is randomly selected among all the covariates. The splitting process continues under the constraint that each terminal node contains at least a fixed number of unique event times. The RSF algorithm computes an ensemble estimate for the cumulative hazard function (CHF), which is used as a predictor. For each terminal node \( h \), let \( t_{(1,h)} < \ldots < t_{(N_h,h)} \) be the distinct ordered uncensored event times, and define \( d_{l,h} \) and \( Y_{l,h} \) as the number of events and individuals at risk at time \( t_{(l,h)} \), for \( l = 1, \ldots, N_h \) respectively. The CHF estimator for the node \( h \) is the Nelson-Aalen estimator

\[
\hat{H}_h(t) = \sum_{t_{(l,h)} \leq t} \frac{d_{l,h}}{Y_{l,h}}.
\]

Each tree provides such a CHF estimate at each terminal node. Let \( \hat{H}_b(t|z) \) denote the cumulative hazard estimate for tree \( b \) conditionnaly on the covariate \( z \). To determine the CHF estimate for individual \( i \) with covariate vector \( z_i \) obtained from the tree \( b \), drop \( z_i \) down the tree. It will fall in a unique terminal node \( h \). So we have \( \hat{H}_b(t|z_i) = \hat{H}_h(t) \) if \( z_i \in h \). Let \( I_{i,b} = 1 \) if \( i \) is an OOB point for \( b \) and 0 otherwise. The OOB ensemble CHF estimator for \( i \) is:

\[
\hat{H}^*_e(t|z_i) = \frac{\sum_{b=1}^{n_{\text{tree}}} I_{i,b} \hat{H}_b(t|z_i)}{\sum_{b=1}^{n_{\text{tree}}} I_{i,b}}.
\]

Note that the above estimator is obtained by avering only over bootstrap samples in which \( i \) is excluded.
4 Comparison of the methods

In order to compare the different variable selection approaches, we apply them to two real data sets, the first concerning breast cancer and the second male fertility. The five following procedures have been compared: the bootstrap Cox stepwise procedure (BSS), the bootstrap Cox Lasso procedure (BLS), the bootstrap Cox randomized Lasso procedure (BRLS) with three different values of $\alpha$ (0.2, 0.4 and 0.6), the bootstrap node-level stabilization procedure (BNLS) and the random survival forest method (RSF).

4.1 Software

We use the methods implemented in R software (CRAN). We adapted the bootstrap to the stepwise Cox algorithm and to the R package penalized. We also modified the R package rpart in order to introduce bootstrap at node-level in the building of the tree (BNLS procedure). Instead of the default exp method which maximises an exponential likelihood, we used as a splitting criterion in rpart the logrank test statistic, following the recommendations of Radespiel-Troger et al. [28]. They showed that the prediction error of the tree is related to the instability of the covariate selection at each node. Compared to other splitting criteria on a real survival data set on gallbladder stones, the logrank test statistic yields the lowest prediction error (evaluated by the Brier score). The R package randomSurvivalForest provides four different splitting rules: logrank, conservation of events, logrank score, and approximate logrank (for more details, see Ishwaran and Kogalur [29]). We used the default splitting criterion i.e. the logrank test statistic. Concerning the number of bootstrapped samples, we take $N = 100$ for the Cox model based procedures, the default value $N = 1000$ for RSF, and $N = 1000$ for the BNLS procedure.

4.2 Comparison criteria

We compare the five procedures on the basis on their prediction performance but also on the usefulness of the model for non-statisticians. The prediction performance is measured by the prediction error rate. A model will be considered as useful if the statistical method involved is the most adapted to the problematic of the clinician and if the results are easy to interprete.

