Confidence sets for model selection by $F$-testing

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Abstract

We introduce the notion of variable selection confidence set (VSCS) for linear regression based on $F$-testing. Our method identifies the most important variables in a principled way that goes beyond simply trusting the single lucky winner based on a model selection criterion. The VSCS extends the usual notion of confidence intervals to the variable selection problem: A VSCS is a set of regression models that contains the true model with a given level of confidence. Although the size of the VSCS properly reflects the model selection uncertainty, without specific assumptions on the true model, the VSCS is typically rather large (unless the number of predictors is small). As a solution, we advocate special attention to the set of lower boundary models (LBMs), which are the most parsimonious models that are not statistically significantly inferior to the full model at a given confidence level. Based on the LBMs, variable importance and measures of co-appearance importance of predictors can be naturally defined.

Key words and phrases: Confidence set, variable selection, linear regression.

1 Introduction

A statistical model can be interpreted as a story about how the data might have been generated by a particular random process. In many empirical analyses, a relevant question is: “Which story is the most plausible?” Sometimes, we are in the fortunate situation where the data strongly support one story, and so the corresponding model may be properly singled out as the “truth” for most purposes. More often than not, however, while we wish to select a single statistical model to describe the data, neither scientific nor statistical considerations can really help us to fix on a single model as the unique right story.

In the literature of model selection, this issue is sometimes referred to as model selection uncertainty (Chatfield, 1995, Draper, 1995, Hoeting et al., 1999, Yuan and Yang, 2005). A wealth of methods is available in the literature of statistics and machine learning for variable selection. However, often it is difficult to declare a single model as...
significantly superior to all possible competing models or even among the best set of models, due to the prevailing effect of model selection uncertainty. The methodology proposed in this paper is not meant to compete with existing model selection methods. Rather, it aims to characterize the intrinsic model selection uncertainty associated with the data at hand and also provide information on variable importance that goes beyond the standard single-final-model approach.

A well-established way to address model selection uncertainty is model averaging. It is now well understood that by weighting the candidate models properly, estimation or prediction can be much improved. See, e.g., Hoeting et al. (1999); Yang (2001); Hjort and Claeskens (2003) and references therein. In our view, a fundamental drawback from selecting a single model, which is not sufficiently dealt with by model averaging, is that when a single set of variables is chosen, a wealth of information is possibly thrown away in three key aspects. One is that alternative stories, possibly equally well supported, are ignored, which may be highly undesirable in terms of scientific understanding of the nature of the data. The second aspect is that it does not give any indication of how reliable the selected model is (in fact, uncertainty measures such as standard errors and confidence intervals based on the final model can be highly misleading). The third issue is that centering on a single model alone fails to provide trustworthy association among the predictors in jointly influencing the response variable.

This paper approaches variable selection from a different perspective by reducing the set of all possible collections of the variables to a smaller set, which we refer to as a variable selection confidence set (VSCS), which contains the true model with a given level of confidence. The methodology developed in this paper honestly reflects variable selection uncertainty. If the data are uninformative, distinguishing between models is difficult, and thus the VSCS may contain a large number of equally interesting models. Conversely, the VSCS tends to be much smaller in the presence of abundant information (when the sample size grows to infinity the VSCS essentially gives out the true model). Our approach goes as follows. We begin with a set of predictors with size smaller than the sample size, possibly after a variable screening or an initial variable selection. Then, we construct an exact VSCS based on $F$-tests. As mentioned already, such a confidence set can be very large. Even so, as will be seen, some sparse model selection methods sometimes produce a model not in the VSCS, in which case, one can be confident that the selected model is too sparse. Next, a very important subset of the VSCS is identified, which we call the lower boundary models (LBMs), defined as as the smallest models that are not statistically significantly inferior to the full model at a given confidence level. Dropping any term(s) in the LBMs would make the reduced model unfit from a hypothesis testing perspective. At the given confidence level, each model in the LBM set tells a well-justified, most parsimonious story. We show that the LBMs contain the information on the true predictors as $n \to \infty$; at the same time the LBMs are computationally more tractable. The set of LBMs can provide very useful information on: How many plausible stories are there to explain the data? Which predictors are definitely needed in most of the stories? Which predictors co-star in most of the stories? The answers to these questions based on the LBMs
provide a realistic and reliable perspective on the variable selection problem at hand.

The idea of confidence set for models has been explored. For example, Shimodaira (1998) advocates the use of a set of models that have AIC values close to the smallest among the candidates based on hypothesis testing. An important recent work is by Hansen et al. (2011) that proposes a notion of model confidence set in a framework that does not directly require the specification of the data generating model. Starting from a given set of models (loosely interpreted), they formulate an approach of an equivalence test on the currently remaining models followed by an elimination rule to remove the worst model when the equivalence test rejects the null hypothesis that the models are all equivalent under the loss of interest. This process is repeated until the equivalence test fails to reject the null. The whole procedure thus produces a subset of the original models that is meant to contain (or equal) the set of models with the best performance under the given loss function. They establish an asymptotic result, showing that this indeed happens at the target confidence level. Their approach is analogous to some step-down procedures for multiple hypothesis testing (e.g., Dudoit et al. (2003), Lehmann and Romano (2006), Ch. 9, or Romano and Wolf (2005)), as is mentioned in Hansen et al. (2011). Although we share the same general motivation with Hansen et al. (2011), the approach in this paper is completely different: We start with a strong linear model assumption and concretely build variable selection confidence sets. The focused framework offers a number of advantages. Firstly, we achieve exactly the specified coverage probability for the globally optimal model, which is difficult to obtain when starting from an arbitrary list of models as in Hansen et al. (2011). Secondly, in our setting the number of predictors, $p$, is allowed to grow with the sample size, $n$, and the number of potentially relevant models can thus be very large. In contrast, the approach of Hansen et al. (2011) is meant to handle only a fixed and small number of models to begin with. Thirdly, in addition to the confidence sets, our approach naturally leads to tools to assess model selection uncertainty (the multiple explanation index) for linear modeling, as well as importance of the predictors or their combination (inclusion and co-inclusion importance plots). Hansen et al. (2011) intend to have a much broader scope of applications (e.g., dealing with multiple forecasts). Not surprisingly, it requires several assumptions on the differences of the losses of the different models and on the behaviors of the equivalence tests and the elimination rules. A detailed discussion on these assumptions will be given in Section 6.

