

Solar: L_0 path averaging for fast and accurate variable selection in high-dimensional data*

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Abstract

We propose a new algorithm for variable selection in high dimensional, large scale data: *subsample-ordered least-angle regression (solar)*. Solar constructs a solution path for a subsample using the L_0 norm and then averages solution paths across subsamples. Path averaging retains the ranking information of the informative variables while averaging out sensitivity to high dimensionality, improving variable selection stability, efficiency, and accuracy. Because it uses the same numerical optimizers as lasso, solar may be generalized to many lasso variants. We prove that: (i) with a high probability, path averaging perfectly separates informative variables from redundant variables on the average L_0 path; (ii) solar variable selection is consistent and accurate; and (iii) the probability that solar omits weak signals is controllable for finite sample size. Using simulations, examples, and real-world data, we demonstrate the following advantages of solar: (i) with less than 1/3 of the lasso computation load, solar outperforms lasso in terms of the sparsity (64-84% reduction in redundant variable selection) and accuracy of variable selection; (ii) compared with the lasso safe/strong rule and variable screening, solar largely avoids selection of redundant variables and rejection of informative variables in the presence of complicated dependence structures and harsh settings of the irrerepresentable condition; (iii) the sparsity and stability of solar conserves residual degrees of freedom for data-splitting hypothesis testing, improving the accuracy of post-selection inference on weak signals with limited n ; (iv) replacing lasso with solar in bootstrap selection (e.g., bolasso or stability selection) produces a multi-layer variable ranking scheme that improves selection sparsity and ranking accuracy with the computation load of only one lasso realization; and (v) solar bootstrap selection is substantially faster (98% lower computation time) than the theoretical maximum speedup for parallelized bootstrap lasso (confirmed by Amdahl's law). The efficiency of bootstrap solar makes cross validation computationally affordable for optimizing the bootstrap selection threshold even in high dimensional, large scale data.

Keywords: Variable selection, sparsity, computation time, complicated dependence structure, lasso rules, irrerepresentable condition, bolasso, subsampling selection, variable screening.

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1. Introduction

Recent innovations to lasso-type algorithms (Efron et al., 2004; Friedman et al., 2007, 2010) have largely addressed selection of redundant variables, rejection of informative variables, and poor performance under high multicollinearity in high dimensional ($p > n$) and large scale data (large p and large n). However, in alleviating old problems, the innovations have revealed new challenges.

Bootstrap variable selection [e.g., Bach (2008), Meinshausen and Bühlmann (2010), Wang et al. (2011), and Mameli et al. (2017)] markedly improves selection sparsity and inference accuracy, yet it requires repeating lasso on hundreds of bootstrap subsamples to average selection or inference results. Moreover, accurate bootstrap selection may require tuning hyperparameters on each subsample (Meinshausen and Bühlmann, 2010). As a result, Xu et al. (2012) and Sections 4.4 and 4.6 below show that bootstrap selection methods exponentially increase computation load, limiting applicability in large scale data such as DNA sequencing, image recognition, fMRI and MRI neuroimaging data, and natural language processing (where both p and n are often over 10,000). More seriously, choosing the bootstrap variable selection threshold, typically implemented by simulation or field experience, remains an unsolved issue. Bach (2008) and Huang et al. (2014, Figure 2) illustrate that a pre-defined threshold may omit informative variables (low power) and select redundant variables (high false discovery rate) in both high and low dimensions.

One strategy to improve lasso selection sparsity without increasing the computation burden is to use a post-selection rule to screen variables selected by lasso. Post-lasso selection rules [e.g., the ‘safe rule’ (Ghaoui et al., 2010) and the ‘strong rule’ (Tibshirani et al., 2012)] are capable of reducing the number of variables to enhance computational efficiency in lasso. However, recent research (Wang et al., 2014; Zeng et al., 2017) and Section 3.2 below suggest both rules may be prone to rejecting informative variables, selecting redundant variables, or proposing repeated modifications (e.g., rejecting a variable in an early round and adding it back in a later round).

Data-splitting hypothesis tests are another way to screen variables selected by lasso (Wasserman and Roeder, 2009; Meinshausen et al., 2009; Romano and DiCiccio, 2019; DiCiccio et al., 2020). The dataset is split in two: one part for variable selection, the other for testing. To improve test power, data splitting is repeated on each bootstrap subsample, raising similar computational concerns as bootstrapping variable selection (Bach, 2008). DiCiccio et al. (2020) also argue that because data splitting reserves some of the data for variable selection, it reduces the degrees of freedom for testing on the remaining data, presenting a challenge to detect weak signals when sample size is limited.

Specifically designed to address the challenges of high dimensional data, the variable screening algorithm (Fan and Lv, 2008; Hall and Miller, 2009; Hall et al., 2009; Li et al., 2012a,b) ranks the absolute values of the unconditional correlations between each covariate and the response variable, selecting only the top-ranked variables. However, Fan and Lv (2008), Barut et al. (2016), and Section 3.2

below show that variable screening also suffers from selection of redundant variables and rejection of informative variables when the dependence structures in the data are complicated.

According to [Friedman et al. \(2001\)](#); [Weisberg \(2004\)](#), forward selection was historically dismissed in high-dimensional spaces due to inefficiency and sensitivity to sampling randomness, multicollinearity, noise, and outliers due to the iterative refitting of the residual. Using simulation, [Tibshirani \(2015\)](#) shows that forward selection may produce similar generalization errors to lasso-type estimators and that forward selection is computationally competitive with lasso in image de-noising, matrix completion, etc. However, [Tibshirani \(2015\)](#) does not suggest any solution to a range of issues for forward selection or lasso (solved by lars), including instability of variable selection, selection of redundant variables, lack of robustness to the irrepresentable condition and complicated dependence structures, or sensitivity to sampling randomness, multicollinearity, noise, and outliers.

1.1. Main results

To address these issues, we propose a new forward selection algorithm: *subsample-ordered least-angle regression (solar)*. We also offer a coordinate-descent generalization of solar.

Solar constructs a solution path for a subsample using the L_0 norm and then averages solution paths across subsamples. Path averaging retains the ranking information of the informative variables while averaging out sensitivity to high dimensionality, improving variable selection stability, efficiency, and accuracy. Because it uses the same numerical optimizers as lasso, solar may be generalized to many lasso variants. Under the [Zhang \(2009\)](#) general framework of forward selection, we prove that: (i) with a high probability, path averaging perfectly separates informative variables from redundant variables on the average L_0 path; (ii) solar variable selection is consistent and accurate under the general framework of forward selection; and (iii) the probability that solar omits weak signals is controllable for finite sample size.

Using simulations, examples, and real-world data, we demonstrate the following advantages of solar: (i) with less than 1/3 of the lasso computation load, solar outperforms lasso in terms of the sparsity (64-84% reduction in redundant variable selection) and accuracy of variable selection; (ii) compared with the lasso safe/strong rule and variable screening, solar largely avoids selection of redundant variables and rejection of informative variables in the presence of complicated dependence structures and harsh settings of the irrepresentable condition; (iii) the sparsity and stability of solar conserves residual degrees of freedom for data-splitting hypothesis testing, improving the accuracy of post-selection inference on weak signals with limited n ; (iv) replacing lasso with solar in bootstrap selection (e.g., bolasso or stability selection) produces a multi-layer variable ranking scheme that improves selection sparsity and ranking accuracy with the computation load of only one lasso realization; and (v) solar bootstrap selection is substantially faster (98% lower computation time) than the theoretical

maximum speedup for parallelized bootstrap lasso (confirmed by Amdahl’s law). The efficiency of bootstrap solar makes cross validation (CV) computationally affordable for optimizing the bootstrap selection threshold even in large scale and high dimensional data. We provide a parallel computing package for solar (`solarpy`) that uses a Python interface and an Intel MKL Fortran/C++ compiler in a supplementary file and dedicated [Github page](#).

The paper is organized as follows. In Section 2, we introduce the solar algorithm, outline the theoretical properties of path averaging and solar, explain the coordinate descent generalization of solar, and discuss generalizations of solar to lasso variants. In Section 3, we use examples to demonstrate the advantages of solar over lasso, the safe/strong rules, and variable screening. In Section 4, we use simulations to demonstrate the advantages of solar over lasso-type algorithms in terms of variable selection sparsity, accuracy, and computation load. In Section 5, we use real-world data to show that the improvements from solar are feasible in the presence complicated dependence structures, while lasso and elastic net—the lasso variant alleged (Zou and Hastie, 2005; Jia and Yu, 2010) to have the best selection accuracy and sparsity under multicollinearity—completely lose sparsity. The proofs of the properties of solar are in Supplementary Material A. The `solarpy` code and raw simulation results are in Supplementary Material B.

2. The Solar algorithm

The key to solar lies in the parameterization of the solution path. For any forward selection method, Zhang (2009, Theorem 2) shows that the earlier a variable enters the solution path, the more likely it is to be informative. Thus, an accurate and stable ordering of variables in the solution path may help to identify the informative variables. Since we focus on accuracy, the only relevant feature of the regression coefficients in the solution path is whether $\beta_i = 0$ at each stage. Thus, solar parameterizes the lasso path (or, more generally, any forward selection path) using the L_0 norm.

Definition 2.1 (L_0 solution path). Define the L_0 **solution path** on (Y, X) to be the order that least angle regression includes variables across all stages. For example, if the least angle regression includes \mathbf{x}_3 at stage 1, \mathbf{x}_1 at stage 2 and \mathbf{x}_2 at stage 3, the corresponding L_0 path is the ordered set $\{\mathbf{x}_3, \mathbf{x}_1, \mathbf{x}_2\}$.

2.1. Solar optimized by least angle regression

The solar algorithm involves two steps: *parameterizing and averaging L_0 paths* and *selecting variables on the average L_0 path*.

2.1.1. L_0 path parameterizing and averaging

The solution path is the foundation of variable (feature) selection in L_p -regularized linear modelling. The first step in solar is to improve the robustness of the solution path to high dimensional issues such as multicollinearity, complicated dependence structures, noise, weak signals, etc. In particular, there are two major concerns.

- **Computation efficiency:** computation load is a central concern in subsampling-based model averaging. Because bootstrap methods (e.g., bootstrap lasso) require hundreds of lasso repetitions to average out variable selection issues in high dimensions, they are computationally expensive with large n and large p . Thus, improving selection performance and reducing the number of repetitions would go a long way to reducing computation load.
- **Averaging efficiency:** the L_1 lasso solution path (solved by lars) is essentially a piecewise linear function $\beta = g(\lambda)$, which is easy to average. By contrast, it is not obvious how to average the L_0 path because it is an ordered set of rankings. If we average the ranks each \mathbf{x}_i enters the path in large p problems, a weak signal (i.e., an \mathbf{x}_i with a small but non-zero $\|\beta_i\|_1$ in the population) may occasionally be ranked at a later stage, returning a large stage value, and exerting outlier influence on the stage value averaging. In other words, to accurately average solution paths using as few subsamples as possible, we need a parameterization method for the L_0 path that is more robust to outliers in ranking the \mathbf{x}_i .

Our solution to these concerns is the \hat{q} method, summarized in Algorithm 1 and illustrated in Figure 1. For solution path averaging, instead of using the stage value each \mathbf{x}_i enters the solution path, which varies in the range $[0, +\infty)$, Algorithm 1 uses \hat{q}_i^k , which normalizes the stage value into the range $[0, 1]$. The concentration inequalities for empirical processes show that averaging \hat{q}_i^k across subsamples converges much faster and is much more stable than averaging the stage values of the \mathbf{x}_i .

After the subsamples are created, lines 4-6 of Algorithm 1 compute \hat{q}^k , which summarizes the order that least angle regression includes each \mathbf{x}_i across all stages (see Figure 1). The unrestricted least angle regression ranks variables by the stage they enter the solution path. As shown in line 6 of Algorithm 1 and Figure 1, variables included at earlier stages have larger \hat{q}_i^k values: the first variable included is assigned 1, the last is assigned $1/\tilde{p}$, while the rejected variables are assigned 0 (which occurs only when $p > n$). Thus, the L_0 solution path is obtained by ranking the \mathbf{x}_i according to their \hat{q}_i^k values.

Zhang (2009, Theorem 2) implies that, on average, variables with the largest \hat{q}_i^k values are more likely to be informative. The \hat{q}_i^k may be sensitive in high-dimensional spaces to multicollinearity, sampling randomness, and noise. In these circumstances, a redundant variable may be included at an early stage in some (Y^k, X^k) subsample. Algorithm 1 reduces the impact of sensitivity in the \hat{q}_i^k by

Algorithm 1: \hat{q} method: parameterizing and averaging L_0 solution paths

input : (Y, X) .