4.2.1 Comparison of the prediction error rates

We compare the stability of the five procedures using a statistical criterion: the prediction error rate, which allows to quantify the prediction performance of the final model selected. To this aim, the original data set is divided into two subsets, the training set and the test set. The procedures, computed on the training set, give a final model which is then
applied on the test set to calculate the prediction error rate. We obtain the prediction error rates in three steps:

1. We graphically chose sensible values of the cut-off $\kappa$ for the methods based on the Cox model and also values of $\lambda$ for BLS and BLRS. For that, considering the whole set, we plot the values of $\kappa$ for each selected covariable for BSS and we plot the values of $\kappa$ with respect to values of $\lambda$ for BLS and BLRS. The aim is to identify the most relevant variables whose frequencies of inclusion $\kappa$ are larger than those of the other covariates whatever the value of $\lambda$. Then we chose values of $\lambda$ and $\kappa$ which maximize the gap between the two subgroups of covariates. For BNLS, the choice of the optimal complexity parameter $cp$ is achieved by cross-validation techniques.

2. We then apply the procedures on the training set with the values of $\lambda$, $\kappa$ and $cp$ chosen at step 1. As far as the methods based on the Cox model are concerned, for each method, we adjust on the training set a simple Cox model including the subset of covariates for which the inclusion frequencies are greater than $\kappa$. This permits to obtain the estimations of the $\beta$ parameter in order to calculate the predicted outcomes.

3. We finally calculate the prediction error rates on the test set for each method.

The most commonly used prediction error rate for survival models is based on the Harrell’s concordance index $C$ [30]. To compute the $C$-index, the observed and predicted outcomes are compared, as follows:

- Form all possible pairs of observations.
- Eliminate those pairs where the shorter survival time is censored, and also pairs where the survival times and the censoring indicators are equal.
- For each permissible pair, count 1 if the shorter survival time has the worse predicted outcome and count 0.5 if the predicted outcomes are tied. Sum over all permissible pairs.
- The $C$-index is the ratio of the latter sum over the number of permissible pairs.

The prediction error rate is defined as $1 - C$ and is in $\{0, 1\}$. Note that a value of 0.5 is not better than random guessing.

For the Cox model based procedures, the prediction for an individual $i$ with covariate vector $Z_i$ is based on the linear predictor $\hat{\beta}'Z_i$. Individual $i$ has worse predicted outcome than $j$ if $\hat{\beta}'Z_i > \hat{\beta}'Z_j$. The predictions of the survival trees based procedures are derived from the CHF estimator. For the RSF procedure, they are derived from the OOB ensemble
CHF estimator of formula (2). Let \( t_1, \ldots, t_m \) be the unique times in the test set. Individual \( i \) is said to have a worse predicted outcome than \( j \) if

\[
\sum_{k=1}^{m} \hat{H}^*_e(t_k|Z_i) > \sum_{k=1}^{m} \hat{H}^*_e(t_k|Z_j).
\]

A total of 30 training sets and test sets were drawn to obtain a sample of error rates. Boxplots of the error rates are presented in order to compare the variability of the prediction performances.

### 4.2.2 Comparison of the selected covariates

We compare the set of covariates selected by the five different approaches applied on the entire set. For BSS, BLS and BRL, we take the values of the cut-off \( \kappa \) and of the penalties \( \lambda \) determined above. The BNLS algorithm provides the final tree which has been pruned with a fixed complexity parameter \( cp \). For the RSF procedure, the impact of the variables is measured by their “variable importance” (VIMP), which is the difference between the prediction error obtained with the original RSF procedure and the prediction error obtained using randomized assignments whenever a split for the considered variable is encountered [10].