The rest of the paper is organized as follows. In Section 2, we present the exact VSCS based on $F$-tests. In Section 3, the subset of LBMs is defined, and the properties of the LBMs and variable importance measures are given. In Section 4, we illustrate the utility of our methods based on two real data sets. Simulation results are in Section 5. A discussion of the closely related work of Hansen et al. (2011) is in Section 6, followed by final remarks in Section 7. The proofs of the main theorems are deferred to a separate Appendix available online.
2 Exact confidence set

2.1 Setup

In this section, we construct an exact confidence set (ECS) in terms of coverage probability and study a related issue of detectability of the terms in the true model. Throughout the paper, we assume a normal regression model for the response variable:

\[ Y_i = \beta_0 + \sum_{j=1}^{p} \beta_j x_{j,i} + \epsilon_i, \quad i = 1, \ldots, n, \]  

(1)

where \( \epsilon_i \) are i.i.d. from \( N(0, \sigma^2) \), for some \( \sigma^2 > 0 \). The predictors are considered to be fixed and the intercept is always included. Some of the coefficients in \( (\beta_1, \ldots, \beta_p) \) are possibly zero. Let \( \gamma^* \) denote the set of indexes of all non-zero terms in the true mean expression. Our main interest is to construct a confidence set of models, \( \widehat{\Gamma} \), such that \( P(\gamma^* \in \widehat{\Gamma}) \geq 1 - \alpha \), for a given \( 0 < \alpha < 1 \). We will use \( \gamma \) as the index of a model, which corresponds to a collection of predictors.

2.2 Exact confidence sets based on F-testing

We use the familiar F-test to look for models that can plausibly be the true model. The full model is assumed to be uniquely fitted by least squares, which is typically appropriate when \( p \) is smaller than \( n \). When \( p \) is larger than \( n \), screening methods may be needed to reduce the number of predictors to be less than \( n \). When one uses the VSCS to assist a model selection method by providing additional information, a “full model” can be constructed to be a super model of the presently selected model by the method (see Section 4.2). Let \( \gamma_f \) denote the full model.

The well-known F-test compares the candidate model \( \gamma \) to the full model \( \gamma_f \). Particularly, \( \gamma \) is rejected when

\[ \widehat{F}(\gamma_f, \gamma) = \frac{(RSS_{\gamma} - RSS_{\gamma_f}) / (df_{\gamma} - df_{\gamma_f})}{RSS_{\gamma_f} / df_{\gamma_f}} > F_{\nu_1, \nu_2} (\alpha), \]  

(2)

where: \( RSS_{\gamma} \) and \( df_{\gamma} \) denote the usual residual sum of squares from fitting \( \gamma \) and the associated degrees of freedom; and \( F_{\nu_1, \nu_2} (\alpha) \) is the upper \( \alpha \) quantile of the F-distribution with \( \nu_1, \nu_2 \) degrees of freedom.

Considering all the subset models from the \( p \) predictors as the candidates models, the variable selection confidence set \( \widehat{\Gamma} \) is defined by the set of all these models that satisfy \( \widehat{F}(\gamma_f, \gamma) \leq F_{(df_{\gamma_f} - df_{\gamma_f}), \nu_f} (\alpha) \). By default, the full model is included in \( \widehat{\Gamma} \). The following result is a direct consequence of this way to construct \( \widehat{\Gamma} \).

**Theorem 2.1** Under the normal model, if the true model is not the full model, we have

\[ P \left( \gamma^* \in \widehat{\Gamma} \right) = 1 - \alpha. \]

When the true model is the full model, \( P \left( \gamma^* \in \widehat{\Gamma} \right) = 1. \)
The result follows trivially from the fact that when $\gamma = \gamma^*$, the $F$-statistic above has a $F(df_{\gamma}, df_{\gamma^*})$ distribution. We call $\hat{\Gamma}$ the exact confidence set (ECS).

The confidence set can be used to check if a given model (e.g., from a selection rule) is too parsimonious. A model in $\hat{\Gamma}$ is said to be $(1 - \alpha)$-SAFE (surviving against $F$-test evaluation). If a model is not $(1 - \alpha)$-SAFE, it most likely misses important predictors. As will be seen, models selected by some popular sparse model selection methods sometimes are not $(1 - \alpha)$-SAFE.

The simple VSCS basically has exact $1 - \alpha$ coverage probability, but its size needs to be discussed. First, note that the largeness of VSCS is necessary in general. Without any condition on the magnitudes of the effects of the predictors, to guarantee the coverage probability, we must include large models because one simply cannot tell apart two nested models between two situations: 1) both are correct (and thus the larger model should not be used); 2) the smaller model is wrong, but the extra terms in the larger one are tiny relative to the sample size. Therefore, $\hat{\Gamma}$ cannot be improved without further conditions on signal strength. Second, from a practical perspective, this VSCS can be too large to be directly very useful beyond checking a model suspected of being overly parsimonious. Therefore, a special subset of the ECS $\hat{\Gamma}$ will be considered in the next section.

### 2.3 ECS after screening or a conservative selection

In various applications, $p$ is larger than $n$. Methods such as Lasso and Scad can still be applied to obtain a sparse model with a relatively small number of predictors. An important issue then is to examine the reliability of the selected model. In this context, VSCS can provide a complementary perspective on which variables and models may be important and thus serve as a tool to support or cast doubt on the selected model.

To construct a VSCS when $p > n$, a variable screening method can be used to sift out unimportant variables and reduce the number of predictors for further consideration to be less than the sample size. In the literature, several screening methods have been proposed with theoretical justifications (see, e.g., Fan and Lv (2008) and Fan and Song (2010)).

Consider a variable screening method $\psi$ that yields a reduced collection of the original predictors, denoted by $\Omega(\psi)$, of size at most $n - 1$. The size is typically substantially smaller than $n$, say of a smaller order. For example, in the sure independence screening procedure of Fan and Lv (2008) based on marginal correlations, the prescribed size of $\Omega(\psi)$ is $d_n = O(n/\log(n))$. Treating $\Omega(\psi)$ as the full model, we can find the ECS as described in the previous subsection and denote it by $\hat{\Gamma}_{\Omega(\psi)}$.

Alternatively, we may consider a set of $L$ high-dimensional model selection methods $\Psi = \{\psi_1, ..., \psi_L\}$ that each produces a model with a choice of a tuning parameter. For our purpose, the tuning parameter for each method is chosen conservatively so that the selected model is more likely to not miss the true predictors (but may include noise variables at the same time). The set $\Psi$ (with the tuning parameters) is said to be collectively over-consistent.
if with probability going to 1 the union of the sets of predictors in the selected models by the $L$ methods, denoted by $\Omega(\Psi)$, contains all the predictors in the true model. Clearly, if any of the model selection methods is actually consistent or over-consistent in selection, then $\Psi$ is collectively over-consistent, but the reverse is not true. Hence the condition is much milder than demanding at least one of the methods to be consistent. Let $\hat{\Gamma}_{\Omega(\Psi)}$ denote the ECS based on $\Omega(\Psi)$ as the full model (assumed to be of size less than $n$).