- 1 divide the original sample equally into K folds and generate K subsamples $\{(Y^k, X^k)\}_{k=1}^K$ by removing one fold in turn from (Y, X) ;
- 2 set $\tilde{p} = \min\{n(K-1)/K, p\}$;
- 3 **for** $k := 1$ to K , *stepsize* = 1 **do**
- 4 run an unrestricted least angle regression (or any forward selection algorithm) on (Y^k, X^k) and record the order of variable inclusion at each stage;
- 5 define $\hat{q}^k = \mathbf{0} \in \mathbb{R}^p$;
- 6 $\forall i, l \in \mathbb{N}^+$, if \mathbf{x}_i is included at stage l and excluded at $l-1$, set $\hat{q}_i^k = (\tilde{p} + 1 - l)/\tilde{p}$, where \hat{q}_i^k is the i^{th} entry of \hat{q}^k ;
- 7 **end**
- 8 $\hat{q} := \frac{1}{K} \sum_{k=1}^K \hat{q}^k$;
- 9 **return** \hat{q}

computing $\hat{q} := \frac{1}{K} \sum_{k=1}^K \hat{q}^k$ and ranking the \mathbf{x}_i according to \hat{q}_i (the i^{th} entry in \hat{q}), to arrive at the average L_0 solution path. The average L_0 solution path is formally defined as follows.

Definition 2.2 (average L_0 solution path). Define the **average L_0 solution path** of least angle regression on $\{(Y^k, X^k)\}_{k=1}^K$ to be the (decreasing) rank order of the \mathbf{x}_i variables based on their corresponding \hat{q}_i values. For example, in Figure 1, the \hat{q}_i for $\mathbf{x}_1, \mathbf{x}_2$ and \mathbf{x}_3 are, respectively, $\hat{q}_1 = 5/6$, $\hat{q}_2 = 4/6$ and $\hat{q}_3 = 3/6$. Thus, the average L_0 solution path may be represented as an ordered set $\{\mathbf{x}_1, \mathbf{x}_2, \mathbf{x}_3\}$.

To justify theoretically the \hat{q} method, we use the Zhang (2009) framework to derive the theoretical properties of path averaging (see Appendix A).

- Under the Zhang (2009) conditions, Lemma A.1 shows that, with a high probability, using \hat{q}_i^k ranking for variable selection on (Y^k, X^k) generates the same theoretical results as the Zhang (2009) forward selection method.
- Under a similar stopping condition to Zhang (2009), Lemma A.2 shows that, with a high probability, there exists a threshold c^k for the L_0 path on (Y^k, X^k) such that $\hat{q}_i^k \geq c^k$ for informative \mathbf{x}_i and $\hat{q}_i^k < c^k$ for redundant \mathbf{x}_i
- Using Lemma A.2, Lemma A.3 shows that there exists a threshold $c = \sum_{i=k}^K c^k / K$ for the average L_0 path such that $\hat{q}_i \geq c$ for informative \mathbf{x}_i and $\hat{q}_i < c$ for redundant \mathbf{x}_i with large probability.

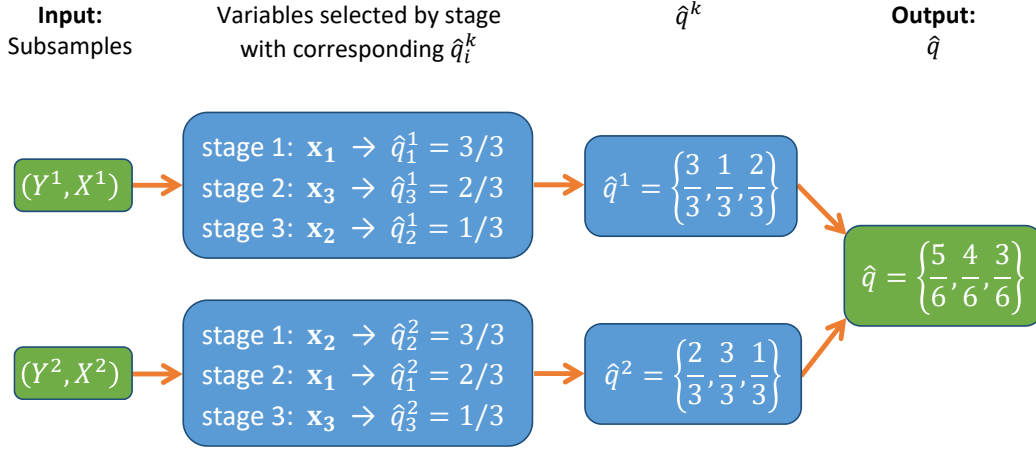


Figure 1: Computation of \hat{q} on 2 subsamples by least angle regression.

2.1.2. Variable selection on the average L_0 path

The solar algorithm is constructed on the average L_0 path and summarized in Algorithm 2. We present solar under the generic framework of forward regression. Solar is easily adapted to least angle regression, forward selection, and backward selection algorithms.

Algorithm 2: Subsample-ordered least-angle regression (solar)

- 1 Randomly select 20% of the sample points as the validation set; denote the remaining points as the training set;
 - 2 Estimate \hat{q} using Algorithm 1 on the training set and compute $Q(c) = \{\mathbf{x}_j \mid \hat{q}_j \geq c, \forall j\}$ for all $c \in \{1, 0.98, \dots, 0.02, 0\}$.
 - 3 Run an OLS regression of each $Q(c)$ on Y using the training set and find c^* , the value of c that minimizes the validation error;
 - 4 Compute the OLS coefficients of $Q(c^*)$ on Y using the whole sample.
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In Algorithm 2, variables are included into forward regression according to their rank order in the average L_0 solution path, represented by $\{Q(c) \mid c = 1, 0.98, \dots, 0\}$ in Algorithm 2. We use \hat{q} from Algorithm 1 to generate a list of variables $Q(c) = \{\mathbf{x}_j \mid \hat{q}_j \geq c, \forall j \leq p\}$. For any $c_1 > c_2$, $Q(c_1) \subset Q(c_2)$, implying a sequence of nested sets $\{Q(c) \mid c = 1, 0.98, \dots, 0\}$. Each c denotes a stage of forward regression. For a given value of c , $Q(c)$ denotes the set of variables with $\|\beta_i\|_0 = 1$ on average and $Q(c) - Q(c - 0.02)$ is the set of variables with $\|\beta_i\|_0$ just turning to 1 at c . Therefore, $\{Q(c) \mid c = 1, 0.98, \dots, 0\}$ is the average L_0 solution path of Definition 2.2. Variables that are more likely to be informative have larger c values in $Q(c)$ and will be selected first by the solar algorithm.

Using the Zhang (2009) framework and Lemmas A.2 and A.3, we derive the following theoretical results for variable selection (see Appendix A).

- Theorem A.2 shows that solar variable selection is L_0 consistent under similar sparse eigenvalue and irrerepresentable conditions used to prove lasso consistency.
- Under similar assumptions to Zhang (2009), Lemmas A.4 and A.5 show that the number of omitted informative \mathbf{x}_i and the probability of selecting at least one redundant \mathbf{x}_i are restricted by sample size, the sparse eigenvalue condition, and the stopping condition.

The key difference between solar and the lasso-type estimators, and the source of the advantages of solar, is solution path averaging.

- Lasso and solar both use the solution path for variable selection. The difference between solar and lasso is that solar averages the solution path. Lasso and its variants focus on optimizing the shrinkage parameter λ (via cross validation), ignoring concerns about the reliability of the lasso path in high dimensions. Optimizing λ on an unreliable path renders variable selection difficult. By contrast, solar prioritizes averaging the solution path, which not only averages out path unreliability in high dimensions, but also ranks all the informative variables at the start of the average L_0 path (as shown in Lemma A.2 and A.3). Hence, with a high probability, the variable selection algorithm needs only to analyze the variables at the start of the average L_0 path, making selection accurate and efficient.
- The difference between solar and lasso-related bootstrap selection (e.g., bolasso) is in how they average the variable selection algorithm. Given the λ value (optimal or not), lasso-related bootstrap selection averages the selection *results* across subsamples. Thus, bootstrap selection requires hundreds of repetitions to average out the instability and redundancy of lasso variable selection (Bach, 2008). By contrast, solar averages solution *paths*, which solves most of the lasso instability and redundancy issues, returning a more reliable average L_0 path. Variable selection along a reliable path substantially reduces the likelihood that solar selects redundant variables or omits informative variables. As a result, solar-related bootstrap selection (e.g., bootstrap solar or solar stability selection) requires only 3-5 repetitions to outperform hundreds of lasso-related bootstrap repetitions (see Section 4.6 below for details).

2.2. Solar optimized by coordinate descent

The solar algorithm can easily be generalized to use coordinate descent. For lasso, least angle regression or coordinate descent generates the same solution path parameterized by the β_i and the shrinkage parameter λ . Thus, to reprogram solar to use coordinate descent, we simply replace Algorithm 1 with Algorithm 3, which records the order of variable selection along the coordinate descent solution path.

Algorithm 3: average L_0 solution path estimation via coordinate descent

input : (Y, X) .

- 1 generate K subsamples $\{(Y^k, X^k)\}_{k=1}^K$ by randomly remove $1/K$ of observations in (Y, X) ;
- 2 set $\tilde{p} = \min\{n_{\text{sub}}, p\}$;
- 3 **for** $k := 1$ to K , $stepsize = 1$ **do**
- 4 denote λ_s as the shrinkage parameter value that coordinate descent lasso selects s variables,
 $\forall s \in [0, \tilde{p}]$;
- 5 run a pathwise coordinate descent for lasso on (Y^k, X^k) , $\forall \lambda \in \{\lambda_0, \lambda_1, \dots, \lambda_{\tilde{p}}, \}$
- 6 record the order of variable inclusion at each $\lambda \in \{\lambda_0, \lambda_1, \dots, \lambda_{\tilde{p}}, \}$;
- 7 define $\hat{q}^k = \mathbf{0} \in \mathbb{R}^p$;
- 8 $\forall i, s \in \mathbf{N}^+$, if \mathbf{x}_i is included at $\lambda = \lambda_s$ and excluded at λ_{s-1} , set $\hat{q}_i^k = (\tilde{p} + 1 - s)/\tilde{p}$, where \hat{q}_i^k is the
 i^{th} entry of \hat{q}^k ;
- 9 **end**
- 10 $\hat{q} := \frac{1}{K} \sum_{k=1}^K \hat{q}^k$;
- 11 **return** \hat{q}

Algorithm 3 serves the same purpose as Algorithm 1: to estimate the average L_0 path. Algorithm 3 uses λ to record the order that each variable enters the path. Consider the example in Figure 2. To re-parameterize the solution path, we denote λ_s to be the λ value that coordinate descent lasso includes s variables, $\forall s \in (0, \min\{n/2, p\}]$, giving a sequence of λ for grid search. In each subsample (Y^k, X^k) , we train a standard pathwise coordinate descent for lasso, allowing λ to increase stepwise within the grid $\{\lambda_1, \dots, \lambda_{\min\{n/2, p\}}\}$, where $\lambda_1 \geq \dots \geq \lambda_{\min\{n/2, p\}}$. In Figure 2, when $\lambda \leq \lambda_3$ at subsample (Y^1, X^1) , all three variables are selected in the solution path, implying that $\hat{q}_i^1 \geq 1/3$ for all variables. When λ increases to λ_2 , only $\{\mathbf{x}_3, \mathbf{x}_1\}$ survive the harsher shrinkage, implying that they should be ranked higher than \mathbf{x}_2 . As a result, $\hat{q}_1^1, \hat{q}_3^1 \geq 2/3$ and $\hat{q}_2^1 = 1/3$. When λ reaches λ_3 , only $\{\mathbf{x}_1\}$ remains, leaving $\hat{q}_1^1 = 3/3$ and $\hat{q}_3^1 = 2/3$. Applying the same method to each subsample produces the same \hat{q} as Algorithm 1.