### 4.3 Results for the breast cancer data set

#### 4.3.1 Description

The breast cancer data set contains the metastasis-free survival times from the study of Vijver et al. [31] who classified a serie of 295 patients with primary breast carcinomas as having a gene-expression signature associated with either a poor or a good prognosis. We restricted the study to the 144 patients who had lymph node positive disease. Vijver et al. [31] evaluated the predictive value of the gene-expression profile of patients for the 70 genes previously determined by Veer et al. [32] based on a supervised learning method. The data set can be found in the \texttt{R} package \texttt{penalized}. Five clinical risk factors and 70 gene expression measurements found to be prognostic for metastasis-free survival have been recorded. The censoring rate is 66%. The variables in the data set are:

- **time**: metastasis-free follow-up time,
- **event**: censoring indicator (1 = metastasis or death; 0 = censored),
- **diam**: diameter of the tumor (two levels),
- **N**: number of affected lymph nodes (two levels),
- **ER**: estrogen receptor status (two levels),
| Procedure | Mean | Standard deviation | Median |
|-----------|------|--------------------|--------|
| BLS       | 0.319| 0.066              | 0.310  |
| BRLS $\alpha = 0.2$ | 0.309| 0.075              | 0.292  |
| BRLS $\alpha = 0.4$ | 0.309| 0.081              | 0.287  |
| BRLS $\alpha = 0.6$ | 0.311| 0.086              | 0.302  |
| BNLS      | 0.376| 0.073              | 0.394  |
| RSF       | 0.279| 0.062              | 0.286  |
| TREE      | 0.389| 0.0741             | 0.382  |

Table 1: Mean, standard deviation and median of error rates for the different procedures: Bootstrap Lasso selection, Bootstrap Randomized Lasso selection (three values of $\alpha$), Bootstrap node-level selection, Random Survival Forest and a single survival tree.

- **grade**: grade of the tumor (three ordered levels),
- **age**: age of the patient at diagnosis,
- **TSPYL5...C20orf46**: gene expression measurements of 70 prognostic genes.

### 4.3.2 Prediction error rates

The Cox stepwise algorithm did not converge for the BSS procedure, probably due to too many covariates and not enough events in the boostrapped samples. As suggested by Meinshausen and Bühlmann [3], we search graphically the penalty $\lambda$ which leads to the best split between covariates in the BLS and BRLS procedures. However, the graphical determination of $\lambda$ is not so easy as in Meinshausen and Bühlmann [3] which presents results on an ad-hoc example. Figure 1 suggests that selecting between four and five covariates seems to be the most relevant whatever the value of $\lambda$ and $\kappa$. So we decide to choose a value of $\lambda = 0.4$ and a value of $\kappa = 0.2$ for BLS and BRLS whatever the value of $\alpha$. The complexity parameter $cp$ of the BNLS procedure is obtained by a ten-fold cross-validation procedure: we chose graphically $cp = 0.002$.

Figure 2 presents the boxplots for the different approaches based on 30 iterations and the numerical summaries are in table 1. As expected, RSF gives the lowest mean, median and standard deviation of the prediction error rates. We verify that the single survival tree has the worse prediction performance, but the BNLS procedure is hardly better. Whatever the value of $\alpha$, the BRLS procedure shows a large dispersion of error rates compared to the BLS procedure which has a standard deviation of 0.06, similar to this of RSF. However, a good compromise between dispersion and mean of error rates is observed for BRLS with $\alpha = 0.2$ or $\alpha = 0.4$. 


1. BLS

2. BRLS, $\alpha = 0.2$

3. BRLS, $\alpha = 0.4$

4. BRLS, $\alpha = 0.6$

Figure 1: Inclusion frequencies $\kappa$ with respect to $\lambda$ for Bootstrap Lasso Selection and Bootstrap Randomized Lasso Selection.
Figure 2: Boxplots of the prediction error rates for seven methods: Bootstrap Lasso selection, Bootstrap Randomized Lasso selection (three values of $\alpha$), Bootstrap node-level selection, Random Survival Forest and a simple survival tree.

4.3.3 Variables selected in the final model

Figure 3 shows the covariates selected by the Cox based procedures BLS and BRLS according to the choice of the inclusion frequency $\kappa$. Only 23 covariates of the 75 initial covariates appear in the selection done by the procedures applied to the bootstrapped samples (i.e. with $\kappa = 0$) and no clinical covariate is selected. For $\kappa = 0.2$, six covariates are selected by BLS whereas only four are selected by the BRLS procedures whatever the value of $\alpha$ (these four covariates are a subset of the previous six). We notice a clear gap between the four first selected variables and the other ones in the BRLS procedures. We can observe that the BLS and BRLS procedures find the same four most relevant covariates but not in the same order: PRC1, ZNF533, QSCN6L1 and IGFBP5.1.