For the result below, we assume that the screening or pre-selection by $\Psi$ is done based on a side data set (e.g., from a previous study) or using a small part of the present data. In real applications, when the sample size is small and there is no side data, variable screening may be done with the full data as done in Fan and Lv (2008), although there might be a bias due to reuse of the same data for both steps (screening and VSCS construction).

**Corollary 2.2** Assume that the screening method $\psi$ satisfies that $\Omega(\psi)$ contains all the variables in $\gamma^*$ with probability going to 1; or $\Psi$ is collectively over-consistent. Then we have

$$\liminf_{n \to \infty} P \left( \gamma^* \in \hat{\Gamma}_{\Omega(\psi)} \right) \geq 1 - \alpha,$$

where $\Omega$ is either $\Omega(\psi)$ or $\Omega(\Psi)$ under the respective condition.

From the above corollary, VSCS can be reliably constructed after an effective variable screening or conservative selection.

### 2.4 Detectability Conditions

Here we try to understand the conditions by which the terms in the true model will eventually not be missed in the models in the ECS. Let $\gamma$ denote a model that misses at least one true term. The $F$-statistic $\hat{F}(\gamma_f, \gamma)$ has a non-central $F$-distribution $F_{(df_{\gamma_f} - df_{\gamma}), df_{\gamma}, \delta_{\gamma}}$, where $\delta_{\gamma}$ is the non-centrality parameter summarizing the overall effect from missing one or more terms. A VSCS method is said to asymptotically detect all the true terms if all the true terms are included in each of the models in the confidence set with probability going to 1. Note that asymptotic detectability does not address the issue of inclusion of unnecessary terms. For the following result, the number of predictors $p$ is allowed to depend on $n$ and thus will be denoted by $p_n$. We assume that $p_n$ is bounded away from $n$ in the sense that $p_n \leq (1 - \varepsilon)n$ for some possibly small $0 < \varepsilon < 1$. Let $p_0$ denote the number of terms in the true model, which is assumed to satisfy that $\log p_0$ is of order $\log n$ and $p_0/p_n \to 0$.

**Theorem 2.3** Let $\Gamma_u$ denote the set of models that miss at least one of the true terms. For the ECS, a necessary condition for asymptotic detectability of the true terms at each $0 < \alpha < 1$ is

$$\min_{\gamma \in \Gamma_u} \frac{\delta_{\gamma}}{\sqrt{df_{\gamma_f} - df_{\gamma}}} \to \infty \text{ as } n \to \infty.$$
Moreover, the true terms are asymptotically detectable if

\[
\min_{\gamma \in \Gamma_u} \delta_{\gamma} \leq \frac{\xi_n + \sqrt{\left(\right)}}{\left(1 + \log \frac{p_n}{df_{\gamma} - df_{\gamma'}}\right)}
\]

is no less than a large enough positive constant for some arbitrarily slowly increasing sequence of \(\xi_n \to \infty\). Furthermore, this condition cannot be generally improved in the sense that there is a setting with \(p_n \to \infty\) such that the condition is necessary for the true terms to be asymptotically detectable.

From the above, when the true model is strong (i.e., each term in the true model is important in its unique contribution for explaining the regression function) in the sense that removing any term(s) in the true model would result in that \(\delta_{\gamma}\) is much larger than \(\sqrt{(df_{\gamma} - df_{\gamma'}) \left(1 + \log \frac{p_n}{df_{\gamma} - df_{\gamma'}}\right)}\), the true predictors should appear in all the ECS models with probability approaching 1. Both the necessary and the sufficient conditions typically hold in the traditional case that the list of models is fixed (which contains the true model), while \(n \to \infty\). They basically mean that each true coefficient in absolute value (assuming that the predictors are more or less standardized) is of an order larger than \(1/\sqrt{n}\), which is just as expected. Note that there is a small gap (at the order of a logarithmic term in \(p_n\)) between the sufficient and necessary conditions in the theorem in case of \(p_n \to \infty\) (although the term \(\xi_n\) is technically needed when the other term in the denominator happens to stay bounded, since it is allowed to approach \(\infty\) arbitrarily slow, it can be basically ignored in the discussion). As will be shown in the proof of the theorem, we can construct a setting where the extra logarithmic term is necessary. Therefore the sufficient condition in the theorem is not generally improvable.

The result confirms an intuitive understanding regarding the effect of \(p_n\). Consider a proper subset model of the true model with at least one true term missing. Under the usual assumption that the true predictors are normalized and not highly correlated, the detectability condition implies that \(n \beta^2 / (\sqrt{p_n} - p_0 \log(p_n)) \to \infty\), where \(\beta\) is any true coefficient. If the true model is sparse and \(p_n\) is only of a slightly larger order than \(p_0\), then this condition is very mild. In contrast, if the full model is too large so that \(p\) is of order \(n\), even if the true model is sparse, the true coefficients have to be much larger to ensure the detectability of all the true terms. Therefore, one should avoid using a really large full model, if possible. This understanding can be exploited to derive empirical rules to construct a “full model” based on a high-dimensional model selection method so as to gain more insight than offered by the selected model alone (see Section 4.2 for an example).

3 The subset of lower boundary models (LBMs)

Let \(\gamma\) be a model in a confidence set \(\hat{\Gamma}\). We say that \(\gamma\) is a lower boundary model if there is no model in \(\hat{\Gamma}\) that is nested within \(\gamma\). Let \(LBM(\hat{\Gamma})\) denote the set of all lower boundary models. From Theorem 2.1, with probability
at least $1 - \alpha$, the true model is a LBM or it contains at least one LBM (as its subset model). Therefore, the set of the lower boundary models can naturally serve as a tool to check if a selected model is over-simplifying (or worse): If it is not on the lower boundary or above, we can confidently say that the model has missed important predictors and we have an idea of what they are. Since some sparse model selection methods are sometimes overly aggressive in finding a parsimonious model (which is not necessarily bad for prediction), for the purpose of model identification beyond predictive performance, an objective check as aforementioned can be very helpful to avoid consequences of a decision based on an excessively simplified description of the data.

When the true model is weak (relative to the sample size and the error variance), $LBM(\hat{\Gamma})$ may involve noise variables. In this case, while it is not quite feasible to identify the true model, all the variables that appear in at least one of the models in $LBM(\hat{\Gamma})$ are justified as necessary at the $1 - \alpha$ confidence level in terms of the need to include them in at least one model that is not statistically significantly different from the full model. When the signal is strong, however, we have the following result.

**Corollary 3.1** Assume that the ECS asymptotically detects all the true terms. As $n \to \infty$, if $\gamma^*$ is not the full model, then $P(LBM(\hat{\Gamma}) = \{\gamma^*\}) \to 1 - \alpha$; if $\gamma^*$ is the full model, then $P(LBM(\hat{\Gamma}) = \{\gamma^*\}) \to 1$.

From the above result, for a large sample size, when the true terms are asymptotically detectable, the true model will be the only LBM at the given confidence level. Also, all the useful variables will not be missed in the LBMs with probability close to one.