2.3. Comparison and generalization to lasso variants

Because solar is trained by least angle regression or coordinate descent, it can easily be extended to several lasso variants:

- ‘Grouped solar’ is invoked by forcing specific variables to be simultaneously selected into the solution path;
- ‘Adaptive solar’ is obtained by weighting variable rankings in the average L_0 path according to their OLS coefficients;

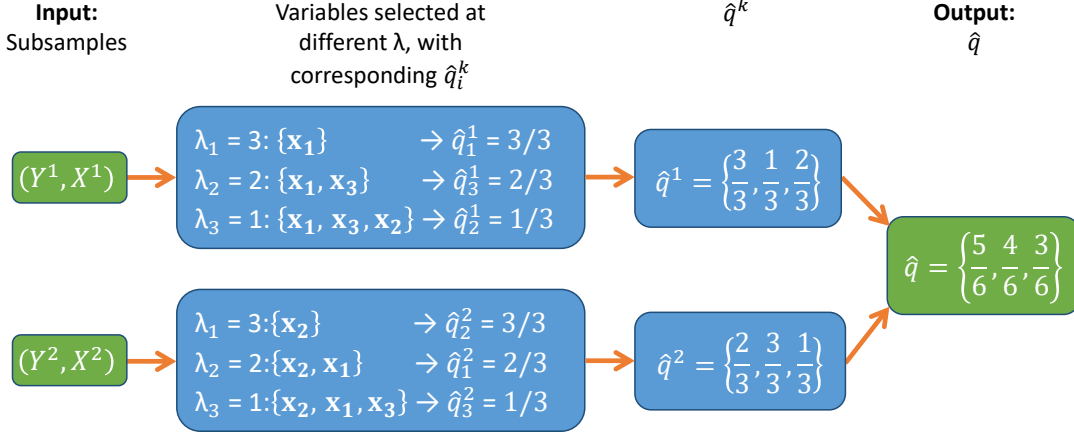


Figure 2: Computation of \hat{q} on 2 subsamples using coordinate descent.

- ‘Solar elastic net’ or ‘fused solar’ is derived by replacing the coordinate descent loss function in Algorithm 3 with the L_1 - L_2 loss

$$\|Y - X\beta\|_2^2 + \lambda^{(1)} \|\beta\|_1 + \lambda^{(2)} \|\beta\|_2^2 \quad (2.1)$$

or fused loss

$$\|Y - X\beta\|_2^2 + \lambda^{(1)} \|\beta\|_1 + \lambda^{(2)} \sum_{j=2}^p |\beta_j - \beta_{j-1}|_1. \quad (2.2)$$

Furthermore, many lasso enhancements (e.g., safe/strong rules, post-lasso hypothesis testing) may be applied to solar because they use the same optimization methods. Rather than competing with the lasso enhancements, solar supplements them by improving variable selection performance and computation speed in large-scale applications.

3. Solar advantages over lasso variants, lasso rules, and variable screening

In this section, we use a series of examples to demonstrate the advantages of the solar algorithm for post-selection hypothesis testing, in the presence of complicated dependence structures, and in terms of its robustness to the *irrepresentable condition* (IRC).

3.1. Post-selection hypothesis testing

A major advantage of solar is its amenability to post-selection testing. Because the lasso tests (Lockhart et al., 2014; Taylor et al., 2014) are based on forward regression, they may be adapted to solar. More interestingly, by replacing lasso with solar in the post-selection tests (e.g, the data-splitting tests) (Wasserman and Roeder, 2009; Meinshausen et al., 2009), we substantially improve testing accuracy for weak signal detection with limited n . We illustrate this point using Example 1.

Example 1. Consider the DGP

$$Y = \mathbf{x}_0 + 2\mathbf{x}_1 + 3\mathbf{x}_2 + 4\mathbf{x}_3 + 5\mathbf{x}_4 + \sum_{j=5}^p 0 \cdot \mathbf{x}_j + e, \quad (3.1)$$

where \mathbf{x}_i , $i = 0, \dots, p$, are standard Gaussian variables with pairwise correlations of 0.5, e is a standard Gaussian noise term, and $p/n = 140/120$.

Following Romano and DiCiccio (2019, Example 4.1) and DiCiccio et al. (2020), we conduct data-splitting tests by randomly separating the data into two portions of 60 observations. In the first round, one portion is used for solar or lasso selection and the other for testing. In the second round, the roles of the two portions are reversed. As a result, the p-values of any given variable are uncorrelated across the two rounds. Thus, we may apply Theorem 3.2 of Romano and DiCiccio (2019) and compute the average p-value across the two rounds to conduct a valid t-test for any selected covariate.

DiCiccio et al. (2020) stresses the importance of retaining residual degrees of freedom to ensure accurate tests. Because solar yields a more sparse and accurate variable selection than lasso (Section 4 below), it conserves residual degrees of freedom, improving the reliability of post-selection p-values. Figure 3 plots the average p-values for the informative variables $\{\mathbf{x}_0, \dots, \mathbf{x}_4\}$ from post-solar and post-lasso data-splitting tests using 100 repetitions. While the solar and lasso p-values are less than 0.05 for the stronger signals $\{\mathbf{x}_1, \dots, \mathbf{x}_4\}$, more than 25% of the lasso p-values exceed 0.05 for the weakest signal \mathbf{x}_0 , implying non-trivial false non-rejection of H_0 . By contrast, the solar p-value boxplot is very compact for \mathbf{x}_0 , with only 5 out of 100 above 0.05. Hence, solar p-values are more reliable for detecting weak signals with small n and large p .

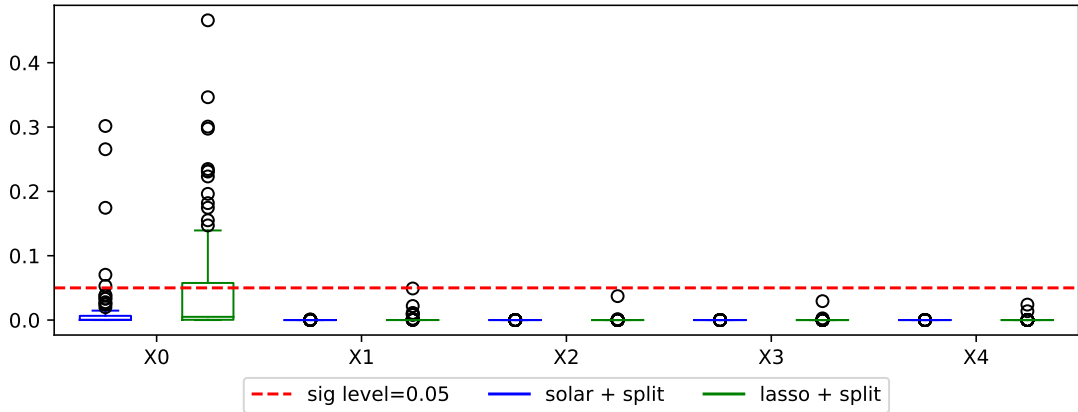


Figure 3: Average p-value boxplots for data-splitting t-tests with solar and lasso.

The solar L_0 path may also assist with the formulation of H_0 for $p > n$. Because conserving residual degrees of freedom is so important, tests on the selection (omission) of redundant (informative) variables trigger decisions on which β_i to test. Zhang (2009, Theorem 2) shows that the earlier a variable enters the L_0 path, the more likely it is informative, implying that variables should be tested

in rank order. Given the solar path is more robust than lasso path to settings of the irrerepresentable condition, sampling noise, multicollinearity, and other issues, it provides more reliable guidance on the order to test the β_i . ■

3.2. Complicated dependence structures

Another advantage of solar is that the average L_0 solution path is more robust to outliers, multicollinearity, and noise in high-dimensional spaces. Thus, solar is likely to be more reliable than other variable selection methods under complicated dependence structures. We illustrate the point with the following two (Bayesian network) examples.

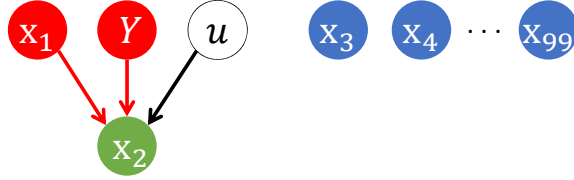


Figure 4: Y is unconditionally uncorrelated with an informative x_1 .

The first example is a common empirical regression problem: *informative variables* that are *unconditionally uncorrelated to Y* in the DGP. In Figure 4, x_1 and x_2 are informative for Y [see the derivation of (3.2)-(3.3)], while x_1 and Y are independent. For example, in biostatistics, concussion (x_1) or a brain tumor (Y) may cause headaches (x_2), implying that concussion history is necessary when attempting to diagnose a brain tumor. In this setting, Example 2a shows that solar is more reliable than post-lasso rules and variable screening.

Example 2a. In Figure 4, there are 100 variables and x_2 is (causally) generated by its parents $\{x_1, Y\}$ as follows,

$$x_2 = \alpha_1 x_1 + \alpha_2 Y + u, \quad (3.2)$$

where x_1 is unconditionally uncorrelated with Y , x_1 and Y are both unconditionally and conditionally uncorrelated with the redundant variables $\{x_3, \dots, x_{99}\}$, $\{\alpha_1, \alpha_2\}$ are population regression coefficients, and u is a Gaussian noise term. If Y is chosen to be the response variable, the population regression equation is

$$Y = -\frac{\alpha_1}{\alpha_2} x_1 + \frac{1}{\alpha_2} x_2 - \frac{1}{\alpha_2} u. \quad (3.3)$$

Note that x_1 and x_2 are both informative for Y . However, since x_1 is unconditionally uncorrelated with Y in the population, the post-lasso rules—such as the strong rule (Tibshirani et al., 2012) and the safe rule (Ghaoui et al., 2010)—may be prone to rejecting x_1 . For a given value of the shrinkage parameter λ in grid search, the base strong rule and the safe rule for lasso to reject a selected variable,

respectively, satisfies (3.4) and (3.5):

$$|\mathbf{x}_i^T Y| < \lambda - \|\mathbf{x}_i\|_2 \|Y\|_2 \frac{\lambda_{max} - \lambda}{\lambda_{max}}; \quad (3.4)$$

$$|\mathbf{x}_i^T Y| < 2\lambda - \lambda_{max}, \quad (3.5)$$

where the \mathbf{x}_i are standardized and λ_{max} is the value of the shrinkage parameter that rejects all the variables. Both rules are based on the unconditional covariance between \mathbf{x}_i and Y . For a given value of λ (typically selected by CV), lasso will likely select \mathbf{x}_1 and \mathbf{x}_2 along with redundant variables from $\{\mathbf{x}_3, \dots, \mathbf{x}_{99}\}$ because the DGP does not violate the IRC. Since $\text{corr}(\mathbf{x}_1, Y) = \text{corr}(\mathbf{x}_3, Y) = \dots = \text{corr}(\mathbf{x}_{99}, Y) = 0$ in the population, the sample value of $|\mathbf{x}_1^T Y|$ will be approximately as small as the $|\mathbf{x}_i^T Y|$ of any redundant variable. Put another way, \mathbf{x}_1 cannot be distinguished from the redundant variables by the value of $|\mathbf{x}_i^T Y|$. To ensure \mathbf{x}_1 is not rejected by (3.4) or (3.5), both $\lambda - \|\mathbf{x}_1\|_2 \|Y\|_2 \frac{\lambda_{max} - \lambda}{\lambda_{max}}$ and $2\lambda - \lambda_{max}$ must be smaller than $|\mathbf{x}_1^T Y|$. However, this will lead to two problems. First, decreasing the right-hand side of (3.4) and (3.5) will reduce the value of λ , implying that lasso will select more redundant variables. Second, since $|\mathbf{x}_1^T Y|$ will be approximately as small as the $|\mathbf{x}_i^T Y|$ of any redundant variable selected by lasso, not rejecting \mathbf{x}_1 (by reducing both right-hand side terms) may result in (3.4) and (3.5) retaining redundant variables.

Variable screening methods (Fan and Lv, 2008) may also be prone to selecting redundant variables. Screening ranks variables decreasingly based on the absolute values of their unconditional correlations to Y , selecting the top w variables (with w selected by CV, bootstrap, or BIC). Since $\text{corr}(\mathbf{x}_2, Y) \neq 0$ in the population, screening will rank \mathbf{x}_2 highly. However, it may not rank \mathbf{x}_1 highly because $\text{corr}(\mathbf{x}_1, Y) = 0$ in the population. Thus, some redundant variables may be ranked between \mathbf{x}_2 and \mathbf{x}_1 , implying that if both \mathbf{x}_1 and \mathbf{x}_2 are selected, screening will select redundant variables.

The average L_0 solution path will not suffer the same problems. For convenience, assume $-\alpha_1/\alpha_2 > 0$ and $p/n = 100/200$ or smaller. For least angle regression, as $\|\beta_2\|_1$ increases at stage 1 (i.e., as \mathbf{x}_2 is ‘partialled out’ of Y), the unconditional correlation between $Y - \beta_2 \mathbf{x}_2$ and \mathbf{x}_1 will increase above 0 significantly while the marginal correlation between $Y - \beta_2 \mathbf{x}_2$ and any redundant variable will remain approximately 0. Thus, in the L_0 solution path and, hence, the average L_0 solution path, \mathbf{x}_1 will be included immediately after \mathbf{x}_2 is included. ■

Fan and Lv (2008) and Barut et al. (2016) propose two solutions for variable screening in situations like Example 2a. However,

- the first approach (Barut et al., 2016, Section 2.2 and 3) assumes the identity of \mathbf{x}_2 is known, which is unlikely to be realistic in practical applications. [In Bayesian networks or probabilistic graph modelling, \mathbf{x}_2 is known as a *collider*; Barut et al. (2016) refer to \mathbf{x}_2 as a *hidden signature* variable and denote it by X_c];

- the second approach (Barut et al., 2016, Section 1 and 2.2) suggests randomly trying out variables to be colliders. The logic is straightforward: randomly trying out a wrong variable (like \mathbf{x}_2) to be a collider is harmless because conditioning on that variable will not make $\text{corr}(Y, \mathbf{x}_1) \neq 0$, nor will it cause the selection of a redundant variable. Moreover, by repeatedly randomly trying out variables, there is a non-zero probability the correct collider will eventually be uncovered, producing a statistically significant $\text{corr}(Y, \mathbf{x}_1) \neq 0$. Since using multiple trials may be inefficient and computationally expensive, especially with high-dimensional data, Barut et al. (2016) suggests trying out several variables simultaneously. However, if $\text{corr}(Y, \mathbf{x}_1) \neq 0$ were discovered after trying out, say, $\{\mathbf{x}_2, \text{other variables}\}$, it would still be necessary to decide which of $\{\mathbf{x}_2, \text{other variables}\}$ are redundant, meaning variable selection is not completed.

