Exploiting Disagreement Between High-Dimensional Variable Selectors for Uncertainty Visualization

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ABSTRACT

We proposed combined selection and uncertainty visualization (CSUV), which visualizes selection uncertainties for covariates in high-dimensional linear regression by exploiting the (dis)agreement among different base selectors. Our proposed method highlights covariates that get selected the most frequently by the different base variable selection methods on subsampled data. The method is generic and can be used with different existing variable selection methods. We demonstrate its performance using real and simulated data. The corresponding R package CSUV is at https://github.com/christineyuen/CSUV, and the graphical tool is also available online via https://csuv.shinyapps.io/csuv.

1. Introduction

Model and variable selection in high-dimensional regression settings have been widely discussed in the past decades, with techniques such as the best subset selection (Beale, Kendall, and Mann 1967), Lasso (Tibshirani 1996), Elastic Net (Zou and Hastie 2005), Group Lasso (Yuan and Lin 2006), SCAD (Fan and Li 2001), MCP (Zhang 2010), and a handful of others having gained widespread popularity. Fan and Lv (2010) provided a detailed review of different variable selection methods in high-dimensional settings. On the uncertainty quantification front, there has been a growing focus on post-selection inference. Van de Geer et al. (2014), Zhang and Zhang (2014) and Javanmard and Montanari (2018) advocated the de-biasing approach, which constructs confidence intervals for covariates by de-sparsifying the Lasso estimators. Lee et al. (2016), Tibshirani et al. (2016), and Tibshirani et al. (2018) proposed a conditional approach which provides confidence intervals for the selected covariates using the distribution of a post-selection estimator conditioning on the selection event. Chatterjee and Lahiri (2011) and Liu and Yu (2013) suggested using bootstrapping on some existing variable selectors.

In this article, we propose a simple approach to assessing and visualizing selection uncertainty in the linear model. We assume that the observed data are the realization of

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

where \( p \) is the number of covariates, \( n \) is the number of observations, and we potentially have \( p > n \). \( X_{ij} \) is the \( j \)th covariate of the \( i \)th observation of \( X \) and \( X \) is a fixed \( n \times p \) design matrix. \( X \) is standardized with each covariate \( X^t \) has \( \sum_{i=1}^{n} X_{ij}^2/n = 1 \).

Furthermore, the model is assumed to be sparse with the set of true covariates \( S = \{ j \in \{1, ..., p \} : \beta_j \neq 0 \} \), \(|S| \ll p \).

Given a dataset, how to select the best variable selection method remains an open and yet very important question to ask. Various theoretical performance guarantees are available for a range of methods (e.g., the irrepresentable condition (Zhao and Yu 2006) is sufficient and almost necessary for the Lasso to be sign consistent) but many of them are not testable in practice; for instance, checking the irrepresentable condition usually requires knowing the true set of covariates. Therefore, this type of theory can be of limited practical use in method selection. To illustrate the uncertainty associated with method selection, let us consider two real-life datasets in Examples 1 and 2.

Example 1 (Riboflavin data). The riboflavin dataset concerns the riboflavin (vitamin B2) production by bacillus subtilis. The response is the logarithm of the riboflavin production rate by bacillus subtilis and the \( p = 4088 \) covariates are the logarithms of the expression levels of 4088 genes. The number of samples is \( n = 71 \ll p \). The dataset is available in the R package lassoglm.

Example 2 (Prostate cancer data, Stamey et al., 1989). The prostate cancer dataset comes from a study that examined the relationship between the level of prostate-specific antigen and \( p = 8 \) clinical measures (logarithm of weight, age, Gleason score, among others) in men who were about to receive a radical prostatectomy. The sample size is \( n = 97 \). The dataset is available in the R package lassoglm.

We process the datasets using five different variable selection methods: the Lasso, Elastic Net, relaxed Lasso (Meinshausen and Yu 2009), Group Lasso (Yuan and Lin 2008), MCP (Zhang 2010), and MCP (Zhang 2010)
MCP, and SCAD in R with default tuning in the corresponding R packages (see Section 5 for more details). The selection results are shown in Figures 1 and 2. Figure 1 shows that for the riboflavin dataset the sets of covariates selected vary significantly among the methods, which makes it difficult to justify the validity of the set of covariates selected using any one method. For the prostate cancer dataset, even though there are only eight covariates to choose from, there is still selection disagreement among the methods (Figure 2).