For the tree-based procedures RSF and BNLS, figure 4 and 5 show that the same most discriminant variable is ZNF533. BNLS selects three other covariates among the first most important variables selected by RSF: COL4A2, PRC1 and N. The selection with a single survival tree procedure exhibits the same first discriminant variable ZNF533 followed by PRC1, RP5.860F19.3, HRASLS, IGFBP5, and SCUBE2. But the additional selected covariates by the CART algorithm never appear in the stable selection procedures. We can also notice that no clinical factor is included in the survival tree built by CART procedure unlike the BNLS and RSF procedures.

As far as the comparison of the Cox based methods and survival trees methods is
Figure 3: Selected covariates with respect to the inclusion frequency $\kappa$ for Bootstrap Lasso Selection and Bootstrap Randomized Lasso Selection.

1. BLS, $\lambda = 0.4$

2. BRLS, $\alpha = 0.2$, $\lambda = 0.4$

3. BRLS, $\alpha = 0.4$, $\lambda = 0.4$

4. BRLS, $\alpha = 0.6$, $\lambda = 0.2$
concerned, we notice similarity between the selected covariates: for example, \texttt{ZNF533} is found in all models. Regarding clinical factors, none was selected in Cox models contrary to the survival trees based procedures. As they can take into account interactions between covariates, tree based procedures show that \texttt{N}, the number of affected lymph nodes and the age of the patient at diagnostic are relevant predictors in metastasis-free survival. Notice that these two factors are also known to be clinically relevant.

4.4 Results for the fertility data set

4.4.1 Description

The fertility data were obtained from 2138 couples consulting for male infertility during the period from 2000 to 2004 at Toulouse Male Sterility Center (TMSC) located in University Public Hospital (France). Patients were followed from entry and during treatment by an andrologist specialist until either discontinuation of treatment or delivery of an alive infant. The maximum follow-up duration is 9 years. The outcome assessment was based on the delivery of an alive infant obtained at TMSC (pregnancies after medical treatment - medicine and/or surgical treatment, or assisted reproductive technologies (ART) - as well as spontaneous pregnancies). The event considered here is the birth of an alive infant and right-censored events correspond to miscarriage or loss to follow-up. The “survival” time is the delay in months from the first visit of the couple to the birth of its alive infant. We will work on the subset of the 1773 couples with covariates without missing values. 40% of the couples succeed in their parental project, leading to a censoring rate of 60%.
In agreement with clinicians, we decided to keep 32 covariates, among which:

- **for the man**: age, clinical investigation including medical histories as: histories of orchitis, epididymitis, urogenital infections (UI), sexually transmitted infection (STI), inguinal hernia, testis trauma, cryptorchidism (testicular maldescent), cancer, as well as testis, epididymis or vas deferens surgery, and clinical examination as: aspects of scrotum, testicular migration (presence of the testes into the scrotum and their position), epididymides and vasa deferentia evaluation, presence of a varicocele or hydrocele, testicular volumes, and if he received a non-ART treatment (both medical and surgical, including pharmacological and hormonal treatment, varicocelectomy, and vas deferens/epididymis surgery).

- **for the woman**: age, tubal factor, ovarian factor, cervical factor, ovulation, and if she received a non-ART treatment (hormonal treatment).

- **for the couple**: fecundity type (primary or secondary), infertility duration, type of ART including IUI (intra-uterine insemination), IVF (in vitro fertilization), ICSI (intracytoplasmic sperm injection) using male sperm cells, and ART with donor sperm.