As mentioned already, when there is no condition at all on the magnitude of the non-zero coefficients, to guarantee the coverage probability, one has no power at all to reject a larger model when two nested models are compared. Therefore, when constructing a VSCS, it is natural to require that if a model is included in $\hat{\Gamma}$, then any larger model is also included. We call such a confidence set expansive. Note that the ECS in the previous subsection is not necessarily always expansive.

Clearly, for an expansive confidence set, all we need to know is the set of lower boundary models. The characteristics of the LBMs can be very informative regarding the roles of the predictors. We discuss several scenarios below.

1. $LBM(\hat{\Gamma})$ is unique (the simplest case): This is a situation where all the predictors in the unique model are clearly important and none of the other predictors is proven to be necessary with the limited information presently available. When the sample size gets much larger, $LBM(\hat{\Gamma})$ may involve more predictors.

2. The size of $LBM(\hat{\Gamma})$ is larger than 1, but small: One sub-scenario is that the models in $LBM(\hat{\Gamma})$ differ only in one or two predictors, in which case the common predictors in the LBMs are clearly important and several predictors are useful but we do not know which one is the best. Another sub-scenario is that the LBMs are
quite different in terms of variable composition, which indicates that various combinations of the predictors can give similar explanation power of the response variable.

3. The size of $LBM(\hat{\Gamma})$ is moderate: This can happen when the number of predictors is not small and a number of predictors are moderately or highly correlated.

4. The size of $LBM(\hat{\Gamma})$ is relatively large: For high-dimensional cases, this may be typical and one cannot realistically find the “true” or best model. Any model selection rule is more or less randomly picking out a model among many possibilities that have similar criterion values.

### 3.1 A Multiple-Explanation Index and Inclusion Importance

Based on the LBMs, we propose some quantities that can be useful for measuring the degree to which multiple models seem to explain the data well and also the importance of a variable. For a set $A$, $|A|$ denotes the size of the set. For a given predictor $x_i$, let $K(x_i)$ be the number of times that $x_i$ appears in the models in $LBM(\hat{\Gamma})$.

**Definition 3.2** The $(1 - \alpha)$-multiple-explanation index (MEI) is

$$MEI = \log |LBM(\hat{\Gamma})|.$$  

The MEI can be as large as the logarithm of the combinatorial number of $p$ choose $\lfloor p/2 \rfloor$, which is roughly $\frac{p}{2} \log(2e)$. Clearly, when $MEI = 0$, there is a single model in $LBM(\hat{\Gamma})$. Basically, MEI describes (on a log-scale) how many most-parsimonious models there are to explain the data at the given confidence level.

**Definition 3.3** The $(1 - \alpha)$-inclusion importance of a predictor $x_j$ is

$$II(x_j) = \frac{K(x_j)}{|LBM(\hat{\Gamma})|}.$$ 

Note that for a predictor that appears in all models in $LBM(\hat{\Gamma})$, its $II$ value is 1. For a predictor that appears in some models in $LBM(\hat{\Gamma})$ but not in others, its inclusion importance is decreased. For such a variable, although we cannot say that it is in the true model confidently, at least one LBM suggests that it is possible. For variables that do not appear in any of the models in $LBM(\hat{\Gamma})$, their $II$ values are 0, which means that at the $1 - \alpha$ level, not enough evidence shows that such a term may be definitely needed. Of course, a variable with $II = 0$ should not be declared useless: only that there is not enough evidence to support that it is useful at the time being.

**Corollary 3.4** Assume that the ECS asymptotically detects all the true terms. Then we have $\liminf_{n \to \infty} P(MEI = 0) \geq 1 - \alpha$ and $\lim_{n \to \infty} P(II(x_j) = 1) = 1$ for all $x_j$ in the true model and $\lim_{n \to \infty} P(II(x_j) > 0) \leq \alpha$ for all $x_j$ not in the true model.
From the above result, in the most positive case that we have sufficient information to learn the true model, MEI is zero. In the toughest case where the signal is weak enough, $II(x_j)$ can be very small for the true variables, as is expected.

In the rest of the paper, we also consider inclusion importance based on the entire confidence set $\hat{\Gamma}$, defined by $\tilde{II} = \tilde{K}(x_j)/|\hat{\Gamma}|$, where the function $\tilde{K}(x_j)$ is the number of times that $x_j$ appears in the models of $\hat{\Gamma}$. When a smaller model is included in $\hat{\Gamma}$, also larger models containing the same predictors plus some other variables tend to be included as well by construction. Therefore, differently from $II(x_j)$ based on the lower boundary set, a predictor cannot be declared relevant just because it appears in $\hat{\Gamma}$. Actually, unimportant predictors tend to have $\tilde{II}$ close to 1/2 (because when expanding from the lower boundary models, given the other added predictors, the predictor being examined may or may not be included). This will be seen clearly from our numerical examples in Sections 4 and 5.

### 3.2 Importance profile and co-importance of predictors

Let $\hat{\Gamma}_\alpha$ and $LBM(\hat{\Gamma}_\alpha)$ denote the $1-\alpha$ confidence set ECS, $\hat{\Gamma}$, and the corresponding lower boundary set, $LBM(\hat{\Gamma})$, respectively. Clearly, when $\alpha \to 1$, $LBM(\hat{\Gamma}_\alpha)$ degenerates to the full model and when $\alpha \to 0$, $LBM(\hat{\Gamma}_\alpha)$ becomes the set of all single term (predictor) models unless some predictor vectors are perfectly orthogonal to the response vector (or so after removing the effects of some other predictors). Tracing the LBMs as $\alpha$ changes between these two extremes can be informative.

We introduce two new graphical tools to study the explanatory role of predictors. The first tool is the predictor $II$ profile plot, which traces the inclusion importance, $II(x_j)$ or $\tilde{II}(x_j)$ of all (or some) predictors, against $\alpha$. By looking at the profile plot, one can inspect whether one or more predictors become more important as the confidence level changes. For instance, a sharp and steady increase in $II$ when the confidence level changes from 99.9% to 95% would suggest that the predictor is highly relevant and should not be missed (see Figures 1 (a) and 3 (a)).