Such disagreement among methods as shown in Figures 1 and 2 is not an exception but a common observation. The distance heat maps in Appendix A.4 show that selection disagreement manifests itself across different simulation settings. Having observed disagreement, one possible way to proceed would be to rank the different models considered (e.g., using cross-validation or an information criterion) and select the highest-ranked one.

However, notwithstanding the usefulness of using a single, well-performing variable selection method, it is tempting to ask whether more can be said regarding the uncertainty of variable selection, based on the disagreement between the methods tested. The similarities and disagreements among the different variable selectors, which is a piece of information not typically used by any one of them, may provide us with some useful insight. For example, in Figure 1 all of the methods select the first three covariates whereas the remaining covariates are selected by some of the methods only. Does it mean that the first three covariates are more likely to be the true covariates? This question is central to this article, and motivates our main development. In this article, we propose a new tool for uncertainty visualization associated with variable selection, termed combined selection and uncertainty visualizer (CSUV). CSUV combines, in a particular way, a number of different base variable selection methods into the solution path of a new variable selector, and illustrates the output of this new selector together with a graphical representation of its uncertainty. It makes use of sets of covariates selected on different subsamples of the data with different variable selection methods. A full description of the proposed method is in Section 3 and 4. The variable selection part of the proposed procedure can be summarized as follows: first, split the data into the training and test sets and fit different variable selection methods on the training set over a grid of tuning parameter values. Estimate the performance of the fitted models on the test set, and retain only the best-performing model. Repeat the process a number of times and select the covariates that appear the most frequently in the collection of the retained fitted models.

The goal of variable selection in our context is purely interpretative: we are interested in visualizing the uncertainty associated with including each individual covariate into any model. The user aware of the data-analytic context can then make their own decision as to whether or not to include each covariate, based on the task at hand (e.g., prediction). This task is facilitated by our approach, as it returns what can be interpreted as a measure of importance of each covariate, and a natural ordering of the covariates according to the certainty of their inclusion. Based on the output of the procedure, the user can select a number of sets of covariates, if this is preferred—for example, by trying different combinations of the "most certain" variables.

An important component of CSUV is a graphical tool designed to visualize the selection uncertainty by using disagreement among the different model fits. See Figure 3 for an example of a graphical output of CSUV. The plot shows the frequency with which each covariate is selected and the variability of the estimated coefficients.

The article is organized as follows. In Section 2, we describe some related work. In Section 3, we discuss the main ideas behind CSUV, and we present the solution path, variable selec-
2. Related Work

One possibility open to analysts when faced with competing fitted models is to select one of them. For example, Chen and Chen (2008) proposed eBIC, an extension of BIC to high-dimensional data which takes into account both the number of unknown parameters and the complexity of the model space. Zhang and Yang (2015) advocated the use of delete-$n/2$ cross-validation to select a method among all the candidate methods.

Model combination with subsampling has been used to improve variable selection performance of a single variable selection method. For example, Bolasso (Bach 2008) fits the Lasso on each bootstrap sample and takes the intersection of all the selections. Wang, Peng, and Dunson (2014) proposed the median selection subset aggregation estimation (MESSAGE) algorithm which aims to perform variable selection on large-$n$ datasets.

The ranking-based variable selection (Baranowski, Chen, and Fryzlewicz 2020) algorithm uses subsampling to identify the set of consistently highly ranked covariates. Stability selection (Meinshausen and Bühlmann, 2010 and Shah and Samworth, 2013) provides control over the finite sample familywise type I errors via subsampling.

Similarly to the methods above, CSUV, our proposal, fits variable selection methods on subsampled data and selects the covariates that appear the most frequently. Unlike these other approaches, however, CSUV makes use of different variable selection methods as we observe that no one method outperforms all other methods in all settings. This brings various advantages, including obtaining access to good model fits from different variable selection methods, and being able to exploit disagreement between the selectors to evaluate selection uncertainty. We elaborate on these points later.

One high-level difference between CSUV and stability selection is the output from both procedures: CSUV is designed to visualise the uncertainty associated with the inclusion of each covariate into (any) linear model, whereas stability selection does not have a similar function. To this end, CSUV uses information from the disagreement between different model fits, a point of view that is absent from stability selection. In addition, CSUV has a mechanism for only considering the disagreement between “good” model fits; this is achieved by filtering out (predictively) “bad” model fits. (Clearly, there would be little point in considering disagreement between a number of unsatisfactory model fits.) This point of view is not present in stability selection, which does not automatically have a mechanism for filtering out unsatisfactory base methods or model fits.