A related example illustrates another common empirical regression problem: *redundant variables* that are *unconditionally correlated to Y* in the DGP. In Figure 5, the problem occurs because \mathbf{x}_3 and Y are determined by common variables. For example, house rent (Y) and food expenditure (\mathbf{x}_3) are both determined by income (\mathbf{x}_1) and saving (\mathbf{x}_2), yet \mathbf{x}_3 is redundant if \mathbf{x}_1 and \mathbf{x}_2 are used to predict Y . In this setting, Example 2b illustrates that the strong rule, base rule, and variable screening methods may struggle to reject the redundant \mathbf{x}_3 even when the IRC is satisfied. By contrast, solar will be less prone to selecting redundant variables.

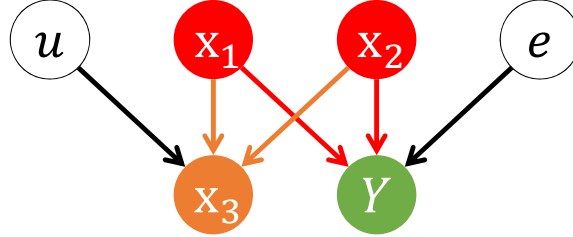


Figure 5: Y is unconditionally correlated with a redundant \mathbf{x}_3 .

Example 2b. Figure 5 depicts the following confounding structure,

$$\begin{cases} \mathbf{x}_3 = \frac{1}{3}\mathbf{x}_1 + \frac{1}{3}\mathbf{x}_2 + \frac{\sqrt{7}}{3}u, \\ Y = \frac{7}{10}\mathbf{x}_1 + \frac{2}{10}\mathbf{x}_2 + \frac{\sqrt{47}}{10}e, \end{cases} \quad (3.6)$$

where \mathbf{x}_1 and \mathbf{x}_2 cause both Y and \mathbf{x}_3 , implying that \mathbf{x}_3 is unconditionally correlated to Y ; \mathbf{x}_1 , \mathbf{x}_2 , u and e are independent; \mathbf{x}_3 is independent from e ; Y is independent from u ; and all variables are standardized.

For large n , when the sample correlations are close to their population values, the sample marginal

correlations to Y are:

$$\begin{aligned}\text{corr}(\mathbf{x}_1, Y) &= 0.7, \\ \text{corr}(\mathbf{x}_3, Y) &= \text{corr}\left(\frac{1}{3}\mathbf{x}_1 + \frac{1}{3}\mathbf{x}_2, \frac{7}{10}\mathbf{x}_1 + \frac{2}{10}\mathbf{x}_2\right) = 0.3, \\ \text{corr}(\mathbf{x}_2, Y) &= 0.2.\end{aligned}\tag{3.7}$$

Because \mathbf{x}_2 ranks below \mathbf{x}_1 and \mathbf{x}_3 in terms of marginal correlations to Y , the variable screening method must select all 3 variables—including the redundant \mathbf{x}_3 —to avoid omitting \mathbf{x}_2 . The base strong rule and safe rule may also have difficulty rejecting \mathbf{x}_3 . Since $\text{corr}(\mathbf{x}_3, Y) > \text{corr}(\mathbf{x}_2, Y)$, if lasso selects \mathbf{x}_3 and \mathbf{x}_2 and the strong (or safe) rule is used to reject \mathbf{x}_3 , \mathbf{x}_2 will also be rejected.

Forward regression, solar, and lasso will not make the same error. Because (3.6) does not violate the IRC, variable-selection consistency of forward regression, lars, and lasso is assured from the theoretical results of Zhang (2009) and Zhao and Yu (2006). In forward regression, \mathbf{x}_1 will be included at the first stage. After controlling for \mathbf{x}_1 , the partial correlations (for large n) of both \mathbf{x}_2 and \mathbf{x}_3 with Y are:

$$\begin{aligned}\text{corr}(\mathbf{x}_2, Y|\mathbf{x}_1) &= \text{corr}\left(\mathbf{x}_2, \frac{2}{10}\mathbf{x}_2\right) = 0.2, \\ \text{corr}(\mathbf{x}_3, Y|\mathbf{x}_1) &= \text{corr}\left(\frac{1}{3}\mathbf{x}_1 + \frac{1}{3}\mathbf{x}_2, \frac{2}{10}\mathbf{x}_2\right) = 0.0667.\end{aligned}\tag{3.8}$$

Because $\text{corr}(\mathbf{x}_2, Y|\mathbf{x}_1) > \text{corr}(\mathbf{x}_3, Y|\mathbf{x}_1)$, forward regression will include \mathbf{x}_2 , not \mathbf{x}_3 , at the second stage. After controlling for \mathbf{x}_1 and \mathbf{x}_2 , remaining variation in Y is due to e , which \mathbf{x}_3 cannot explain. Thus, CV or BIC will terminate forward regression after the second stage and \mathbf{x}_3 will not be selected. Similarly, because solar relies on the average L_0 path, it will include \mathbf{x}_1 and \mathbf{x}_2 but not \mathbf{x}_3 . ■

Essentially, the strong rule, safe rule, and variable screening struggle in Examples 2a and 2b because they rely on unconditional correlations to Y , whereas informative variables in regression analysis are defined in terms of conditional correlations. In many scenarios, unconditional and conditional correlations are aligned. However, when they are not, variable selection based conditional correlation is better placed to select the informative variables.

Fan and Lv (2008) propose to redeem variable screening on Y by first selecting variables with high unconditional correlations to Y and then running a lasso of the residuals on the dropped variables. By contrast, solar completes variable selection in a single pass of conditional correlation ranking, reducing computational costs. Moreover, the Fan and Lv (2008) approach does not solve Example 2b type problems. At the first step, variables with high unconditional correlations to Y will be selected, including the redundant \mathbf{x}_3 . Selecting redundant variables will be more serious when Y has multiple \mathbf{x}_3 -like siblings and in complicated dependence structures where multicollinearity results in inaccurate estimates of the coefficients and standard errors in finite samples. In short, solar is likely to be more

computationally efficient and better at variable selection in settings with complicated dependence structures.

3.3. Robustness to the IRC

Solar is more robust than lasso to different settings of the IRC. The IRC is considered to be sufficient and almost necessary for accurate lasso variable selection (Zhang, 2009). Here, we ignore lasso rules and variable screening since, as discussed above, their selection accuracy may be compromised by reliance on unconditional correlations to Y . We define the IRC as in Zhang (2009).

Definition 3.1 (IRC). Given $F \subset \{1, \dots, p\}$, define X_F to be the $n \times |F|$ matrix with only the full set of informative variables. Define

$$\mu(F) = \max \left\{ \left\| \left((X_F)^T X_F \right)^{-1} (X_F)^T \mathbf{x}_j \right\|_1 \mid \forall j \notin F \right\}.$$

Given a constant $0 < \eta \leq 1$, the *strong* irrerepresentable condition is satisfied if $\mu(F) \leq 1 - \eta$ and the *weak* irrerepresentable condition is satisfied if $\mu(F) < 1$. ■

Example 3. Modify the DGP in Example 2b to match the Zhao and Yu (2006) simulations. Thus, $n = 200$, $p = 50$, and $\{\mathbf{x}_0, \dots, \mathbf{x}_4, \mathbf{x}_6, \dots, \mathbf{x}_{50}\}$ are generated from a zero-mean, unit-variance multivariate Gaussian distribution, where all the correlation coefficients are 0.5. The DGP of Y and \mathbf{x}_5 is

$$\begin{cases} \mathbf{x}_5 = \omega \mathbf{x}_0 + \omega \mathbf{x}_1 + \gamma \cdot \sqrt{1 - 2\omega^2} \\ Y = 2\mathbf{x}_0 + 3\mathbf{x}_1 + 4\mathbf{x}_2 + 5\mathbf{x}_3 + 6\mathbf{x}_4 + e \end{cases} \quad (3.9)$$

where $\omega \in \mathbb{R}$, while γ and e are both standard Gaussian noise terms, independent from one another and all the other variables. Compared with Example 2b, this DGP increases the challenge of accurate selection by increasing the number of redundant variables from 1 to 46, $\{\mathbf{x}_5, \dots, \mathbf{x}_{50}\}$. This DGP also makes it straightforward to control the IRC through ω , which affects the value of $\mu(F)$.

In (3.9), the IRC only affects the redundant \mathbf{x}_5 . Hence, we focus on the probability of incorrectly selecting \mathbf{x}_5 in 200 repetitions. By setting ω to either 1/4, 1/3, or 1/2, the population value of $\mu(F)$ changes, respectively, to 1/2, 2/3, or 1, gradually increasing the difficulty of rejecting the redundant \mathbf{x}_5 .

Figure 6 displays the simulation results. When $\mu(F) = 1/2$, lasso wrongly includes \mathbf{x}_5 with probability more than 0.25. By contrast, the probability solar selects \mathbf{x}_5 is less than 0.1. When $\mu(F)$ increases to 2/3, the probability lasso includes \mathbf{x}_5 increases to around 0.3. When $\mu(F)$ increases to 1 in the population and strong IRC is violated, the probability lasso includes \mathbf{x}_5 rises to almost 0.5. By contrast, the probability solar includes \mathbf{x}_5 only increases incrementally and stays much lower than the lasso probability. The results illustrate that solar is more robust to different settings of the IRC. ■

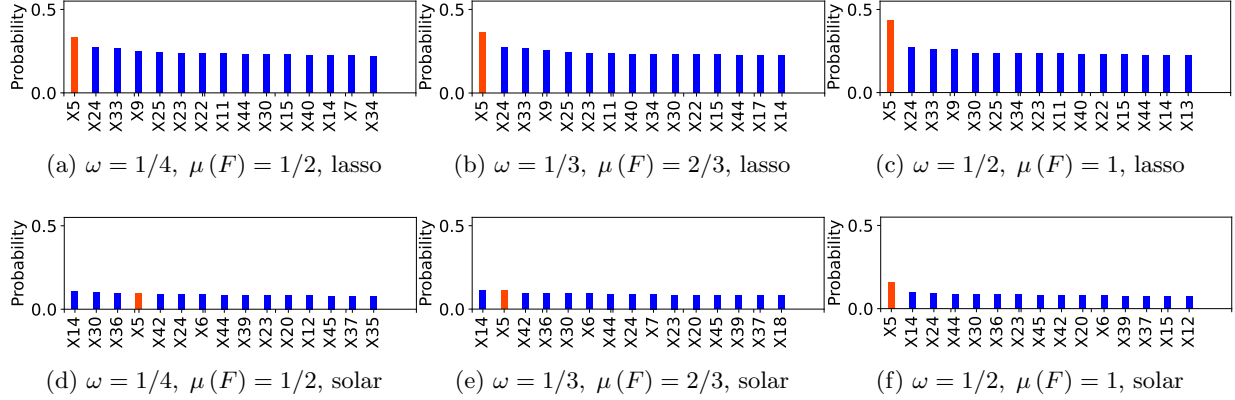


Figure 6: Probability of including redundant variables (top 15) in simulation 2 (\mathbf{x}_5 in orange).

4. Solar advantages over subsample variable selection

In this section, we shift our focus to simulation. We demonstrate that: (i) solar offers significant improvements over lasso-type algorithms in terms of variable selection sparsity and accuracy; (ii) replacing lasso with solar in bootstrap selection drastically reduces the computation load (i.e., runtime). We choose the simulation settings so that, as far as possible, the comparisons are fair, representative, and generalizable. Our overall goal is to enable *ceteris paribus* comparisons between solar and state-of-the-art lasso algorithms.