The second tool is the co-inclusion importance ($CII$) plot. The $CII$ plot displays the co-importance of variable pairs $\{x_j, x_k\}$, $j, k = 1, \ldots, p$. Let $K(x_j, x_k)$ denote the number of models in the $LBM(\hat{\Gamma}_\alpha)$ including both $x_j$ and $x_k$. The co-importance of $x_j$ and $x_k$ is defined by

$$CII(x_j, x_k) = \frac{K(x_j, x_k)}{K(x_j) + K(x_k) - K(x_j, x_k)}$$

if $K(x_j, x_k) > 0$, and $CII(x_j, x_k) = 0$ if $K(x_j, x_k) = 0$. Note that the denominator counts all the models in $LBM(\hat{\Gamma}_\alpha)$ that include either $x_j$ or $x_k$. Thus, clearly from the above definition we have $0 \leq CII(x_j, x_k) \leq 1$, where the case $CII(x_j, x_k) = 1$ occurs if whenever one of $x_j$ and $x_k$ appears in a model in $LBM(\hat{\Gamma}_\alpha)$, so does the
other. For the example of genetic data in Section 4.2, we display co-inclusion importance using a graph display where the nodes represent variables and the thickness of the edges is proportional to $CII$ values (see Figures 1 (b) and 3 (b)).

It is well-known that high correlations between predictors complicate the correct selection of the true predictors. Model selection methods often exclude predictors that are highly correlated with ones that are already in a model, whether or not they should be included from a different angle. Some methods such as Elastic Net (Zou and Hastie, 2005) are proposed to alleviate the problem. The examination of the LBMs can offer insight on the question of whether two predictors should co-appear or not. Although we focused on co-inclusion of two predictors, the idea also works for a set of three predictors or more, although a graphical representation becomes difficult.

4 Real data examples

4.1 Prostate Cancer Data

To illustrate our method, we consider the benchmark data set from a study of prostate cancer (Stamey et al., 1989). Tibshirani (1996), Zou and Hastie (2005), Li and Lin (2010) studied these data among other authors in the model selection literature. The predictors are eight clinical measures: log(cancer volume) (lcavol), log(prostate weight) (lweight), age, the logarithm of the amount of benign prostatic hyperplasia (lbph), seminal vesicle invasion (svi), log(capsular penetration) (lcp), Gleason score (gleason) and percentage Gleason score 4 or 5 (pgg45). The response is the logarithm of prostate-specific antigen (lpsa). In Table 1, we show size and relative frequency of the predictors for the ECSs at the 95, 99 and 99.9% confidence levels. The size of the ECS is clearly monotone in $\alpha$.

At the 99 and 99.9% levels, lcavol, lweight, lbph and svi appear in more than half of the sets, yielding only 2 and 3 LBMs respectively. At the 95% confidence level, lcavol, lweight and svi appear in all the models in the ECS and there is a single LBM containing these predictors.

<table>
<thead>
<tr>
<th>$(1 - \alpha)$%</th>
<th>ECS size</th>
<th>MEI</th>
<th>lcavol</th>
<th>lweight</th>
<th>age</th>
<th>lbph</th>
<th>svi</th>
<th>lcp</th>
<th>gleason</th>
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<td>1.00</td>
<td>0.74</td>
<td>0.48</td>
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<td>0.79</td>
<td>0.43</td>
<td>0.40</td>
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</tr>
<tr>
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<td>1.00</td>
<td>1.00</td>
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<td>0.50</td>
<td>1.00</td>
<td>0.50</td>
<td>0.50</td>
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</tr>
</tbody>
</table>

Table 1: Exact confidence sets (ECSs) for the prostate cancer data. The columns represent confidence level $(1 - \alpha)$%, size of $\hat{\Gamma}$ (ECS size), multiple explanation index (MEI), and relative frequency of the predictors in the ECS (columns 4-11).

In Table 2, we show the lower boundary models for the 95 and 99% confidence levels, and models selected using the following methods: AIC and BIC, Lasso and Scad (Fan and Li, 2001). To compute Lasso and Scad we use the R package *ncvreg* (available at http://cran.r-project.org). The tuning parameters for Lasso and Scad are chosen by 10-fold cross validation. In this example, all the considered selection procedures turned out to be SAFE at
the 95% confidence level, since the selected models were found in the exact confidence set. Our \( II \) statistic shows 4 variables appearing at least once in the lower boundary at the 99% confidence level (lca\( v \)ol, l\( w \)eight, lbp\( h \) and svi), suggesting that such variables are indeed relevant. At the 95% confidence level, only lca\( v \)ol, l\( w \)eight and svi are relevant for a parsimonious story of the underlying process. The other selection methods tend to agree on the importance of these variables.

\[ \alpha = 0.01 \]

\begin{array}{cccccccc}
\text{Term} & \text{LBMs} & II & \text{LBMs} & II & \text{AIC} & \text{BIC} & \text{Lasso} & \text{Scad} \\
\hline
\text{lca\( v \)ol} & 1 & 1 & 1.00 & 1 & 1.00 & 1 & 1 & 1 & 1 \\
l\( w \)eight & 1 & 0 & 0.50 & 1 & 1.00 & 1 & 1 & 1 & 1 \\
age & 0 & 0 & 0.00 & 0 & 0.00 & 1 & 0 & 0 & 1 \\
lbp\( h \) & 0 & 1 & 0.50 & 0 & 0.00 & 1 & 0 & 1 & 1 \\
svi & 0 & 1 & 0.50 & 1 & 1.00 & 1 & 1 & 1 & 1 \\
lcp & 0 & 0 & 0.00 & 0 & 0.00 & 0 & 0 & 0 & 0 \\
gleason & 0 & 0 & 0.00 & 0 & 0.00 & 0 & 0 & 0 & 0 \\
pgg45 & 0 & 0 & 0.00 & 0 & 0.00 & 0 & 0 & 1 & 1 \\
\end{array}

Table 2: Lower boundary models and model selection for the prostate cancer data. We list the lower boundary models (LBMs) (1=predictor included, 0=predictor not included) computed for \( \alpha = 0.01, 0.05 \), and variable inclusion importance (\( II \)) for each predictor. The last columns show the models selected using AIC, BIC, Lasso and Scad (1=predictor included, 0=predictor not included); For the AIC and BIC we used exhaustive search for all possible models; for Lasso and Scad we used 10-fold cross validated tuning parameters.

In Figure 1 (a), we show the inclusion importance profiles for the variables computed at the 95% level. Besides the models in the lower boundary, the ECS also includes all the models obtained by expanding from the lower boundary models; therefore, for the ECS we have \( \widetilde{II} = 1/2 \) for \( \alpha \) sufficiently close to 0. As \( \alpha \) increases, we observe different behaviors of the predictors. The importance of lca\( v \)ol, l\( w \)eight, and svi increases rapidly as \( \alpha \) grows reaching the limit value \( \widetilde{II} = 1 \), for \( \alpha \) larger than 0.035. In contrast, the importance of age, lbp\( h \), lcp, gleason and pgg45 converges to 0.5, meaning that we have no sufficient information to declare such variables important when \( \alpha \) gets larger than 0.035. When \( \alpha \) is between 0 and 0.035, lbp\( h \) and pgg45 appear to be moderately relevant. In Figure 1 (b), we show the co-inclusion importance graph for the variables computed at the 99% level. The nodes correspond to individual variables, while the thickness of the edges is proportional to the co-inclusion importance statistic, \( CII \), defined in Section 3.2. The graph emphasizes pairwise occurrence of variables lca\( v \)ol, l\( w \)eight, svi and lbp\( h \) in the lower boundary. It suggests that at the 99% level, in terms of explaining the variability in the prostate-specific antigen most parsimoniously, svi and lbp\( h \) appear together, and they serve as an alternative to l\( w \)eight.