Adaptive regression by mixing (ARM, Yang 2001) and its variation, adaptive regression by mixing with screening (ARMS, Yuan and Yang 2005), aggregate fits from different methods by estimating weights through subsampling.

Variable selection deviation measures (VSD, Nan and Yang 2014) aim to provide a sense of how trustworthy a set of selected covariates is. The VSD of a target model $m$ is the weighted cardinality of the symmetric difference between $m$ and each candidate model. Nan and Yang (2014) suggested using the sets of fitted models on the solution paths from the Lasso, SCAD and MCP as candidate models and the weight of each candidate model is calculated based on information criteria or ARM. The simulation results in Nan and Yang (2014) showed that a large VSD compared to the size of the target model means that the target model is not trustworthy, but a small VSD does not
necessarily mean that the target model is close to the true model. Yang and Yang (2017) proposed to select a set of covariates that minimizes the total Hamming distance with all the candidate models in terms of VSD (we refer to this method as VSD-minimizing in the remainder of the article). The authors also propose using different thresholds, where the threshold of 0.5 is equivalent to minimizing the standard Hamming distance. For variable selection method combinations that do not involve subsampling, Tsai and Hsiao (2010), Mares, Wang, and Guo (2016), and Pohjalainen, Rasänen, and Kadioglu (2015) provided empirical results on combining sets of selected covariates from different variable selection methods by intersection, union and/or some other set operations.

Both our method and VSD use resampling and different variable selection methods to provide an assessment of how good the final set of covariate selection is. VSD focuses on the whole model fit. Our method focuses on the uncertainty of individual covariates and a graphical tool is designed to illustrate these uncertainties. In terms of methodology detail, our method combines the sets of covariates selected in resampling fits whereas VSD combines sets of covariates selected on the solution path when fitting using all the data. Resampling data are only used in VSD for calculating the weight of each set of covariates. In our simulation study we compare the variable selection performance of our method to the VSD-minimizing method proposed by Yang and Yang (2017), as it is the method the most similar to CSUV. The simulation results in Section 5 show that in general our method outperforms the VSD-minimizing model.

Hahn and Carvalho (2015) provided an informative review of Bayesian approaches to variable selection in linear models, a formalism in which the data-analytic output consists of probabilities assigned to different models given data. Therefore, these approaches provide natural measures of model uncertainty and should be seen as an alternative to the approach taken in CSUV and other frequentist methods. As reviewed in Hahn and Carvalho (2015), two potential weaknesses of the Bayesian approach in this context are: subjectivity in the choices of the various priors, and the fact that posterior sampling is usually computationally intensive. CSUV attempts to mitigate both these issues by minimizing user involvement in the selection of its parameters, and by using computationally efficient constituent methods.

3. CSUV Solution Path Methodology

3.1. The CSUV Algorithm

The first goal of this article is to use the similarity of fits from different methods to obtain the ranking of covariates, alternatively referred to as a solution path, which can then (if desired) be used for the purpose of variable selection. CSUV achieves this by following the simple aggregation principles below.

- Step 1: fit the data using different variable selection methods.
- Step 2: record the percentage of times a covariate \( X_j \) is selected among the different methods. Denote it by \( \theta_j \).
- Step 3: rank the covariates according to decreasing \( \theta_j \)'s. If variable selection is required, select those that correspond to the highest \( \theta_j \)'s, for example, \( \{ X_j : \theta_j \geq 0.5 \} \).

At the same time, CSUV uses the following additional mechanisms.

- Only include the fitted models that exhibit good performance (see Section 3.2.2).
- Repeat the fitting on subsampled data.

This is done, respectively, to discourage CSUV from focusing on poorly performing methods, and to incorporate the variability in selection caused by the variability in data.

Different variable selection methods optimize different objective functions. In the case of regularized regression, the difference among methods is usually in terms of the penalty. If a covariate is selected by the majority of methods, then it means the covariate is chosen to minimize many different objective functions. We expect that a true covariate \( j \) should have a high \( \theta_j \), i.e. it should frequently be chosen regardless of the objective function used.