4.1. Simulation competitors

We consider a subset of lasso-type algorithms for comparison to solar. Firstly, some lasso modifications (e.g., fused lasso, grouped lasso) are designed to solve specific empirical problems that are not relevant to our paper. Secondly, it may be difficult to determine how much some variants outperform lasso.¹ Since both solar and lasso may be evaluated via least angle regression and coordinate descent, many other lasso modifications can be directly applied to solar, as discussed in Section 2.3. We do not consider information criteria for shrinkage parameter tuning. Pedregosa et al. (2011) points out that information criteria are over-optimistic and require a proper estimation of the degrees of freedom for the solution. Moreover, information criteria are derived asymptotically and tend to break down when the problem is badly conditioned (e.g., $p > n$).²

¹For example, while Jia and Yu (2010) show numerically that elastic net has slightly better variable-selection accuracy than lasso, they also find that “when the lasso does not select the true model, it is more likely that the elastic net does not select the true model either” (a point we verify in Section 5). While simulations in Zou (2006) show that adaptive lasso outperforms lasso when $p/n < 1$, it requires first computing the OLS estimates of all \mathbf{x}_i coefficients, which is difficult when $p/n > 1$.

²See https://scikit-learn.org/stable/modules/linear_model#lasso.html for details.

Solar competes with 10-fold, cross-validated lasso (denoted ‘lasso’ for short), following the [Friedman et al. \(2001\)](#) simulations that show 10 folds balances the bias-variance trade-off in CV error minimization. We set the number of generated subsamples (K) in Algorithm 1 to 3 since $K > 3$ has only negligible effects. Because least-angle regression and coordinate descent yield similar selection results for solar and lasso, we combine the lars and coordinate descent results for solar and report only the runtime for lars lasso (see Supplementary Material B).

We also include bootstrap selection algorithms in the comparisons. A bootstrap selection repeats lasso multiple times across bootstrap subsamples to produce a set of averaged (or accumulated) selection results. Given the similarities among lasso bootstrap selection methods, we choose the [Bach \(2008\)](#) bootstrap lasso (*bolasso*) to compete with solar. [Bach \(2008\)](#) proposes two bolasso algorithms: bolasso-H and bolasso-S; both are competitors in the simulations. Bolasso-H selects only variables that are selected in all bootstrap subsamples, i.e., the subsample selection frequency threshold, $f = 1$. Bolasso-S selects variables that are selected in 90% of the bootstrap subsamples ($f = 0.9$). [Bach \(2008\)](#) finds that bolasso selection and prediction performance improves with the number of subsamples. To ensure a vigorous challenge for solar, we set the number of bootstrap subsamples in bolasso to 256, the maximum in the [Bach \(2008\)](#) simulations. To return an accurate selection result, bolasso requires optimizing the λ value on each bootstrap subsample.³ Since the optimal value of λ on each (sub)sample may vary substantially in high dimensions, we use 10-fold CV to tune λ in each bootstrap subsample ([Friedman et al., 2001](#)).

We also consider bootstrap solar selection (*bsolar*), which executes solar on each bootstrap subsample and computes the selection frequency for each variable across all bootstrap subsamples. To ensure that any performance difference is due solely to replacing lasso with solar in the bootstrap selection system, bolasso and bsolar use the same subsample selection frequency threshold. Thus, we evaluate 2 versions of bsolar: bsolar-H ($f = 1$) and bsolar-S ($f = 0.9$). We use the notation bsolar- m H and bsolar- m S, where m is the number of subsamples used to compute the selection frequency.

4.2. Simulation settings

The DGP for the simulations is as follows. The p covariates in $X \in \mathbb{R}^{n \times p}$ are generated from a zero-mean, multivariate Gaussian distribution, with all off-diagonal elements in the covariance matrix equal to 0.5. The first 5 variables in X are informative; the remaining $p - 5$ variables are redundant. The response variable $Y \in \mathbb{R}^{n \times 1}$ is:

$$Y = 2\mathbf{x}_0 + 3\mathbf{x}_1 + 4\mathbf{x}_2 + 5\mathbf{x}_3 + 6\mathbf{x}_4 + e, \quad (4.1)$$

³The hyperparameter optimization on bootstrap subsamples is crucial for different bootstrap selection methods. For example, if λ is unnecessarily large on more than 10% of the bootstrap subsamples, lasso may wrongly drop some informative variables in those subsamples, implying that bolasso-H and bolasso-S will omit informative variables.

where $e \in \mathbb{R}^{n \times 1}$ is a standard Gaussian noise term. All data points are independently and identically distributed. Each \mathbf{x}_i , $i = 1, \dots, p$, is independent from the noise term e , which is standard Gaussian. Simulations are repeated 200 times with fixed Python random generators across simulations.

We vary the data dimensions p/n as follows. In the first block of simulations, p/n approaches 0 from above, corresponding to the classical $p < n$ setting. In the second block, p/n approaches 1 from above, corresponding to high dimensional settings. In the third block, $p/n = 2$ as $\log(p)/n$ slowly approaches 0, corresponding to ultrahigh dimensional settings, i.e., where $(p - n) \rightarrow \infty$.

We compare the performance of solar and lasso in terms of sparsity and accuracy of variable selection as well as runtime. Sparsity is measured by the mean number of variables selected. Discovery accuracy is measured by the mean number of *informative* variables selected. Purge accuracy is measured by the mean number of *redundant* variables selected (equal to sparsity minus discovery accuracy). Runtime is measured by mean CPU time. The raw simulation results are available in Supplementary Material B.

4.3. Programming languages, parallelization, and hardware

To ensure a credible comparison between solar and the lasso competitors, we choose the hardware and software to maximize the computation speed of lasso. We show that, even under the ideal computation environment for lasso, solar exhibits a substantial runtime advantage.

To maximize computation speed, we use `Numpy`, `Scipy`, and `Cython`—all well-known for performance and speed—to outsource all numerical and matrix operations to the Intel Math Kernel Library, currently the fastest and most accurate C++/Fortran library for CPU numerical operations.

To reduce the possibility of CPU and RAM bottlenecks in parallel computing of lasso and bolasso, we code in Python rather than R.⁴ Given the simulations repeat solar, lasso, and bolasso many times to arrive at representative performance measures, choosing Python over R also mitigates the impact of hardware limitations. Computations are executed using an Intel Xeon W-3245 CPU with a base frequency of 3.2GHz and 64GB RAM, further reducing the possibility of CPU-RAM bottlenecks.

To guarantee the programming quality of the lasso implementation, we source lasso and bolasso from the Sci-kit learn library (Pedregosa et al., 2011) of efficient machine-learning tools.⁵ Used widely in research and industry, Sci-kit learn also uses `Numpy`, `Scipy`, and `Cython` to delegate all numerical and matrix operations to Fortran/C++.

Lastly, to optimize computation and avoid large overheads, we implement multi-core parallelization. Each realization of lasso requires 10 repetitions of lars or coordinate descent to compute the CV error

⁴Donoho (2017) claims: “R has the well-known reputation of being less scalable than Python to large problem sizes.”

⁵Detail is available at <https://scikit-learn.org/stable/>.

of each λ value. Thus, we design a parallel architecture to assign one repetition per CPU core, maximizing the computation speed for lasso.

4.4. Comparison of sparsity, accuracy, and time complexity

Table 1 summarizes average selection performance.⁶ While all competitors always include the 5 informative variables, solar outperforms lasso in terms of sparsity in every p/n scenario, implying superior ability to limit the selection of redundant variables. Notably, as $p/n \rightarrow 1$, lasso sparsity deteriorates while solar sparsity improves, further confirming the advantage of path averaging. While the sparsity of all competitors deteriorates as $\log(p)/n \rightarrow 0$, solar maintains a clear advantage over lasso.

Table 1: Simulation results for sparsity and accuracy.

	$p/n \rightarrow 0$			$p/n \rightarrow 1$			$\log(p)/n \rightarrow 0$		
	$\frac{100}{100}$	$\frac{100}{150}$	$\frac{100}{200}$	$\frac{150}{100}$	$\frac{200}{150}$	$\frac{250}{200}$	$\frac{400}{200}$	$\frac{800}{400}$	$\frac{1200}{600}$
<i>mean number of selected variables</i>									
lasso	20.4	19.5	19.7	23.1	24.1	27.2	30.7	36.7	37.4
solar	10.5	9.3	9.1	10.7	9.8	8.7	11.4	16.1	18.5
bolasso-S	5.5	6.4	6.5	5.5	6.4	6.5	5.7	6.6	7.6
bolasso-H	5	5	5	5	5	5	5	5	5
bsolar-3S/3H	5.4	5.2	5.1	5.4	5.2	5.1	5.3	5.8	6
bsolar-5S/5H	5.2	5.1	5	5.2	5.1	5	5.1	5.2	5.4
bsolar-10S	5.2	5.1	5	5.2	5.1	5	5.1	5.2	5.3
bsolar-10H	5	5	5	5	5	5	5	5	5.1
<i>mean number of selected informative variables</i>									
lasso	5	5	5	5	5	5	5	5	5
solar	5	5	5	5	5	5	5	5	5
bolasso-S/H	5	5	5	5	5	5	5	5	5
bsolar-3S/3H/5S/5H/10S/10H	5	5	5	5	5	5	5	5	5

Table 1 also reveals several advantages of solar over lasso in bootstrap selection.

- In terms of variable selection, bolasso-S stands out with the poorest sparsity; the others perform almost identically.
- Solar and bsolar display a considerable computational advantage. We show in Section 4.6 that solar imposes less than 1/3 of the lasso computation load, implying that bsolar-3 has the same computation load as lasso. Given bolasso requires 256 subsample lasso repetitions while bsolar-3 has the same computation load as one lasso realization, bsolar reduces subsample repetitions by 99% relative to bolasso (assuming a time complexity measure like $O(n^2)$ and $p > n$ for lasso).

⁶Detailed histograms are available in Supplementary Material B.

- Similar findings apply to the comparison between bsolar and lasso stability selection. Bsolar, bolasso-H, and stability selection ($f > 0.9$) push their selection sparsity and accuracy to the maximum (on average selecting all the informative variables while rarely including a redundant variable). However, lasso stability selection implements 100 subsample repetitions while bsolar-3 requires only 3. Even though the size of the bootstrap subsample in stability selection is $n/2$ (substantially smaller than the bootstrap sample size of bsolar, which is n), the time complexity analysis of [Meinshausen and Bühlmann \(2010\)](#) still implies that bsolar-3 produces a reduction of at least 67-82% in computation load relative to lasso stability selection. Such a computation time reduction is crucial in large scale applications like DNA sequencing, natural language processing, image processing, and MRI neuro-imaging.⁷ The computation reduction may be even larger if the application requires a ‘group’, ‘fused’, or ‘elastic net’ variant (discussed in [Section 2.3](#)).

4.5. Explanation of the efficiency difference between bolasso and bsolar

The efficiency of bsolar is due to a unique two-layer variable ranking scheme. While bolasso and bsolar both generate bootstrap subsamples, the two-layer procedure in bsolar reduces the number of bootstrap repetitions without compromising selection performance. Specifically,

- bsolar executes [Algorithm 1](#) (or [3](#)) on each bootstrap subsample and ranks variables using the average L_0 path, which we call the *internal ranking*. The internal ranking identifies the strongest signals on each bootstrap subsample.
- bsolar collates the internal ranking results to produce an overall ranking, which we call the *external ranking*. The external ranking identifies the strongest signals across the majority of bootstrap subsamples.

The two-layer method has several advantages over one-layer ranking methods inherent in bolasso and lasso stability selection ([Fan and Lv, 2008](#); [Hall et al., 2009](#); [Hall and Miller, 2009](#); [Li et al., 2012a,b](#)).

- First, one-layer methods rank variables on the whole sample. By contrast, the internal ranking uses the average L_0 path, which, as discussed in [Section 2.1](#), improves robustness to multicollinearity, noise, and sample size.
- Second, as shown in [Section 3.2](#), internal ranking avoids issues caused by complicated dependence structures that other (unconditional) ranking methods cannot.

⁷For example, each MRI image (observation) often contains more than 10^6 pixels (candidate variables), and the total data size can easily exceed 1TB even with limited n .

- Most important, multi-layer ranking reduces the number of bootstrap repetitions without compromising accuracy. One-layer methods select variables immediately after ranking. Our method performs a second external ranking that, by detecting persistent signals, is more tolerant of subsample variation: if \mathbf{x}_i is wrongly selected or omitted in the internal ranking, there is still a large probability that the mistake will be corrected in the external ranking. While stability selection and bolasso require, respectively, 100 and 256 repetitions to average out lasso selection issues, bsolar requires only 3-10 bootstrap repetitions to confirm the solar variable ranking.

Table 2: Subsample variable selection frequencies for bolasso and bsolar-10.