### 4.2 Bardet-Biedl syndrome genetic data

We apply our methods to gene expression data from the micro-array experiments of mammalian eye tissue of 120 twelve-week-old male rats (Scheetz et al., 2006). The outcome of interest is the expression of TRIM32, a gene
which has been shown to cause Bardet-Biedl syndrome (Chiang et al., 2006), a genetic disease of multiple organ systems, including the retina. The micro-arrays contain over 31,042 different probe sets. For each probe, gene expression is measured on a logarithmic scale. Following the pre-processing steps in (Huang et al., 2008, Scheetz et al., 2006), we selected 18,976 of the 31,042 probe sets on the array as they “exhibited sufficient signal for reliable analysis and at least 2-fold variation in expression”; then we restricted our attention to the 3,000 probes with the largest variance.

Example 1: Marginal correlation and Lasso screening. Here we consider statistical screening, which is routinely applied on micro-array data when no biological hypothesis is available. Following Huang et al. (2008), we selected 200 variables with the strongest correlation with TRIM32; then we used penalized regression to select a smaller subset of predictors. For illustration purposes, we considered the Lasso method to carry out the latter step using the R package ncvreg. We computed models \( \Gamma = \{ \gamma_1, \ldots, \gamma_{100} \} \) along the Lasso solution path corresponding to a grid of 100 Lasso regularization parameters and selected the best model \( \hat{\gamma}^* \in \Gamma \) using 5-fold cross-validation consisting of 18 predictors. We then built the “full model” \( \gamma_f \) by moving along the lasso path and taking the largest model on the path with the number of predictors \( \tilde{p} \) such that

\[
\frac{\hat{\delta}_{\text{max}}}{\sqrt{\tilde{p} - p^* + 1)}(1 + \log \tilde{p} - \log (\tilde{p} - p^* + 1)) > C
\]

where \( \hat{\delta} = \max_j \{ \hat{t}_j^2 \} \) is an estimated upper-bound for the non-centrality parameter when one term is missing, \( \hat{t}_j \).
The $t$-statistics for the individual variables in the lasso model and $C$ is a constant representing the necessary signal-to-noise ratio to detect the true terms in the sense of Theorem 2.3. The left hand side in (4) represents an approximated upper bound to the detectability condition in Theorem 2.3; for example, with $C = 3$, we end up with a full model with 21 predictors. The rationale for the above choice of full model is that if we are to trust the lasso model at all, using a larger full model than given above may even make the strongest term in the lasso model undetectable.

In Figure 2 (a), we show the $p$-values corresponding to the $F$-test comparisons between the full model against the candidate sub-models along the Lasso path. The upper region in the plot contains 95%-SAFE models along the Lasso path, while the bottom part of the plot contains models that are unsafe due to the overly aggressive Lasso selections, which miss one or more important variables. Of particular interest are the model closest to the boundary (circled on in Figure 2 (a)), since they represent the most parsimonious Lasso path model within the ECS. Such models include the following probes: 1370429_at, 1374106_at, 1379971_at, 1383110_at, 1383673_at, 1383996_at, and 1389584_at, which alone explain approximately 70% of the variability in TRIM32. The probes selected by such parsimonious models overlap with the selection obtained by the adaptive Lasso and adaptive Scad methods described Huang et al. (2008). The best fitting ECS model on the Lasso path (also circled in Figure 2 (a)) accounts for 74% of the variability in TRIM32, but contains almost twice as many variables. In Figure 2 (b) we show the inclusion importance of predictors at the 95% confidence level. It shows that except two predictors that appear on at least 40% of the LBMs, the other predictors have rather low II values, which reflects the fact that there are many roughly equally plausible models with different compositions of the predictors that can explain TRIM32. From the plot, we may need to admit that the task of identifying the best model at the current sample size is just infeasible.

Example 2: Biological screening. In this example, we consider as potential predictors expression in 11 probes with significant linkage to the known retinal disease genes Bbs1, Bbs4, Bbs8, Opn1sw, Pedh15, Pde6a, Pex1, Pex7, Rdh12 and Rdp4 in (Sheetz et al., 2006) (probe ids 1384603_at, 1383417_at, 1383007_at, 1378416_at, 1388025_at, 1378408_at, 1393426_at, 1376595_at, 1379784_at, 1382949_at, 1371762_at, ). Figure 3 (a) shows the marginal inclusion importance profile plot for $\alpha$ ranging from 0 to 0.1. The most important genes appear to be Bbs8, Bbs4, Pex7 and Opn1sw for all considered confidence levels. We remark that Bbs4 and Bbs8 are known to be related to the Bardet-Biedl syndrome, since they belong to the so-called BBS group. Also Opn1sw is reputed as important since it represents a non-contiguously regulated gene encoding proteins related to the disease. Figure 3 (b) shows the co-inclusion importance graph, where the thickness of the edges represents values of the co-inclusion importance statistic $CII$ defined in Section (3.2) (edges corresponding to $CII \leq 0.2$ are omitted for clarity). It offers information unavailable in the marginal $\widetilde{II}$ plot or from an usual model selection process. The totally isolated predictors in the graph are weak on their own and also do not appear to have any potential jointly with another
Figure 2: SAFE models on the Lasso path: (a) p-values based on the F-test of the full model $\gamma_f$ with 21 predictors against smaller models on the Lasso path; the models above the line are safe at the 95% confidence level. (b) Inclusion importance of predictors (II) computed from the lower boundary models.

There is some evidence to support that Rdh12, Opn1sw and Pex1 are useful, which is already seen from Figure 3 (a). But Figure 3 (b) further shows that Rdh12 and Pex1 tend to influence the response by appearing together, but they are not connected with Opn1sw, meaning that $\{\text{Rdh12, Pex1}\}$ and Opn1sw have competing (rather than synergistic) effects. Note also that while Bbs4 and Opn1sw have the same high II value (0.62), their CI value is very small (the connection is very light), which says that their effects are redundant if appearing together. Such information may help the biologist to gain more insight on the problem.