The CSUV solution path procedure, detailed in Algorithm 1, can be summarized as follows. First, randomly split the data into training and test sets, and fit different variable selection methods on the training set over a grid of regularization parameters (see Section 3.2.4 for more details). Then, use the test set to calculate the performance of the fitted models and retain the best model (see Section 3.2.2 for more details on performance measures). Repeat the process many times. Order the covariates according to the relative same sign frequency \( \tau_j \), which we now define.

Definition 1 (Relative same sign frequency \( \tau_j \)). Assume a set of fitted models \( \mathcal{M} \). The relative same sign frequency of covariate \( X_j \) is defined as

\[
\tau_j = \frac{1}{|\mathcal{M}|} \max \left( \sum_{\mathcal{M}_k \in \mathcal{M}} \frac{1}{\beta_{j, \mathcal{M}_k}^{>0}}, \sum_{\mathcal{M}_k \in \mathcal{M}} \frac{1}{\beta_{j, \mathcal{M}_k}^{<0}} \right),
\]

where \( \beta_{j, \mathcal{M}_k}^{>0} \) is the estimated coefficient of the \( j \)th covariate on the fitted model \( \mathcal{M}_k \in \mathcal{M} \), and \( 1_x \) is the indicator function.

Algorithm 1 involves repeated fits on subsamples of data, and this can be computationally expensive. Fortunately, the algorithm can easily be parallelized by running iterations on different cores/machines, which makes it feasible for high-dimensional data analysis.

3.2. Specifications for CSUV Variable Selection

Algorithm 1 provides a general framework for the CSUV solution path and variable selection approach. Here we discuss how its parameters should or may be set for practical use.

3.2.1. Coefficient Estimation

If Algorithm 1 is tasked with variable selection, it only selects a set of covariates without estimating the \( \beta \) coefficients. In our implementation we use ordinary least-square (OLS) to estimate the \( \beta \) coefficients on the selected set \( \mathcal{S} \) using the full set of data to form the final fitted model. If the number of covariates selected is larger than the number of observations, then the default option in our code is to use ridge regression to estimate the coefficients by cross-validation (with the default cross-validation setting
Algorithm 1 CSUV solution path (and variable selection)

Input: variable selection methods $A_1, \ldots, A_R$ with the corresponding generation of the grid of regularization parameters; $n$ observations with $p$ covariates $X$ and response $Y$; number of repetitions $B$; percentage of data used in training set $w$; performance measure; (for variable selection only) frequency threshold $t$.

Output: ranking of covariates; (for variable selection only) set of selected covariates $S$.

1: for $b$ in $[1, \ldots, B]$ do
2: Randomly assign $w\%$ of the observations as training data with labels $l_{\text{train}}^b$ and the rest as test data with label $l_{\text{test}}^b$. Fit data with label $l_{\text{train}}^b$ using $A_1, \ldots, A_R$ over grids of $K_p$ different values of the corresponding regularization parameters, $r \in \{1, \ldots, R\}$. For each method $A_r$, denote the fitted models as $\hat{M}_{r,1}^b, \ldots, \hat{M}_{r,K_r}^b$, and the set of covariates selected by each fitted model as $S_{\hat{M}_{r,k}^b} = \{ j : \hat{\beta}_{j,k}^b \neq 0 \}, k \in \{1, \ldots, K_r\}$ where $\hat{\beta}_{j,k}^b$ are the estimated coefficients from the method $A_r$.
3: Remove any duplication within each method in terms of variable selection to get $S_{\hat{M}_{1,k}^b}, \ldots, S_{\hat{M}_{K_p,k}^b}$ such that for each $r, S_{\hat{M}_{r,k}^b} \neq S_{\hat{M}_{r,k'}^b} \forall k \neq k' \in \{1, \ldots, K_r\}$. Record the sets of covariates selected by each fitted model $S_{\hat{M}_{1,k}^b}, \ldots, S_{\hat{M}_{K_p,k}^b}, S_{\hat{M}_{r,k}^b}, r \neq k \neq \hat{r}$ and re-index as $S_{\hat{M}_{1,k}^b}, \ldots, S_{\hat{M}_{K_p,k}^b}$, where $K^b$ is the number of fitted models recorded.
4: If the number of selected covariates $|S_{\hat{M}_{r,k}^b}| < |l_{\text{train}}^b|$, refit the selected set of covariates $S_{\hat{M}_{r,k}^b}$ using ordinary least squares (OLS), to get the fitted models $\hat{M}_{r,1}^b, \ldots, \hat{M}_{r,K_r}^b$ with the estimated coefficients $\hat{\beta}_{j,k}^{\hat{M}_{r,k}^b}$. Otherwise, set $\hat{\beta}_{j,k}^{\hat{M}_{r,k}^b} = \hat{\beta}_{j,k}^b$.
5: Use data with label $l_{\text{test}}^b$ to estimate the performance of each fitted model $\hat{M}_{r,k}^b$ from Step 4 according to the given performance measure. Retain the best fitted model and denote it as $\hat{M}_{r,1}^b$.
6: end for
7: Denote the set of retained fitted models by $\mathcal{M} = \{\hat{M}_{1,1}^b, \ldots, \hat{M}_{K_p,1}^b\}$.
8: Calculate the relative same sign frequency $\tau_j$ for each variable $j$, which defines the ranking of covariates.
9: If variable selection is required, select the covariates $\hat{S}$ defined by