(a) bolasso		(b) bsolar-10	
frequency	variables	frequency	variables
≥ 1.00	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0$	≥ 1.00	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0$
≥ 0.88	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{28}$	≥ 0.10	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{91}, \mathbf{x}_{71}$
≥ 0.84	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{28}, \mathbf{x}_{71}$	$= 0$	all other variables
≥ 0.76	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{28}, \mathbf{x}_{71}, \mathbf{x}_{91}$		
≥ 0.70	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{28}, \mathbf{x}_{71}, \mathbf{x}_{91}, \mathbf{x}_{94}$		
≥ 0.69	$\mathbf{x}_4, \mathbf{x}_3, \mathbf{x}_2, \mathbf{x}_1, \mathbf{x}_0, \mathbf{x}_{28}, \mathbf{x}_{71}, \mathbf{x}_{91}, \mathbf{x}_{94}, \mathbf{x}_{70}, \mathbf{x}_{40}$		
\vdots	\vdots		

As shown in Table 2, bsolar produces a shorter and more accurate list of subsample variable selection frequencies. Table 2a breaks down the subsample selection frequency list from 256 subsamples for one bolasso realization with $p/n = 100/200$. Due to the length of the list, we report only subsample selection frequencies ≥ 0.69 . With only one layer of ranking, bolasso is unable to separate informative from redundant variables even with 256 subsample repetitions. The frequency discrepancy for bolasso between the highest-ranking redundant (\mathbf{x}_{28}) and the lowest-ranking informative variable (\mathbf{x}_0) is only 0.12. By contrast, Table 2b shows bsolar-10 returns a much shorter list with a frequency discrepancy between the highest-ranking redundant (\mathbf{x}_{91}) and the lowest-ranking informative variable (\mathbf{x}_0) of 0.9.

To increase the discrepancy between the lowest ranked informative and highest ranked redundant variables for bolasso, Bach (2008) suggests raising the number of subsample repetitions. However, increasing repetitions will further raise the bolasso computation load in high-dimensional spaces, increasing the advantage of bsolar.

4.6. Computation time comparison

The time complexity of an algorithm indicates only how computation time changes as data size (parameterized by n , p , and K) increases. Time complexity analysis omits many other computation parameters (such as hardware specification), suggesting it may substantially underestimate compu-

tation time difference of two algorithms in real-world problems. Hence, in this section we compare computation efficiency in terms of CPU time.

Since the computation load for lars or coordinate descent on a given sample is fixed, we may use the number of lars or coordinate descents to approximate the computation load for solar and lasso. For comparison, we compute solar with K subsamples and lasso with K -fold CV. As shown in Algorithm 1 and 3, solar computes one lars or coordinate descent on each subsample (X^k, Y^k) , which implies $K = 3$ lars or coordinate descents to compute \hat{q} and one more pass to compute c^* for variable selection. Lasso requires computing $K = 10$ lars or coordinate descents to optimize the tuning parameter and, given the optimal tuning parameter, one more pass on the full sample to select variables. Thus, the solar computation load is less than 1/3 that of lasso.

Given the computation loads for lasso and solar, we can work out the differences between bolasso and bsolar using the number of subsample repetitions (SR). Bolasso repeats lasso 256 times while bsolar-3 repeats solar only 3 times to obtain similar sparsity, bsolar-3 has approximately the same computation load as lasso.

Table 3: Simulation results for parallel computation time (mean runtime in seconds).

	$p/n \rightarrow 0$			$p/n \rightarrow 1$			$\log(p)/n \rightarrow 0$		
	$\frac{100}{100}$	$\frac{100}{150}$	$\frac{100}{200}$	$\frac{150}{100}$	$\frac{200}{150}$	$\frac{250}{200}$	$\frac{400}{200}$	$\frac{800}{400}$	$\frac{1200}{600}$
bsolar-3	0.05	0.07	0.08	0.06	0.08	0.12	0.32	0.51	1.04
bolasso (lars, 256 SR)	9.52	12.49	10.61	10.01	13.92	19.72	23.10	184.59	502.56
bolasso (cd, 256 SR)	13.49	60.51	60.35	13.92	16.85	20.17	27.73	100.58	308.12

Table 3 shows the average runtimes. In general, bsolar-3 has a much shorter runtime than bolasso. For low $n \times p$ (the first 5 columns), bsolar-3 runtime is roughly 0.5-1% of bolasso runtime (assuming bolasso is solved by lars), consistent with the time complexity estimation. As n and p increase, parallel computation of bolasso is progressively more difficult to coordinate. This is primarily because the CPU must simultaneously generate 10-16 (the number of CPU cores) data matrices $X \in \mathbb{R}^{1200 \times 600}$, each of which must then be bootstrapped into 256 sub-matrices $X_{sub} \in \mathbb{R}^{1080 \times 600}$ for each CPU core to read again in parallel. The volume of data is more than the CPU can read from RAM in a single pass. As a result, the runtime differences are even more pronounced as p and n increase. The 256 subsample repetitions (2,816 lars or coordinate descent repetitions in total) render the bolasso selection algorithms computationally infeasible even with moderate p and n . By contrast, bsolar-3 requires only 9 realizations of lars or coordinate descent. Due to the lighter computational load and CPU usage, bsolar-3 parallel computing is much easier to coordinate. It follows that the computation time difference will be more substantial if the number of CPU cores is below 8.

4.6.1. Comparison with previous lasso computation research

We demonstrate the solar computation advantages in bootstrap selection in two steps. First, we show that, for given computation resources, our bolasso implementation almost attains the theoretical maximum speedup for lasso parallelization. Second, we show that bsolar is substantially faster than our bolasso package. Thus, for given computation resources, bsolar speed substantially exceeds the theoretical maximum speed of bolasso.

Given the same convergence criteria (tolerance for optimization and number of iterations), number of CV folds ($K = 10$), and number of λ s in the grid search (100), the time complexity of lasso is mostly determined by n , p , and the pairwise correlations among the covariates ($corr$). For the purposes of comparison, we consider a Gaussian regression with $p/n = 1000/100$ and $corr = 0.5$.

- Using a 2.8GHz frequency, 2-core Intel Xeon CPU, [Friedman et al. \(2010\)](#), Table 1) reports an average runtime of 0.07 seconds per pathwise coordinate descent realization (with covariance pre-computed for updating).⁸
- Using an Intel Xeon W-3245 CPU with 3.2GHz frequency and 16 cores, the average runtime for the coordinate descent bolasso package (with covariance pre-computed automatically) is 41.92 seconds for 2,816 pathwise coordinate descent realizations (i.e., 256 realizations of 10-fold, cross-validated lasso), giving an average runtime of 0.014 seconds per pathwise coordinate descent realization.

Thus, with a similar CPU frequency and 14 additional cores, our lasso implementation produces an average speedup of $0.07/0.014 = 5.0$ times over [Friedman et al. \(2010\)](#) for each pathwise coordinate descent realization.

The bolasso code and the [Friedman et al. \(2010\)](#) code use the same design: 10 (parallelizable) pathwise coordinate descent repetitions to optimize λ followed by a final (non-parallelizable) step to compute β . Roughly 11% of the total computations (I/O, code interpretation to C++/Fortran, data generation, etc., matrix manipulation, and the step to compute β) are not parallelizable. Thus, the maximum speedup (given n and p) according to Amdahl's law is:

$$\frac{1}{\rho + (1 - \rho)/s} = \frac{1}{0.11 + (1 - 0.11)/(16/2)} \approx 4.5, \quad (4.2)$$

where ρ is the proportion of computation that is not parallelizable and s is the computation speedup for the parallelizable proportion (i.e., the core number multiple). Given that our CPU base frequency is also higher than [Friedman et al. \(2010\)](#) (3.2GHz over 2.8GHz), we adjust the maximum speedup by the frequency multiple (3.2/2.8), resulting in a final maximum speedup of $4.5 \times 3.2/2.8 \approx 5.2$, or

⁸The [Friedman et al. \(2010\)](#) package is coded in R with all numerical computations executed in Fortran/C++.

4% faster than our speedup of 5.0. Hence, given the core number and CPU frequency, our coordinate descent bolasso package achieves almost 96% of the maximum possible speedup.

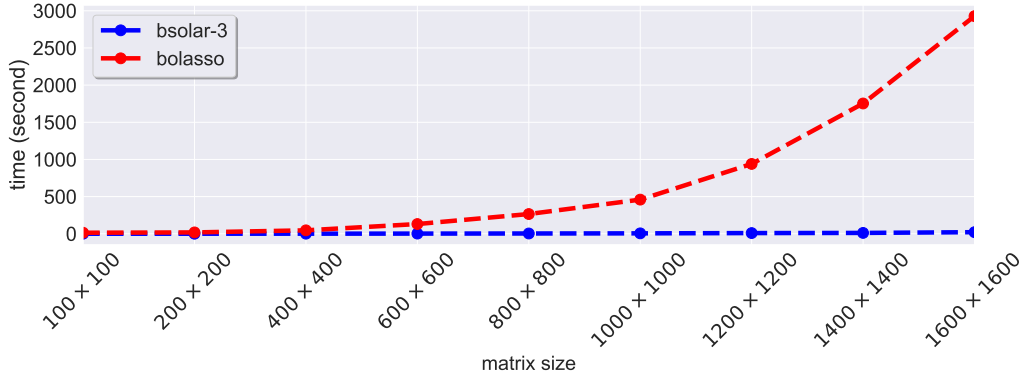


Figure 7: Average runtime (per pathwise coordinate descent) comparison for different X matrix sizes.

As shown in Figure 7, bsolar easily surpasses the theoretical maximum speedup for parallelizing lasso-type estimators as the size ($n \times p$) of the X matrix increases. As matrix size increases, the optimized bolasso package runtime rises exponentially while bsolar runtime increases linearly, confirming the bsolar-3 advantage for high-dimensional data.

4.6.2. Implication of solar computation advantages

The efficiency of bsolar computation solves the issue of choosing a bootstrap variable selection threshold. [Bach \(2008\)](#) and [Meinshausen and Bühlmann \(2010\)](#) claim that choosing pre-defined values for the selection threshold ($f = 1$, 0.9 , or $f \in [0.6, 0.9]$) will return similarly sparse results. However, [Bach \(2008\)](#) and [Huang et al. \(2014\)](#) show that predefined thresholds may cause problems for variable selection. With $p/n = 50/500$, $\text{sd}(e) = 3$, and true signal strength around 2, [Huang et al. \(2014\)](#) finds a 50% false discovery rate with bootstrap selection methods, suggesting that the threshold still requires data-driven tuning. Moreover, the large number of bootstrap repetitions makes traditional parameter tuning methods (such as cross validation) computationally infeasible for stability selection or bolasso. By contrast, the bsolar algorithm is efficient with even large n and p : it is possible to tune f by cross validation with a runtime of less than 6 seconds for $p/n = 1200/600$.

5. Real-world data: Sydney house price prediction

To demonstrate that the improvements from solar are empirically feasible, we apply solar to real-world data. The real-world data reflect both the $p/n \rightarrow 0$ scenarios as well as the challenging IRC settings, complicated dependence structures, and grouping effects typical of data in the social sciences.

The database is assembled from multiple sources. The primary source comprises real estate market transaction data for 11,974 Sydney, Australia, houses sold in 2010, including price and house attribute

information (GIS coordinates, property address, bedrooms, bathrooms, car spaces, etc.). Each property is GIS-matched with: neighbourhood information from 2011 census data by Statistical Area Level 1 (the smallest census area in Australia, comprising at most 200 people or 60 households); 2010 and 2011 crime data by suburb; 2010 geo-spatial information on topology, climate, pollution, and aircraft noise; 2009 primary and secondary school standardized test performance data; and 2010 Sydney traffic and public transport data (bus routes, train stations, and ferry wharfs). We predict house price with a linear model of characteristics based on house attribute, census, crime, geo-spatial, school, and transport data.

Using an ensemble of Bayes network learning algorithms for data cleaning, we reject variables with both very low conditional and unconditional correlations to house price. The remaining 57 variables are listed in the first column of Table 4.⁹ The remaining variables fall into 5 broad categories: house attributes, distance to local amenities (public transport, shopping, etc.), neighbourhood socioeconomic data, localized administrative and crime data, and local school quality. Pairwise correlations among all 57 covariates indicate, not surprisingly, severe multicollinearity and grouping effects, implying a harsh IRC setting.¹⁰ Thus, heuristically increasing the value of the tuning parameter in lasso-type estimators (e.g., using the one-sd or the ‘elbow’ rule) is unlikely to be useful since it may trigger further grouping effects and the random dropping of variables.

Table 4 shows the selection comparison across the elastic net, lasso, and solar. With all variables in linear form, both lasso and elastic net lose sparsity, likely due to the complicated dependence structures and severe multicollinearity in the data, accordant with Jia and Yu (2010). By contrast, solar returns a much sparser model, with only 9 variables selected from 57. Very similar results are found with the variables in log form, hinting that solar possesses superior selection sparsity and robustness to a change in functional form. More importantly, solar variable selection outperforms the lasso-type estimators in terms of the balance between sparsity and prediction power. While pruning 25-48 variables from the elastic net and lasso selections, the post-selection regression R^2 for solar falls by just 3-5%.