4.4.2 Prediction error rates

As for the breast cancer data set, we chose $\lambda$ and $\kappa$ graphically. A first sight of figure 6 reveals that the contribution of the BRLS procedure with regard to the BLS procedure
| Procedure       | Mean  | Standard deviation | Median |
|-----------------|-------|--------------------|--------|
| BSS             | 0.453 | 0.017              | 0.454  |
| BLS             | 0.471 | 0.017              | 0.470  |
| BRLS $\alpha = 0.2$ | 0.483 | 0.013              | 0.482  |
| BRLS $\alpha = 0.4$ | 0.484 | 0.013              | 0.482  |
| BRLS $\alpha = 0.6$ | 0.482 | 0.015              | 0.480  |
| BNLS            | 0.447 | 0.021              | 0.446  |
| RSF             | 0.413 | 0.017              | 0.414  |
| STEP            | 0.451 | 0.015              | 0.450  |
| TREE            | 0.445 | 0.018              | 0.443  |

Table 2: Mean, standard deviation and median of error rates for the different procedures: Bootstrap Stepwise selection, Booststrap Lasso selection, Bootstrap Randomized Lasso selection (three values of $\alpha$), Bootstrap node-level selection, Random Survival Forest, Cox stepwise selection and a single survival tree.

for the choice of the cut-off values is not obvious. For BLS and BRLS, only two variables seem to be the most relevant factors whatever the value of $\lambda$. But for BLS and BRLS with $\alpha = 0.6$, five other covariates follow the two first when $\lambda = 0.4$, for a value of $\kappa = 0.3$ whereas for BRLS with $\alpha = 0.2$ and $\alpha = 0.4$, no group of variables seems relevant. So we decided to choose a fixed penalty $\lambda = 0.4$ for a value of $\kappa = 0.3$ for BLS and BRLS.

For the BSS procedure, we decide to choose a value of $\kappa = 0.5$ which seems to split the set of covariates in two distinct subgroups as suggested by figure 8. Thus, the first eight variables seem to be the most relevant to predict the duration to the birth. For the BNLS procedure, we observe by a 10-fold cross-validation procedure that the optimal value of the complexity parameter is $cp = 0.0035$.

The boxplots of the prediction error rates for the five procedures are presented in figure 7 and table 2 shows the summary statistics. Notice that the mean and median values obtained are higher (between 0.41 and 0.45) than those obtained from the data set on breast cancer which reflects the difficulty to predict the delay to the birth. However, the variabilities of the error rates are lower. As seen previously, the RSF method gives the best predictive model. However, it appears that RSF is similar in dispersion with other procedures. Moreover, we can notice that the BNLS procedure is not better than a single survival tree and the same remark can be done between the BSS procedure compared to a single stepwise Cox regression. These results can be explained by the size of the sample, which is sufficient to produce low error rates without bootstrapping. Regarding BLS and BRLS, these procedures show less variation in error rates but their means and medians are close to 0.5 which suggests that these models do no better prediction than random guessing.
1. BLS

2. BRLS, $\alpha = 0.2$

3. BRLS, $\alpha = 0.4$

4. BRLS, $\alpha = 0.6$

Figure 6: Inclusion frequencies $\kappa$ with respect to $\lambda$ for Bootstrap Lasso Selection and Bootstrap Randomized Lasso Selection.
4.4.3 Selected variables in the final model

If we compare the covariates selected by the different approaches, we can see in figure 8 that the first four selected covariates do not differ for the BLS and BRLS procedures for a value of $\kappa = 0.3$: we find tubal factor, IUI, sperm donor and infertility duration. The additional covariates included by BLS are epididymis, varicocelis, inguinal hernia, fecundity type, testicular trauma and testicular volume. For the BSS procedure, for a value of $\kappa = 0.5$, we find in the selected covariates the first four covariates selected by BLS but also female age, testicular volume, male treatment, varicocelis, testicular trauma, scrotum, female treatment and epididymis. We observe also that the BSS procedure includes more variables than the BLS and BRLS procedures, which leads to a lower prediction error rate.