$$\hat{S} = \{ j : \tau_j \geq t \}.$$ 

from the glmnet R package; however, this should be treated as an exception mode as the number of covariates selected is almost always much smaller than the number of observations.

3.2.2. Performance Measure

Step 5 of Algorithm 1 aims to select the fitted models based on their variable selection performance. In general, in attempting to select fitted models or methods with good variable selection performance, it is common to use prediction measures such as MSE or information criteria such as BIC or eBIC. Theoretically, BIC is consistent in model identification when $p$ is fixed and eBIC is consistent in high-dimensional settings (Chen and Chen 2008). Our empirical experiments, however, show that when using BIC or eBIC as performance measures in Algorithm 1, the resulting fitted models tend to select too few covariates so the final selection by CSUV omits too many true covariates. By contrast, using MSE as the performance measure in Algorithm 1 in our simulation settings provides good variable selection performance. Although MSE measures prediction rather than variable selection performance, MSE is often used for variable selection methods such as in selecting tuning parameter $\lambda$ for SCAD (Fan and Li 2001).

3.2.3. Percentage of Data Used in Training

Following Yang (2001), Yuan and Yang (2005), and Zhang and Yang (2015), we use $w\% = 50\%$ of the data for fitting and the remaining $50\%$ for testing by default; this splitting ratio attempts to ensure a sufficiently large sample size for both. Stability selection (Meinshausen and Bühlmann 2010; Shah and Samworth 2013) also uses the same splitting ratio although their rationale is that subsampling with such a ratio behaves similarly to bootstrapping. Empirically, we found that departures from $50:50$ were occasionally beneficial; many procedures display at least some sensitivity to the training/test split, so this phenomenon in our case is neither unique nor unexpected. For example, we applied CSUV in a higher-dimensional case involving $p = 1000$ (with $n = 100$) and the $75:25$ split was preferred there—possibly because it was particularly important for the algorithm to maximize the size of the training sample when there were so many variables. On the other hand, when $n$ was large in relation to $p$ ($n = 1000$ and $p = 100$), the performance of CSUV with three different splits ratios ($25:75, 50:50,$ and $75:25$) was similar, with $25:75$ and $50:50$ performing slightly better than $75:25,$ quite possibly because with such a large $n$, even $25:75$ was seen as sufficient for training.

3.2.4. Constituent Variable Selection Methods

CSUV is designed to be generic so that any variable selection methods can be used as the constituent methods $A_1, \ldots, A_R$ in CSUV. Ideally, all the methods $A_r$ should have good variable selection performance, and there should be some variability among the methods in terms of false selection. The base methods should also be computationally efficient they are fitted on subsampled data multiple times. In this article, we choose the Lasso, MCP and SCAD to be the default constituent methods as they are optimizing different objective functions. Methods like Elastic Net or relaxed Lasso are not used by default as they are relatively similar to Lasso. Our default base methods are also computationally feasible in high-dimensional settings with efficient fitting algorithms available, and there is also a default way to compute the grid of regularization parameters to consider. For example, the R package ncvreg for MCP and SCAD by default computes a sequence of parameters $\lambda$ with equal spacing on the log scale and of length 100, starting from the smallest value 0.001. We do not consider two-stage methods such as the adaptive Lasso (Zou 2006) as they are relatively slow, or methods without default parameter tuning in R (for example, the Dantzig selector, Candes and Tao 2007) as they
make comparison with other methods like delete-n/2 cross-validation more complicated.