6. Conclusion

In this paper we propose the solar (subsample-ordered least-angle regression) algorithm for high-dimensional data. Solar constructs solution paths using the L_0 norm and averages the resulting solution paths across subsamples, reducing sensitivity to high dimensionality while improving variable selection stability, efficiency, and accuracy. We prove that L_0 path averaging separates informative from redundant variables, that solar variable selection is consistent, and that the probability solar

⁹Due to the 200GB size of the database, Supplementary Material B only includes data for the 57 variables.

¹⁰Correlations and IRC are also reported in Supplementary Material B.

omits weak signals is controllable for finite sample size.

Through simulations, examples, and real-world data, we demonstrate that, with a considerable reduction in computation load, solar yields substantial improvements over lasso in terms of the sparsity, stability, and accuracy of variable selection. We also find that solar largely avoids selection of redundant variables and rejection of informative variables in the presence of complicated dependence structures and harsh settings of the irrepresentable condition while conserving residual degrees of freedom for hypothesis testing. Relative to bootstrap lasso, bootstrapping solar improves selection sparsity and ranking accuracy and, for given computation resources, is substantially faster.

Detection of weak signals is a potential weakness in solar, although relative to lasso the difference is very slight. Nonetheless, we are working on an extension to solar, the double-bootstrap solar (DBsolar), which, if early results are any indication, promises to enable solar accurately to detect variables with weak signals.

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Table 4: Variable selection results for linear and log house price models.

Variable	Description	elastic net		lasso		solar	
		linear	log	linear	log	linear	log
Bedrooms	property, number of bedrooms	✓	✓	✓	✓	✓	✓
Baths	property, number of bathrooms	✓	✓	✓	✓	✓	✓
Parking	property, number of parking spaces	✓	✓	✓	✓	✓	✓
AreaSize	property, land size	✓	✓	✓	✓		
Airport	distance, nearest airport	✓	✓	✓	✓		
Beach	distance, nearest beach	✓	✓	✓	✓	✓	✓
Boundary	distance, nearest suburb boundary	✓	✓	✓	✓		
Cemetery	distance, nearest cemetery	✓		✓			
Child care	distance, nearest child-care centre	✓	✓	✓	✓		✓
Club	distance, nearest club	✓	✓	✓	✓		
Community facility	distance, nearest community facility	✓	✓				
Gaol	distance, nearest gaol	✓	✓			✓	✓
Golf course	distance, nearest golf course	✓	✓	✓	✓		
High	distance, nearest high school	✓	✓	✓	✓		
Hospital	distance, nearest general hospital	✓	✓		✓		
Library	distance, nearest library	✓		✓			
Medical	distance, nearest medical centre	✓	✓		✓		
Museum	distance, nearest museum	✓	✓	✓	✓		
Park	distance, nearest park	✓	✓	✓			
PO	distance, nearest post office	✓	✓		✓		
Police	distance, nearest police station	✓	✓	✓	✓		
Pre-school	distance, nearest preschool	✓	✓	✓	✓		
Primary	distance, nearest primary school	✓	✓	✓	✓		
Primary High	distance, nearest primary-high school	✓	✓	✓	✓		
Rubbish	distance, nearest rubbish incinerator	✓	✓	✓			
Sewage	distance, nearest sewage treatment	✓					
SportsCenter	distance, nearest sports centre	✓	✓	✓	✓		
SportsCourtField	distance, nearest sports court/field	✓		✓	✓		
Station	distance, nearest train station	✓		✓			
Swimming	distance, nearest swimming pool	✓	✓	✓	✓		
Tertiary	distance, nearest tertiary school	✓	✓	✓	✓		
Mortgage	SA1, mean mortgage repayment (log)	✓	✓	✓	✓	✓	✓
Rent	SA1, mean rent (log)	✓	✓	✓	✓	✓	✓
Income	SA1, mean family income (log)	✓	✓	✓	✓	✓	✓
Income (personal)	SA1, mean personal income (log)	✓					
Household size	SA1, mean household size	✓	✓	✓	✓		
Household density	SA1, mean persons to bedroom ratio	✓	✓	✓	✓		
Age	SA1, mean age	✓	✓	✓	✓		✓
English spoken	SA1, percent English at home	✓		✓			
Australian born	SA1, percent Australian-born	✓		✓			
Suburb area	suburb area	✓		✓	✓		
Population	suburb population	✓	✓		✓		
TVO2010	suburb total violent offences, 2010	✓					
TPO2010	suburb total property offences, 2010	✓	✓		✓		
TVO2009	suburb total violent offences, 2009	✓	✓	✓			
TPO2009	suburb total property offences, 2009	✓	✓				
ICSEA	local school, socio-educational advantage	✓	✓	✓	✓	✓	✓
ReadingY3	local school, year 3 mean reading score	✓	✓	✓	✓		
WritingY3	local school, year 3 mean writing score	✓	✓	✓	✓		
SpellingY3	local school, year 3 mean spelling score	✓	✓	✓			
GrammarY3	local school, year 3 mean grammar score	✓		✓			
NumeracyY3	local school, year 3 mean numeracy score	✓	✓	✓	✓		
ReadingY5	local school, year 5 mean reading score	✓					
WritingY5	local school, year 5 mean writing score	✓	✓	✓			
SpellingY5	local school, year 5 mean spelling score	✓	✓	✓			
GrammarY5	local school, year 5 mean grammar score	✓	✓	✓			
NumeracyY5	local school, year 5 mean numeracy score	✓					
	Number of variables selected	57	45	44	36	9	11
	post-selection OLS R^2	0.55	0.76	0.55	0.76	0.50	0.73
	Sample size				11,974		

Appendix A L_0 path ranking accuracy and variable selection consistency

For generality, we derive the theoretical properties of solar under the general forward selection framework of Zhang (2009, Figure 1). Our proof method is summarized as follows. Under various settings and assumptions, Tropp (2004), Yuan and Lin (2007), Wainwright (2009), Zhang (2009), and Ing and Lai (2011) have shown: (i) forward selection is consistent in different modes, and (ii) informative variables are ranked higher at earlier stages of the solution path than redundant variables. Since the L_0 path on each solar subsample is essentially the re-parameterized forward selection path, (i) and (ii) can be applied directly to the L_0 path on each solar subsample. As a result, we can build a probabilistic lower bound to show that, on average, (i) and (ii) also hold for the average L_0 path and for the variable selection result on the average L_0 path (including solar variable selection). We follow the approach of Tropp (2004), Wainwright (2009), and Zhang (2009), because their assumptions and methods of analysis are similar to the theoretical analysis of lasso-type estimators. We follow the Zhang (2009) notation.

Definition A.1. Consider the regression model $Y = X\beta + \mathbf{e}$,

1. $Y = [y_1, \dots, y_n]^T \in \mathbb{R}^{n \times 1}$ is the response variable.
2. The data matrix is $X = [\mathbf{x}_1, \dots, \mathbf{x}_p] \in \mathbb{R}^{n \times p}$ with columns $\mathbf{x}_j \in \mathbb{R}^{n \times 1}$, $\forall j = 1, \dots, p$, and rows $X_{i,\cdot} \in \mathbb{R}^{1 \times p}$, $\forall i = 1, \dots, n$. $\mathbf{e} \in \mathbb{R}^{n \times 1}$ is a stochastic noise term.
3. The regression coefficients of the data generating process (DGP) are $\bar{\beta} = [\bar{\beta}_1, \dots, \bar{\beta}_{p_1}, \mathbf{0}]^T \in \mathbb{R}^{p \times 1}$, where the first p_1 entries are not 0.
4. The support, $\forall \beta \in \mathbb{R}^{p \times 1}$, is $\text{supp}(\beta) = \{j : \beta_j \neq 0\}$.
5. Given $X \in \mathbb{R}^{n \times p}$ and $F \subset \{1, \dots, p\}$,

$$\hat{\beta}_X(F, Y) = \underset{\beta \in \mathbb{R}^{p \times 1}}{\text{argmin}} \frac{1}{n} \|X\beta - Y\|_2^2 \quad \text{subject to} \quad \text{supp}(\beta) \subset F.$$

That is, $\hat{\beta}_X(F, Y)$ is the least squares solution with coefficients restricted to F .

6. $|F|$ is the cardinality of F while $\bar{F} - F$ is the difference of sets \bar{F} and F .
7. X_F is an $n \times |F|$ matrix with columns $\mathbf{x}_j \in [\mathbf{x}_0, \dots, \mathbf{x}_p]$ with $j \in F$ arranged in ascending order.
8. To introduce the irrepresentable condition and sparse eigenvalue condition, define

$$\mu_X(F) = \max_{j \in \bar{F}} \left\| (X_F^T X_F)^{-1} X_F^T \mathbf{x}_j \right\|_1$$

and

$$\rho_X(F) = \inf \left\{ \frac{1}{n} \|X\beta\|_2^2 / \|\beta\|_2^2 : \text{supp}(\beta) \subset F \right\}$$

9. (The ϵ stopping rule.) The [Zhang \(2009, Figure 1\)](#) framework is also known as orthogonal matching pursuit (OMP). Prior to stage l , forward selection finds the unselected variable

$$\mathbf{x}^{(l)} = \operatorname{argmax}_{\mathbf{x}_j} \left| \mathbf{x}_j^T u^{(l-1)} \right|, \quad \text{for all unselected } \mathbf{x}_j,$$

where $u^{(l-1)}$ is the forward regression residual of stage $l-1$. If $|(\mathbf{x}^{(l)})^T u^{(l-1)}| > \epsilon$, the forward selection loop will select $\mathbf{x}^{(l)}$, compute $u^{(l)}$ and move to stage $l+1$; otherwise, the forward selection loop will stop and report the regression coefficients and selected variables on and before stage $l-1$.

We also adopt the [Zhang \(2009\)](#) assumptions:

- [A1] each \mathbf{x}_i is normalized such that $\|\mathbf{x}_j\|_2^2/n = 1$, $\forall j = 1, \dots, p$;
- [A2] the $\bar{\beta}$ is sparse: $\exists \bar{\beta} \in \mathbb{R}^{p \times 1}$ with $\bar{F} = \operatorname{supp}(\bar{\beta})$ such that $\mathbf{E}(Y) = X\bar{\beta} = [X_{1,\cdot}, \bar{\beta}, \dots, X_{n,\cdot}, \bar{\beta}]^T$;
- [A3] the irrepresentable condition ([Tropp, 2004](#)): $\mu_X(\bar{F}) < 1$; the sparse eigenvalue condition ([Wainwright, 2009](#)): $\rho_X(\bar{F}) > 0$.
- [A4] there exists a $\sigma > 0$ such that $Y = [Y_1, \dots, Y_n]$ are independent (but not necessarily identically distributed) sub-Gaussians with $\mathbb{E}(Y_i) = X_{i,\cdot} \bar{\beta}$ and $\mathbb{E}_{Y_i}(e^{t(Y_i - \mathbb{E}(Y_i))}) \leq e^{\sigma^2 t^2/2}$, $\forall t \in \mathbb{R}$ and $\forall i \in \{1, \dots, n\}$.

A4 implies that Y can be either bounded or unbounded. **A2** and **A4** imply that the regression noise in the DGP, $[e_1, \dots, e_n]^T$, are independent sub-Gaussians.

The proof of variable selection accuracy and variable ranking accuracy is specified in the following steps.

Step 1 : re-parameterize the ϵ stopping rule via \hat{q}^k

[Zhang \(2009, Theorem 1\)](#) shows that the probability of omitting informative variables is bounded on a finite sample.