We can notice that the cut-off value $\kappa = 0.3$ appears very clearly in the graphs of figure 8 for the BRLS procedures with $\alpha = 0.4$ and $\alpha = 0.6$, suggesting that the BRLS procedure is a good stable variable selection procedure. However, only four covariates are selected, which leads to poor prediction performances.

If we adjust a single Cox stepwise regression, we find the same covariates selected by the BSS procedure except for the variable scrotum. However, epididymis, testicular trauma and male treatment are not statistically significant at the 5% level in the Cox stepwise model.

Regarding the tree-based RSF and BNLS procedures, we observe in figures 8 and 9...
Figure 8: First part: Selected variables for Bootstrap Stepwise Selection, Bootstrap Lasso Selection, Bootstrap Randomized Lasso Selection and importance of variables for Random Survival Forest.
5. **BRLS** $\alpha = 0.6$, $\lambda = 0.2$

6. **RSF**

Figure 8: Second part: Selected variables for Bootstrap Stepwise Selection, Bootstrap Lasso Selection, Bootstrap Randomized Lasso Selection and importance of variables for Random Survival Forest.

Figure 9: The final tree obtained from the Bootstrap Node-Level selection
that the selected covariates are substantially different from the most relevant covariates selected by the Cox based procedures. The first split, which corresponds to female age, is the same for the two types of procedures but the following splits are different. This may be explained by the fact that the tree based procedures take into account interactions contrary to the Cox model based procedures. Moreover, the covariates selected for the BNLS procedure are found in the most important variables in RSF. The variables selected by a single survival tree are almost the same than those selected by BNLS, which can be explained by the large sample size of the fertility data set.

5 Discussion

5.1 Computational requirements

On a 2.4GHz processor, for the fertility data set with 1773 individuals and 32 covariates, running the RSF procedure is 200 times longer than running a single survival tree from the CART procedure (which takes about 1 second), whereas the stepwise algorithm takes about 10 seconds. Compared to the RSF method, the BLS and BRLS procedures have similar computing times. The BNLS procedure is much longer than RSF (4 times) as a bootstrap is realized at each node. Finally, the BSS procedure, for which a bootstrap sample is used at each stepwise, is the most expensive procedure in time (8 times longer than RSF). Thus, the running times seem reasonable for each procedure taken separately. However, a systematic comparison of errors becomes heavy, as well as the running time to exhibit the optimal complexity parameter \( cp \) in the cross-validation procedure for BNLS, and also the penalty \( \lambda \) for the BLS and BRLS methods.

5.2 Prediction error rates

Compared to the data set on breast cancer, the prediction error rates obtained from the fertility data set are higher, suggesting that data on fertility are more complex and difficult to study and that finding a good prediction model is a difficult task. Moreover, the sizes of the two data sets are very different, which could lead to observe discrepancy in the results. However, we find from the two data sets that the RSF procedure is the best procedure for prediction and that it allows to exhibit the most relevant variables to explain the survival durations. The Cox model and RSF have been previously compared by Omurlu et al. [33] using the Harrell’concordance index. On the basis on Monte Carlo simulations, they show that the Cox model has the best predictive performance whatever the size of the sample \((n = 50, 100, 250, 500)\). However, on a real data set on breast cancer, they find that RSF has the lowest prediction error rate. These contradictory results are not surprising if we consider the fact that the simulated data have been generated from a Cox model. Moreover, the RSF procedure is easy to use and does not require the choice of tuning values as do the BSS, BLS, BRLS and BNLS procedures. However, even if
the selected covariates are identified and sorted by their importance, as no final tree is
provided, the RSF results stay a black-box not easy to interpret and use for clinicians.

We find that the BRLS procedure, whatever the value of $\alpha$, does not seem to improve
the BLS procedure. Although the gap between the selected variables appears more clearly
for BRLS, the final model does explain these results. Moreover, the good results obtained
with this method by Meinshausen and Bühlmann [3] may come from the fact that they
presented an ad-hoc example. As noticed by these authors, we find on our data sets that
choosing a value of $\kappa$ equal to 1, as suggested by Bach [4], is too restrictive.