CSUV can tolerate duplicated or very similar methods, although it is not recommended due to the computational time. In our experience, including duplicated or very similar methods tends not to affect the variable selection performance. In our simulations, when methods that usually select similar sets to the Lasso (such as the Elastic Net or relaxed Lasso) are included, the performance of CSUV does not change much.

A suggestion made by a referee is to incorporate a randomized method (or a selection thereof) as a constituent variable selector in CSUV, to increase the diversity of the constituent methods. In a randomized method, each model uses a random subset of variables, chosen according to a particular recipe (akin to a Bayesian stochastic search). However, it is not obvious to us how to select reasonable random models in a computationally efficient way, and we therefore leave this interesting idea for future work.

### 3.2.5. Number of Repetitions

The number of repetitions $B$ should be large enough to stabilize the value of $\tau_j$ and at the same time it should not be too large so that Algorithm 1 can be run within a reasonable time. We use $B = 100$ in our experiments.

### 3.2.6. Frequency Threshold

The frequency threshold $t$ features in Step 9 of Algorithm 1 and is used when variable selection is required. The CSUV method described by Algorithm 1 and using $t = 1/2$ is denoted by CSUV-m, where the "m" in CSUV-m stands for median, because selecting covariates with $\tau_j \geq 1/2$ is equivalent to selecting covariates with a nonzero median in $M$. The choice $t = 1/2$ is optimal in the sense of minimizing certain distance functions; the details are in Propositions 1 and 2 in the appendix. Empirically, even though CSUV-m strikes a good balance between false inclusion and false omission, compared to other variable selection methods it usually includes many fewer false covariates, with the tradeoff being that it occasionally omits some true covariates. When the analyst's focus is on performance criteria other than variable selection, for example on prediction, they may want to select more covariates. This can be done by considering other thresholds $t$ on the sign frequency $\tau_j$ or a threshold on the model size which we introduce next.

### 3.3. Solution Path and Alternative Variable Selection

Algorithm 1 generates a solution path (formally defined below) by ordering covariates from the highest to the lowest relative same sign frequency $\tau_j$. This solution path can be regarded as a series of nested sets of covariates with increasing model sizes.

**Definition 2 (CSUV solution path).** The CSUV solution path orders covariates so that

$$R_j < R_f$$

if $\tau_j > \tau_f$ or $(\tau_j = \tau_f$ and $|\hat{\beta}_j| > |\hat{\beta}_f|)$

where $R_j$ is the position of covariate $j$ on the solution path, $\tau_j$ is the relative same sign frequency calculated in Step 9 of Algorithm 1 and $\hat{\beta}_j$ is the average of the estimated coefficients in $M$ in Step 7 of Algorithm 1.

The standardization of the design matrix in Equation (1) ensures the comparison of the size of the estimated coefficients is meaningful. The nested character of the solution path makes it easy to entertain alternatives to CSUV-m. In particular, consider replacing line 9 of Algorithm 1 by $\hat{S} = \{j : R_j \leq s\}$, where $s$ is a given size thresholds which specifies the model size. In the particular implementation of CSUV described in this article, we set the size threshold $s$ equal to the median size of the selected sets in $M$ in Step 7 of Algorithm 1 and we define CSUV with this threshold as CSUV-s.

### 4. CSUV Visualization of Uncertainty

#### 4.1. Graphical Component of CSUV

In this section, we introduce the graphical component of CSUV, a tool designed to illustrate the variable selection and estimation uncertainty. An example of a plot is shown in Figure 3 and the graphical tool is available interactively at https://csuv.shinyapps.io/csuv and in the R package CSUV (Yuen 2021). It has the following ingredients.

**Box plots that visualize the estimated coefficient uncertainty.** Each box plot corresponds to a covariate $X_j$ and shows the lower and the upper quartiles of the empirical distributions of the estimated coefficients. Figure 3 shows the unconditional distributions for each covariate. The CSUV package also has a conditional option, in which the box plots show the distributions conditional on the estimated coefficients being nonzero.