Theorem A.1. ([Zhang, 2009](#)) *Consider the forward selection algorithm with Assumption 1 satisfied. Given any $\eta \in (0, 1)$, with probability larger than $1 - \eta$, if the ϵ stopping criterion stops forward selection at stage l , satisfying*

$$\epsilon > \frac{1}{1 - \mu_X(\bar{F})} \sigma \sqrt{2 \ln(4p/\eta)} \quad (\text{A.1})$$

and

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\epsilon}{\rho_X(\bar{F}) \cdot \sqrt{n}}, \quad (\text{A.2})$$

then when the procedure stops at stage l ,

$$\overline{F} = F^{(l-1)},$$

where $F^{(l-1)}$ is the set of variable selected at stage $l - 1$. ■

As shown in Definition A.1, Zhang (2009) executes each forward selection stage using ϵ . By contrast, we re-parameterize forward selection stages (in Algorithm 1) and the stopping rule (in Algorithm 2) using \hat{q} . Given the assumption $\epsilon > \frac{1}{1-\mu_X(\overline{F})}\sigma\sqrt{2\ln(4p/\eta)}$, we can find an equivalent stopping criterion based on \hat{q} as follows,

- Assume that, on subsample (Y^k, X^k) , $\epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ stops the forward selection at stage l^k ;
- from line 6 of Algorithm 1, variables selected before stage l^k must have \hat{q}_i^k values $> (\tilde{p} + 1 - l^k) / \tilde{p}$, where $\tilde{p} = \min\{n(K-1)/K, p\}$
- hence, on the k^{th} subsample, the stopping rule

forward selection stops at stage l^k

is equivalent to the stopping rule

$$\text{forward selection only selects the variables } \left\{ \mathbf{x}_j : \hat{q}_j^k > (\tilde{p} + 1 - l^k) / \tilde{p} \right\}.$$

Zhang (2009) also assumes that, when forward selection stops at stage l^k on subsample (Y^k, X^k) ,

$$\epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}, \exists \eta \in (0, 1).$$

The assumption plays a key role for true signal recovery. Since solar does not explicitly use ϵ , we need to re-parameterize the assumption before applying the Zhang (2009) result. Specifically, denote

$$\omega^{(l^k)} = \left| \left(\mathbf{x}^{(l^k)} \right)^T u^{(l^k)} \right|, \quad (\text{A.3})$$

where $u^{(l^k)}$ is the regression residual and $\mathbf{x}^{(l^k)}$ the variable selected at stage l^k . Solar selects variables on the L_0 and average L_0 paths based on \hat{q}^k and \hat{q} . Hence, we can analyze the selection decisions of solar by examining the absolute co-movement between $u^{(l^k)}$ and $\mathbf{x}^{(l^k)}$ on the L_0 path, which is identical to the ϵ assumption. This can be shown as follows.

- Assume the ϵ stopping rule stops forward selection at stage l^k and $\epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$.

We must have

$$\omega^{(l^k-1)} > \epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)} \quad (\text{A.4})$$

Hence, stopping forward selection based on $\epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ implies

$$\omega^{(l^k-1)} > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}.$$

- Assume $\omega^{(\cdot)} > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ is not violated until stage l^k . This implies that we have $\omega^{(l^k-1)} > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$. Besides, since $\omega^{(\cdot)} > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ for the first $l^k - 1$ stages, there exists some ϵ^* such that

$$\epsilon^* \in \left(\frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}, \min_{i < l^k} \omega^{(i)} \right) \quad (\text{A.5})$$

If we equip the ϵ stopping rule with the ϵ^* , the ϵ stopping rule will also stop forward selection at stage l^k .

Hence, ‘the $\epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ stopping rule stops forward selection at stage l^k ’ is equivalent to ‘the last variable that forward selection selects before stop has its ω larger than $\frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}$ ’. As such, we can re-parameterize the stopping rule on (Y^k, X^k) based on \hat{q}_i^k as follows

Definition A.2. (The ϵ and \hat{q}^k stopping assumptions)

- We refer to the [Zhang \(2009\)](#) assumption on ϵ stopping rule as *the ϵ stopping assumption*:

$$\text{when forward selection stops at stage } l^k \text{ on subsample } (Y^k, X^k), \quad (\text{A.6})$$

$$\exists \eta \in (0, 1) \text{ such that } \epsilon > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}.$$

- Assume forward selection only selects $\{\mathbf{x}_j : \hat{q}_j^k > \frac{\tilde{p}+1-l^k}{\tilde{p}}\}$ on subsample (Y^k, X^k) . We define

$$\omega^{(l^k)} = \left| \left(\mathbf{x}^{(l^k)} \right)^T \mathbf{u}^{(l^k)} \right|.$$

For simplicity, we denote ω as $\omega^{(\cdot)}$ of the last variable that forward selection selects.

- we refer to the following equivalent assumption for solar as *the \hat{q}^k stopping assumption*:

$$\begin{aligned} &\text{when forward selection only selects } \left\{ \mathbf{x}_j : \hat{q}_j^k > \frac{\tilde{p}+1-l^k}{\tilde{p}} \right\} \text{ on } (Y^k, X^k), \\ &\exists \eta \in (0, 1) \text{ such that } \omega > \frac{1}{1-\mu_X(F)}\sigma\sqrt{2\ln(4p/\eta)}. \end{aligned} \quad (\text{A.7})$$

Based on Definition A.2, we can re-parameterize Theorem A.1 into Lemma A.1.

Lemma A.1. *Consider the forward selection algorithm on the k th subsample (Y^k, X^k) with Assumption 1 satisfied. With probability larger than $1 - \eta$, if (A.7) is satisfied and*

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}},$$

then

$$\bar{F} = \left\{ \mathbf{x}_j : \hat{q}_i^k > c^k \right\}.$$

where $c^k = (\tilde{p} + 1 - l^k) / \tilde{p}$.

Proof. Lemma A.1 is derived by replacing the ϵ stopping assumption with the \hat{q}^k stopping assumption. Note that

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}} \implies \min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\epsilon}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}},$$

since (A.5) implies that $\exists \epsilon > \frac{1}{1-\mu_X(F)} \sigma \sqrt{2 \ln(4p/\eta)}$ such that $\omega > \epsilon$. We also replace n in (A.2) with $n(K-1)/K$ since each subsample randomly drops $1/K$ of the original sample points in Algorithm 1. \square

Step 2 : averaging the solution paths

Since we assume the nonzero $\bar{\beta}_i$ are the first p_1 components of $\bar{\beta}$, \mathbf{x} can be rewritten as

Lemma A.2. *Consider the forward selection algorithm on the k th subsample (Y^k, X^k) with Assumption 1 satisfied. With probability less than η , if (A.7) is satisfied and*

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}},$$

then

$$\begin{cases} \hat{q}_j^k \leq c^k, \forall j \leq p_1 \\ \hat{q}_j^k > c^k, \forall j > p_1. \end{cases}$$

where $c^k = (\tilde{p} + 1 - l^k) / \tilde{p}$.

Lemma A.2 directly implies that, with high probability, you can find a threshold value c^k on the L_0 path that perfectly separates the informative from the redundant variables.

To accommodate multiple subsamples in the average L_0 path, we define the \hat{q} stopping rule by slightly modifying (A.7).

Definition A.3. (The assumption for the \hat{q} stopping rule). We refer to the following rule as *the \hat{q} stopping assumption* for the average L_0 path:

$$\begin{aligned} &\text{when forward selection only selects } \left\{ \mathbf{x}_j : \hat{q}_j^k > (\tilde{p} + 1 - l^k) / \tilde{p} \right\} \text{ on } (Y^k, X^k), \\ &\exists \eta \in (0, 1/K) \text{ such that } \omega > \frac{1}{1 - \mu_X(F)} \sigma \sqrt{2 \ln \left(\frac{4p}{K\eta} \right)}. \end{aligned} \quad (\text{A.8})$$

Lemma A.3 follows from the \hat{q} stopping assumption.

Lemma A.3. Consider the forward selection algorithm on the average L_0 path with Assumption 1 satisfied. With probability less than η , if (A.8) is satisfied and

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}},$$

then

$$\begin{cases} \frac{1}{K} \sum \hat{q}_i^k = \hat{q}_i > c^*, \forall i \leq p_1 \\ \frac{1}{K} \sum \hat{q}_i^k = \hat{q}_i \leq c^*, \forall i > p_1 \end{cases}$$

where $c^* = \frac{1}{K} \sum_k c^k / K$ and $c^k = (\tilde{p} + 1 - l^k) / \tilde{p}$.

Proof. The proof is a direct result from Lemma 2. If we apply the c stopping rule, Lemma 2 implies that

$$Pr \left\{ \hat{q}_i^k \leq c^k, \forall i \leq p_1 \text{ and } \hat{q}_j^k > c^k, \forall j > p_1 \right\} \leq \eta / K. \quad (\text{A.9})$$

Since, for multiple events A_i , $Pr \{ \cap_i A_i \} \leq \sum_i Pr \{ A_i \}$, we have

$$Pr \left\{ \sum_{k=1}^K \hat{q}_i^k \leq \sum_{k=1}^K c^k, \forall i \leq p_1 \text{ and } \sum_{k=1}^K \hat{q}_j^k > \sum_{k=1}^K c^k, \forall j > p_1 \right\} \leq \eta. \quad (\text{A.10})$$

Since, $\hat{q}_i = \frac{1}{K} \sum \hat{q}_i^k$ and $c^* = \frac{1}{K} \sum c^k$, we have

$$Pr \{ \hat{q}_i \leq c^*, \forall i \leq p_1 \text{ and } \hat{q}_j > c^*, \forall j > p_1 \} \leq \eta. \quad (\text{A.11})$$

□

Lemma A.3 directly implies that, with high probability, you can find a threshold value c^* on the average L_0 path that perfectly separates the informative from the redundant variables.

Step 3 : variable selection consistency

Based on Lemma A.3, Theorem A.2 on variable selection consistency is straightforward.

Theorem A.2. *Consider the forward selection algorithm on the average L_0 path with Assumption 1 satisfied, noise σ independent of n . Assume that the strong irrepresentable condition holds. For each sample size n , denote $F(n)$ as the index set of selected variables when forward selection stops with $\omega \geq n^{s/2}$, $\forall s \in (0, 1]$, and $\bar{F}(n)$ as the corresponding index set of informative variables. We have*

$$Pr(F(n) \neq \bar{F}(n)) \leq \exp\left(-\frac{n^s}{\log(n)}\right)$$

if

$$p(n) \leq \exp\left(\frac{n^s}{\log(n)}\right),$$

and

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3n^{(s-1)/2}}{\rho_X(\bar{F}(n))}$$

where $p(n)$ is the total dimension of variable as n increases.

Proof. When n is sufficiently large, the assumptions

$$\omega^{(l^k)} = \left| \left(\mathbf{x}^{(l^k)} \right)^T u^{(l^k)} \right| > \frac{1}{1 - \mu_X(F)} \sigma \sqrt{2 \ln(4p/\eta)}$$

and

$$\min_{j \in \bar{F}} |\bar{\beta}_j| \geq \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}}$$

hold with $\eta = \exp(-n^s/\log(n))$. Thus, Theorem A.2 follows from Lemmas A.2 and A.3. \square

Step 4 : probability of omitting weak signals

Zhang (2009, Theorem 2) shows that, if

$$\min_{j \in \bar{F}} |\bar{\beta}_j| < \frac{3\omega}{\rho_X(\bar{F}) \cdot \sqrt{n(K-1)/K}},$$

the probability of selecting at least one redundant \mathbf{x}_i and the number of omitted weak signals are still bounded by sample size, the sparse eigenvalue condition, and the stopping condition. Using the same procedure as step 1, we can rewrite Zhang (2009, Theorem 2) into Lemma A.4.

Lemma A.4. *Consider the forward selection algorithm on the k th subsample (Y^k, X^k) with Assumption 1 satisfied. With probability larger than $1 - \eta$, if*

$$\omega > \frac{1}{1 - \mu_X(\bar{F})} \sigma \sqrt{2 \ln(4p/\eta)} \quad (\text{A.12})$$

then when the procedure stops at stage l , the following claims are true:

$$\left\{ \begin{aligned} & \{\mathbf{x}_j : \hat{q}_i^k > c^k\} \subset \overline{F} \\ & \left| |\{\mathbf{x}_j : \hat{q}_i^k > c^k\} - \overline{F}| \leq 2 \left| \left\{ j \in \overline{F} : \left| \bar{\beta}_j < \frac{3\omega}{\rho_X(\overline{F}) \cdot \sqrt{n(K-1)/K}} \right| \right\} \right| \right| \end{aligned} \right.$$

where $F^{(l-1)}$ is the set of variable selected at stage $l-1$. ■

Using the same method as Lemma A.3, Lemma A.5 follows from the \hat{q} stopping assumption, showing that selection errors are strictly restricted by sample size, the sparse eigenvalue condition, and the stopping condition on the average path.

Lemma A.5. *Consider the forward selection algorithm on the average L_0 path with Assumption 1 satisfied. With probability less than η , if (A.8) is satisfied, then*

$$\left\{ \begin{aligned} & \{\mathbf{x}_j : \hat{q}^k > c^*\} \subset \overline{F} \\ & \left| |\{\mathbf{x}_j : \hat{q}^k > c^*\} - \overline{F}| > 2 \left| \left\{ j \in \overline{F} : \left| \bar{\beta}_j < \frac{3\omega}{\rho_X(\overline{F}) \cdot \sqrt{n(K-1)/K}} \right| \right\} \right| \right| \end{aligned} \right.$$

where $c^* = \frac{1}{K} \sum_k c^k / K$ and $c^k = (\tilde{p} + 1 - l^k) / \tilde{p}$.