Even if some authors [12, 13] showed that the bootstrap method adapted to the
stepwise algorithm improves the stability of the variable selection, our results suggest
no improvement with the BSS procedure. As a matter of fact, the BSS algorithm does
not converge on the breast cancer data set containing too few events with respect to
the number of the covariates. On the other side, for the fertility data set whose size is
sufficient to assure the convergence of the algorithm, it does not give better results than
a single stepwise Cox model.

As far as the BNLS procedure is concerned, it seems not perform better than a single
survival tree on the two data sets, contrary to the results found by Dannegger [6]. It can
be explained by the fact that it is difficult to tune a sensitive value for the complexity
parameter by cross-validation. It may also result from the sufficient size of the fertility
data set.

5.3 Selected variables in the final model

The breast cancer data set was originally analyzed by Vijver et al. [31] who included in
a multivariate Cox regression model the clinical factors and a prognosis variable based
on the gene profile. Indeed, the 70 genes expressions were not included as independent
covariates in the Cox model, as we do in our analysis. We notice that the procedures
based on the Cox model selected no clinical factors in the breast cancer data set. It
can result from the fact that the genes which were introduced in the model are those
whose expression is the most correlated positively or negatively with the breast cancer
survival duration among the multitude of genes studied. Thus, these genes have a more
important weight in the multivariate analysis than clinical factors. In the BNLS and
RSF procedures, which take naturally into account interactions between covariates, two
clinical factors appears in the selected set of covariates. As it is not possible to include
interactions in the Cox model in a high-dimensional context, an alternative would be
to force the inclusion of some clinical factors clearly identified by the clinicians in the
procedures based on the Cox model.

Regarding the fertility data set, we obtain different final models with the procedures
based on the Cox model and those based on survival trees. However, all the selected
covariates are relevant clinical factors in reproductive issues. Indeed, the BSS, BNLS
and RSF procedures showed that female age is an important factor, with a cut-off value
of 34.5 years, which is similar to results observed in other studies [34 – 36]. Moreover, the infertility duration is selected by all the procedures, with a cut-off of 33.5 months in the BNLS method. It is not surprising as it is well-known that duration of infertility is predictive in the occurrence of pregnancy [37]. It is also interesting to notice that the BLS, BSS and BNLS procedures found a relationship between the live-birth and male clinical factors whereas BRLS included very few covariates which are essentially limited to female aspect and ART treatment. In most of the publications about reproductive issue in infertile couples, authors analysed the survival event of live-birth according only to female aspect and type of ART treatment [38 – 41]. However, we observe in our study that varicocelis is included in most of the procedures, which can be explained by the fact that varicocelis has an impact on an impaired spermatogenesis [42, 43]. The same observation could be done for the testicular volume which seems to be an important variable in the BNLS procedure, but with less impact in the other final models.

6 Concluding remarks

As far as the breast cancer data set is concerned, the objective is reasonably well achieved. The method based on randomized Lasso selection combined with bootstrapping provides prediction error rates very similar to those obtained by the RSF method, in association with an easily interpretable final model for the medical field. Regarding the procedure that combines bootstrap and survival trees, this method especially allows to identify interactions between the relevant covariates.

On the contrary, the data set on fertility seems to be much more difficult to analyze. The bootstrapping adapted to the various procedures based on the Cox model or on survival trees seems not improve the standard procedures (single Cox stepwise selection or single survival tree). Moreover, the wide variations observed in the prediction error rates may be due to the weak predictive performance of the selected covariates.

Finally, these results suggest that the Cox and tree based procedures should be performed in a complementary way to identify the most relevant covariates and provide to clinicians a stable and reliable model. Each procedure shows indeed a particular interest, either in terms of its prediction performance, either in the selection of the relevant covariates.
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