In addition, users desiring more details of the empirical distribution of estimated coefficients are able to superimpose the corresponding violin plots on the box plots in the CSUV package. The extent of the whiskers, set by default to the 5 and 95 percentiles of each estimated coefficient, can be adjusted in the CSUV package. The median value of each estimated coefficient is shown as a horizontal line in each box (blue in the color version). The box plots are ordered according to the solution path (Definition 2).

**Shaded background representing $\tau_j$.** The background behind each box plot is shaded according to the relative same sign frequency $\tau_j$ of the corresponding covariate. The darker the color, the higher the value of $[100%\tau_j/10]$. The actual value of $\tau_j$ is displayed in black underneath the boxplots.

**Lines showing the cut-off points for variable selection by the various versions of CSUV.** CSUV-m selects all covariates to the left of the solid vertical line; CSUV-s selects all those to the left of the dotted vertical line.

Figure 3 illustrates (dis)agreement between the constituent variable selectors, in the sense of showing the range of estimates for each linear parameter obtained for each subsample, for each method that wins a prediction competition on that subsample. This ensures that only "good" methods are evaluated (there is little point in including less adequate methods for the purpose of uncertainty visualization, if a better method can be run instead). If the chosen "good" methods exhibit large variation in terms of the parameter value associated with a given covariate, then this can be viewed as a sign of high uncertainty as to whether the covariate should enter any linear model for the data (given that
Our simulations involving synthetic data use five different linear models, details of which are in Section A.6.4. Our real datasets are described below.

**Example 3 (Boston housing data, Harrison Jr and Rubinfeld, 1978).** The dataset consists of the median value of owner-occupied homes as response and $p = 13$ covariates (crime rate, proportion of residential land, etc). Number of observations is $n = 506$. The dataset is publicly available in R with the MASS package. For each simulation, half of the observations (randomized) are used as the training data and the other half are used as the test set; $m = 100$ repetitions.

**Example 4 (Modified riboflavin data).** Here, we re-examine the riboflavin dataset introduced in Example 1. In order to assess the variable selection performance, we randomly permute all but 10 of the 4088 covariates in the riboflavin dataset across all the observations. The same permutation is used for all permuted covariates to keep the original dependence structure among them. The set of 10 unpermuted covariates is chosen randomly among the 200 covariates with the highest marginal correlation with the response. In the simulation results, we refer the 10 unpermuted covariates as the “true” covariates. We repeat the process $m = 100$ times with random selection of the 10 unpermuted covariates.

### 5. Simulation Study

#### 5.1. Settings

In this section, we evaluate the variable selection performance of CSUV. We consider CSUV with different sets of constituent methods: (i) Lasso, MCP, and SCAD (default); (ii) Lasso, Elastic Net, relaxed Lasso, SCAD; (iii) Lasso, Elastic Net, relaxed Lasso, MCP, SCAD, and trees; and (iv) MCP only. The first set is our primary interest. When we mention CSUV without specifying the corresponding constituent methods, we implicitly assume that this set of methods is used. The second and the third combinations are used to verify the claim that adding some similar or inappropriate methods does not affect the performance too much. The fourth set is used to verify the claim that using more constituent methods in general provides better results. We use MCP here because in the majority of the simulation settings it has the best variable selection performance among the individual variable selection methods in terms of the F-measure and the number of false classifications.

We use publicly available R packages for the implementation of the constituent methods (Lasso, Elastic Net, relaxed Lasso, MCP, SCAD, and tree) used in CSUV. Further details are in Section A.6.1. We use eBIC and delete-$n/2$ cross-validation as the major competitors to CSUV. We use eBIC instead of BIC as eBIC is designed for high-dimensional data. We also include the simulation results of each constituent method (Lasso, Elastic Net, relaxed Lasso, MCP, SCAD, and VSD), VSD-minimizing method (Yang and Yang 2017) and BIC for readers’ reference. Further details are in Section A.6.2.

For the datasets for which we know the true sets of covariates (i.e. simulated data and the modified real datasets), we compare the variable selection performance among different methods by the F-measure, the number of false positives (FP), number of false negatives (FN) and the total number of variable selection errors (FP+FN). Although our main focus is variable selection performance, we also compute the prediction mean square error (MSE) on test set data and the coefficient estimation error ($l_1$ and $l_2$) for CSUV and the comparing methods. Further details are in Section A.6.3.