In Table 3, we show lower boundary models at the 95% confidence (models LBM1–LBM8), and inclusion importance statistics. For comparison purposes, we also report 5-fold cross-validated Lasso, Scad and Mcp models selected. Due to pronounced noise in the data, Lasso, Scad and Mcp generate quite different models for different cross-validation runs; for illustration purposes, we show a single instance. We also show the AIC and BIC models computed by exhaustive search. For each model we report p-values from the F-test defined in Section 2.2 and $R^2$ obtained from a ordinary least square fit. The lower boundary models contain 4 to 6 variables emphasizing various combinations of predictors equally useful in explaining TRIM32. All the LBMs give $R^2$ near 50%, while the full model with 11 variables yields $R^2 = 55\%$. Genes Bbs4, Bbs8, Opn1sw and Pex7 are included in most of LBMs. The same genes also appear frequently in the other selected models. Note that the Mcp and BIC models fall outside the confidence set, so those models cannot be trusted at the 95% confidence level. This is not surprising since BIC and Mcp criteria are known to generating overly sparse selections, so here they are likely missing at least one important variable.
<table>
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<tr>
<th></th>
<th>Abca1</th>
<th>Bbs1</th>
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<th>Bbs8</th>
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<th>sw</th>
<th>Podh15</th>
<th>Pde6a</th>
<th>Pex1</th>
<th>Pex7</th>
<th>Rdh12</th>
<th>Rdpl1</th>
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<th>R²</th>
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</table>

| II    | 0.12  | 0.25 | 0.62 | 1.00 | 0.62 | 0.00 | 0.37   | 0.75  | 0.25 | 0.50 |
|       | 0.42  | 0.45 | 0.94 | 1.00 | 0.78 | 0.44 | 0.48   | 0.57  | 0.48 | 0.68 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 100.00    | 0.55 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 23.23     | 0.51 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 58.16     | 0.49 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 0.42      | 0.44 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 45.30     | 0.53 |

|       |       |      |      |      |      |      |       |       |      |      |       |      | 4.70      | 0.48 |

Table 3: Model selection for the Bardet-Biedl data at the 95% confidence level: Lower boundary models (LBM1–LBM8), inclusion importance statistics for the entire confidence set ($\hat{II}$) and the lower boundary ($\tilde{II}$), full, Lasso, Scad, Mcp, AIC and BIC selections; X denotes “selected” and (*) indicates models outside the confidence set (unSAFE). For each model we include percent p-values for the F-test with the full model (p-val) and coefficients of determination based on ordinary least squares fits ($R^2$). Lasso, Scad and Mcp are computed using 5-fold cross-validated hyper-parameters. AIC and BIC models are computed by exhaustive search.

5 Monte Carlo simulations

We sample $n$ covariate vectors in the design matrix from a multivariate normal distribution with mean zero and covariance matrix $\Sigma$. For each covariate vector, we compute the corresponding response $y = x'\beta + \epsilon$, where $\epsilon$ is sampled from $N(0, \sigma^2)$. We study the following setups, which cover a variety of scenarios concerning the magnitude of the relevant predictors and their correlation:

- **Model 1.** The first $k = p/2$ have the same size and the rest is equal to zero: $\beta_j = 1, j = 1, \ldots, k$ and $\beta_j = 0, j = k+1, \ldots, p$. The correlation between the $i$th and $j$th covariates is $\Sigma_{ij} = \rho^{|i-j|}$, $0 \leq \rho < 1$.

- **Model 2.** The first $k = p/2$ coefficients have decreasing size and the rest is equal to zero: $\beta_j = 1/j, j = 1, \ldots, k$ and $\beta_j = 0, j = k+1, \ldots, p$. The correlation between the $i$th and $j$th covariates is $\Sigma_{ij} = \rho^{|i-j|}$, $0 \leq \rho < 1$.

- **Model 3.** Same structure for the coefficients as in Model 1. However, half of the predictors with zero and nonzero coefficients have $\Sigma_{ij} = \rho \not= 0$. The remaining pairwise correlations are zero.

5.1 MC Example 1: ECS and LBM set size

In Table 4, we show Monte Carlo estimates for the ECS size, LBM set size and average number of variables for the lower boundary models based on different choices of $p$, $\rho$ and $\alpha$. The number of models in the ECS is monotone in $\alpha$ with smaller values of $\alpha$ corresponding to larger confidence sets. A similar behavior occurs for the LBM set size when the predictors are orthogonal and all the non-zero coefficients have the same size (Model 1). However, when some of the coefficients are small relative to the others (Model 2), we do not have a monotonicity in $\alpha$. 

Figure 3: Inclusion and co-inclusion importance at the 95% confidence level for the the Bardet-Biedl micro-array data. (a) Inclusion importance profile of predictors ($\tilde{I}$) computed on the entire confidence set. (b) 95% confidence co-inclusion importance graph with edges representing values the co-inclusion statistic $CII$ defined in Section 3.2 (edges corresponding to $CII < 0.2$ are omitted for clarity).

Note that while the size of the ECS increases rapidly in $p$, that of the LBM set remains relatively small. This is important in light of Theorem 3.1 since the boundary models contain sufficient information about the variables in the true model. In the worst case, in terms of signal-to-noise ratio (Model 2, $p = 12$, $\rho = 0.7$), the boundary set has less than 12 models. This suggests that although the size of all the models in the ECS may be huge without further restrictions when $p$ is large, computing the LBMs can still be managed for a moderately large number of predictors.

Finally, note that the number of predictors in the LBMs grows with $\alpha$. When the confidence level increases, the LBMs are more parsimonious. If the confidence level is small, there are only a few but relatively large LBMs. In the presence of relatively small coefficients and a large correlation, the LBMs are numerous but they contain fewer predictors.

5.2 MC Example 2: importance profile of predictors

This example illustrates the behavior of the ECS and LBM set, given different significance levels. We consider a sequence of equally spaced values for $\alpha$ ranging from 0.001 to 0.1 and draw 100 Monte Carlo samples of size $n = 100$ from Model 3 with $p = 8$ predictors. Only the first four predictors have nonzero coefficients $\beta_1 = \beta_2 = \beta_3 = \beta_4 = 1$. While $x_1$, $x_2$, $x_5$ and $x_6$ orthogonal to all the other predictors, $x_3$, $x_4$, $x_7$ and $x_8$ are moderately correlated with correlation $\rho = 0.5$.

Figure 4 (a) shows the Monte Carlo averages of the predictors’ inclusion importance profile plots (solid line)
Table 4: Monte Carlo estimates for size of the exact confidence set (ECS), lower boundary model (LBM) set and average size of LBMs. The results are based on 500 Monte Carlo samples of size $n = 100$ from Models 1 and 2, for different choices for different confidence levels (Conf.), predictors’ correlation ($\rho$) and number of predictors ($p$). Monte Carlo standard errors are smaller than 0.01.

6 Comparison with the work of Hansen et al. (2011)

As mentioned in the Introduction, a recent paper by Hansen et al. (2011) is closely related to our work. Both indeed share the same goal of providing a confidence set of models to give the data analyst a proper sense on how far the information in the data can allow him/her to go in terms of identification of the best model. They also share the same view that the size of the confidence set is a valuable indicator of the degree of model selection uncertainty. However, the approaches in the two papers are drastically different. Below we discuss the differences.