### Table 1. Model summary: performance of CSUV and methods it compares with.

| methods      | f | FP+FN | FP | FN | pred.err | size | 11.diff | l2.diff |
|--------------|---|------|---|---|---------|------|---------|---------|
| lasso        | 0.39 | 25.02 | 24.45 | 0.57 | 2.83 | 4.48 | 1.12 | 30.95 |
| elastic net  | 0.33 | 32.64 | 32.14 | 0.5 | 3.01 | 5.32 | 1.24 | 38.71 |
| relaxed lasso| 0.58 | 12.29 | 11.35 | 0.94 | 2.87 | 3.98 | 1.09 | 17.48 |
| mcp          | 0.7 | 4.86 | 3.32 | 1.54 | 2.93 | 3.2 | 1.13 | 8.84 |
| scad          | 0.63 | 8.04 | 6.79 | 1.25 | 2.91 | 3.05 | 1.09 | 12.6 |
| vsd          | 0.72 | 3.2 | 0.48 | 2.72 | 3.3 | 3.38 | 1.38 | 4.82 |
| bic          | 0.64 | 10.18 | 8.98 | 1.2 | 2.85 | 3.29 | 1.06 | 14.85 |
| eBIC         | 0.71 | 5.4 | 4.02 | 1.37 | 2.87 | 3.04 | 1.08 | 9.71 |
| CV           | 0.54 | 17.25 | 16.61 | 0.64 | 2.72 | 3.74 | 1.01 | 23.03 |
| CSUV.m       | 0.67 | 7.2 | 0.65 | 0.25 | 3.23 | 2.76 | 1.1 | 5.66 |
| CSUV.s       | 0.7 | 5.92 | 4.69 | 1.23 | 2.82 | 3.2 | 1.04 | 10.52 |
| CSUV.m25     | 0.56 | 4.53 | 0.09 | 4.26 | 5.25 | 4.36 | 1.74 | 2.9 |
| CSUV.m75     | 0.79 | 2.98 | 1.77 | 1.22 | 2.73 | 2.55 | 0.95 | 7.61 |
| CSUV.m.all   | 0.77 | 2.75 | 0.78 | 1.97 | 3.16 | 2.75 | 1.09 | 5.88 |
| CSUV.m.mcp   | 0.68 | 3.34 | 0.15 | 3.19 | 4.31 | 3.47 | 1.38 | 4.02 |
| CSUV.m.with.tree | 0.77 | 2.75 | 0.77 | 1.98 | 3.18 | 2.75 | 1.09 | 5.85 |
CSUV vs other final model selection procedures. In the majority of settings, CSUV-m has a better variable selection performance than eBIC, delete-\(n/2\) cross-validation and VSD in terms of FP+FN and the F-measure, and a better coefficient estimation performance in terms of the \(l_1\) loss. For example, out of the 48 simulation settings in which we know the true set of covariates, CSUV-m has a higher F-measure in, respectively, 45, 41, and 32 of the settings compared to delete-\(n/2\) cv, eBIC, and VSD. CSUV-m usually selects the smallest set of covariates when compared with eBIC, delete-\(n/2\) cross-validation and the individual variable selection methods. In some cases like model 4 parameter setting 6, it selects a much smaller set of covariates than the truth. While this worsens the prediction performance of CSUV-m and we may view it as its limitation, it may be due to the limitation of variable selection as a whole: other methods which select much larger sets of covariates usually include a few more true covariates but inevitably also many more false ones. They may perform better than CSUV-m in terms of prediction, but CSUV-m in general outperforms them in terms of variable selection.

The performance of CSUV-s is more difficult to draw conclusions on. Overall, CSUV-s appears better than delete-\(n/2\) cross-validation in terms of variable selection, but roughly comparable to eBIC.

One encouraging result about CSUV is that in many simulation settings like model 2, CSUV-m outperforms not only the final model selection procedures, but also all individual constituent methods in terms of the F-measure and FP+FN. In some simulation settings, CSUV performs better than the best individual variable selection method in both prediction and the F-measure. For example in model 2, there are several parameter settings (e.g., 2) in which the MSE of CSUV is lower and the F-measure is higher than all individual variable selection methods.

For the variable selection performance on the real data, both versions of CSUV perform very well on the riboflavin data example. CSUV-s has the best performance in terms of the F-measure and the total variable selection error.

**Supplementary Material**

Appendix: Some optimality results on CSUV-m; heat maps to illustrate selection disagreement; CSUV assessment of uncertainty; further details of simulation results.

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