6.1 Modeling assumptions

In Hansen et al. (2011), the true data generating model is not specified and the candidate models provide estimations/predictions that are assessed in terms of a chosen loss function. This flexible framework targets various potential applications. Our work, however, starts with a clearly specified full model in the normal regression setting and the issue is on which subset model to use.

Given the setup in this paper, no more assumptions are needed for our construction of the variable selection confidence set. For Hansen et al. (2011), to proceed, an essential assumption is made (on page 458): the mean of the loss difference between any two models stays the same over time. This seems to be very restrictive. For the usual regression data, conditional on the design matrix, even for the true model, the losses at the observations typically...
Figure 4: Inclusion importance ($II$) profile plots simulated from Model 3. The solid and dashed black lines represent Monte Carlo means and 95% confidence bands, respectively. The lighter lines show 100 Monte Carlo realizations. The plots are based on 50 Monte Carlo samples of size $n = 100$ from Model 3, where predictors $x_1, x_2, x_5, x_6$ are orthogonal to all the predictors, while $x_3, x_4, x_7, x_8$ have pairwise correlation $\rho = 0.5$.

are distinct and the mean loss differences are expected to depend on the cases. It is unclear to us how the setup in Section 3.2.1 of Hansen et al. (2011) satisfies the assumption. For the forecasting problem with time series data, the assumption is even less likely to hold because for each reasonable model, the forecast error distribution should have smaller and smaller variability as more information becomes available, which makes the assumption highly unlikely to be met. Furthermore, the demand that for each pair of models, the mean loss difference is either zero or a nonzero constant seems at odds with applications we are familiar with. When multiple models are considered, in general, besides the best model(s), there are models that overfit (e.g., a model with a few extra parameters than the true model or a nonparametric model when the true model is parametric) and there are also under-fitting models. In the traditional truth-fixed analysis, the mean loss difference may indeed be of the constant order (but unlikely to be constant as mentioned earlier) when one of the models is incorrect. But when comparing the true model and an overfitting model or two overfitting models, the mean loss difference actually converges to zero (e.g.,
at order $1/n$ multiplied by the number of extra parameters between the nested models under the squared error loss). Therefore the mean loss difference assumption in Hansen et al. (2011) also immediately rules out applications where overfitting models exist, and it may have over-simplified the nature of different performances of the candidate models.

6.2 Asymptotic or exact coverage

Given the restrictive framework in this paper, the exact confidence set offers finite-sample coverage guarantee. When the set of lower boundary models are considered, we have given only asymptotic results on the containment of the true model and related quantities under an additional condition on the signal strength. In fact, it is easy to show that without such a condition, it is simply impossible to make any sensible correct statement on whether the true model is a lower boundary model. Hansen et al. (2011) starts with an asymptotic result on the behavior of their confidence set (Theorem 1). Later, in the pursuit of a finite-sample coverage probability, a coherency condition is needed to relate the equivalence test and the elimination rule. However, the example of tests from $t$-statistics does not seem to actually work: Proposition 1 is valid only when $c$ in its proof is a constant that should be chosen for the desired coverage probability. When it is estimated based on bootstrap, however, it becomes random and is dependent on the same data on which the max $T$ statistic is constructed. Since the coherency is an exact requirement, the asymptotic justification of the bootstrap method does not appear to be sufficient for deriving a non-asymptotic confidence set. Thus it remains to be seen how their sought finite-sample performance can be achieved for an implementable procedure.

6.3 Handling high-dimensional problems

In our work, the number of predictors, $p$, is allowed to grow with $n$ to capture the challenge in high-dimensional regression. In such a case, there are possibly many models (among the $2^p$ choices) that are hard to be distinguished by testing or whatever method. Our idea is to use the set of the lower boundary models to properly reflect the reality. Although it is not explicitly stated in Hansen et al. (2011), the number of models considered there is fixed for the theoretical results. Indeed, from the proof of Theorem 1 in their paper, if the number of models grows with $n$, since there are unbounded number of events that involve a sequence of correct rejections of the model equivalence hypotheses, each followed by a correct deletion of a poor model before the final incorrect rejection of a best model, the probability of incorrectly rejecting a best model cannot be guaranteed to go to zero as $n \to \infty$ under their Assumption 1. Similarly, the probability of retaining poor models does not necessarily converge to 0. This issue with the number of candidate models being large relative to the sample size seems to present a fundamental challenge to the current methodology in Hansen et al. (2011).
7 Concluding remarks

For reasonably complicated high-dimensional data, it is usually unrealistic to expect a unique model to stand out as the “true” or best model. Rather a number of models are more or less equally supported by the data. In such a situation, it is better to be aware of the top models for a deeper understanding the relationship between the response and the predictors. In this work, we have demonstrated the usefulness of having a variable selection confidence set from multiple aspects. Specifically, the examination on whether a model selected is \((1 - \alpha)\text{-SAFE}\), the inclusion importance and co-inclusion importance all provide valuable information unavailable in the single selected model, no matter what model selection criterion is used.

Statistical estimation/prediction or inference based on more than one model is by no means a new topic. Indeed, model averaging, as mentioned in Section 1, tries to reduce the uncertainty associated with a forced choice of a single model. For example, Burnham and Anderson (2002) advocate the use of Akaike weights for assessing strengths of the candidate models and for model averaging. While such model selection criterion-based weights do provide an intuitive view on the relative usefulnesses of the candidate models, more work is needed to understand how the weights can be interpreted pertaining to reliably selecting the most important variables.

We have shown that the VSCS can be used as a model selection diagnostic tool. To examine a model selected by a sparse modeling method, one can first come up with a super model by moving further along its solution path and adding a few predictors recommended by some other model selection methods. If the model is not 95%-SAFE, then there is strong reason to doubt the soundness of the set of predictors in the model. Furthermore, by comparing it with the LBMs, one has a good idea of which important predictors are missed. Of course, the outcome of the diagnostic process is much more informative when a negative result is reached.

It should be pointed out that although we can always quickly check whether one or a few models selected by certain methods are in the ECS or not, when \(p\) is large, it is computationally challenging to go over all the subset models to identify the entire ECS without further conditions. With that, in the numerical work of this paper, we have limited our scope to manageable sets of candidate models with \(p\) relatively small (possibly after a variable screening). Although the size of the ECS may grow fast in \(p\), the number of LBMs does not grow as much and can be often computed, as seen in the illustrations given in Section 5. Nevertheless, the computation to list out all the LBMs can still be costly. We have planned to seriously examine the computation issues of ECS and LBMs in the future.

Acknowledgement

We sincerely thank the two reviewers and the AE for their very helpful comments and suggestions for improving our work. In particular, the reference of Hansen et al. (2011) that they brought to our attention for comparison
and discussion is appreciated.

References


