STRUCTURED, SPARSE REGRESSION WITH APPLICATION TO HIV DRUG RESISTANCE

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We introduce a new version of forward stepwise regression. Our modification finds solutions to regression problems where the selected predictors appear in a structured pattern, with respect to a predefined distance measure over the candidate predictors. Our method is motivated by the problem of predicting HIV-1 drug resistance from protein sequences. We find that our method improves the interpretability of drug resistance while producing comparable predictive accuracy to standard methods. We also demonstrate our method in a simulation study and present some theoretical results and connections.

1. Introduction. About twenty antiretroviral drugs are currently available for the treatment of human immunodeficiency virus type 1 (HIV-1). The great majority of these function by inhibiting the activity of various proteins produced by the HIV-1 virus, effectively impairing the virus’ ability to reproduce. Resistance to these drugs develops when a mutation changes the structure of the target protein enough to frustrate the drug while still maintaining the function of the protein. HIV-1 is capable of rapid mutation, and is thus often able to adapt to antiretroviral therapy. Understanding the genetic basis for this developed resistance would allow more effective development of new drugs, as well as more informed prescription of the currently available drugs.

Sequencing HIV-1 proteins can be done reliably, and well-designed in-vitro experiments are available for testing the resistance of a particular strain of HIV-1 to drugs; see Petropoulos et al. (2000) and Zhang et al. (2005). We approach this problem using regression. This problem setting leads us to build models to predict drug resistance using mutations in the amino acid sequence of the target proteins. We desire models that are easy to interpret and take into account properties of proteins and amino acids. In particular, it is well known that proteins generally function using areas called active sites, that are, simply, areas of the sequence where the protein binds or otherwise interacts with other molecules. This fact leads us to believe that important mutations will tend to be clustered around such sites.

Protein sequences can be thought to have two layers of structure: the primary sequence consisting of a single string of adjacent amino acids, and a secondary

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structure created by protein folding. We can measure the distance between amino acids in a protein sequence roughly using the differences in position in the primary sequence. When the protein’s folding structure is known, three-dimensional distance can be calculated for any two amino acid positions. But even when the structure of the protein is unknown, because of the continuity of the primary sequence, clustering in three-dimensional space generally corresponds to clustering in the protein primary sequence.

We therefore build models for predicting resistance from mutations that have the following two properties: (1) **Sparsity**—a model that uses only a few mutations is easier to interpret and apply. (2) **Structure**—following the concept of active sites, we wish to use mutations that are clustered in the protein primary sequence. Note that this second property is desirable in other applications. For example, Liu, Lin and Ghosh (2007) use genetic pathways to model the genetic influences on prostate cancer. These pathways can be modeled as a structure on individual genes. In this paper we introduce a variable selection method that builds regression models that satisfy these two properties.

Forward stepwise regression and the lasso are two popular automatic variable selection techniques that are effective at finding sparse regression models. Given data \((X_1, Y_1), \ldots, (X_n, Y_n)\) where \(Y_i \in \mathbb{R}\) and \(X_i \in \mathbb{R}^p\), the lasso \(\hat{\beta}_\text{lasso}\) estimator due to Tibshirani (1996) minimizes

\[
\sum_{i=1}^{n} (Y_i - X_i^T \beta)^2 + \lambda \| \beta \|_1,
\]

where \(\| \beta \|_1 = \sum_j |\beta_j|\) and \(\lambda > 0\) is a tuning parameter which controls the amount of regularization. Forward stepwise regression is a greedy method that adds one predictor, that is, one element \(X_i\), at a time. Both produce sparse solutions, meaning that \(\hat{\beta}_j = 0\) for most \(j\). Sparse solutions are attractive both computationally and for interpretation.

Recent results show that both methods yield estimators with good properties. See Bunea, Tsybakov and Wegkamp (2007), Greenshtein and Ritov (2004), Wainwright (2007) for results on the lasso, and Barron et al. (2008) for results on forward stepwise regression. These papers show that, under weak conditions, both approaches yield predictors that are \(O(n^{-1/4})\) close to the optimal sparse linear predictor. Moreover, this rate cannot be improved. In our application, extra information is available—we expect nonzero \(\beta_j\)’s to cluster together. In this case, we would like to add an additional constraint to the regression.

In this paper we introduce a modification of forward stepwise regression that encourages the selection of new predictors that are “close”—with respect to a distance measure over the predictors—to those already included in the model. We show that our method, Clustered and Sparse Regression (CaSpaR), is useful in regression problems where we desire both a sparse and structured solution.
2. Data. The Stanford HIV drug resistance database described in Rhee et al. (2003) is a large data set of HIV-1 protease sequences, along with resistance phenotypes for up to seven different protease inhibitor (PI) drugs for each sequence. This database is a combination of smaller data sets collected in different clinical trials. Since both the genotyping and phenotyping experiments are well standardized, such a joining of data will not give rise to significant heterogeneity-in-sample concerns. Each protease protein sequence is 99 amino acids long. The phenotypes are obtained from in-vitro experiments, and are measured in terms of number of multiples of standard dose of drug needed to suppress virus reproduction.

We can cast the problem of connecting genotype to phenotype as a regression problem by treating each mutation as a predictor. Previous studies by Rhee et al. (2006) and Beerenwinkel et al. (2003) have used most modern sparse regression and classification techniques to attack this problem. We seek a model that will take into account protein active sites.

3. CaSpaR. We first introduce the usual regression setting. We have an $n \times p$ data matrix $X$ and $n \times 1$ response vector $Y$. We use the usual linear model

\[ Y = X\beta + \varepsilon. \]

Define the support of $\beta$ by

\[ \text{supp}(\beta) = \{ j : \beta_j \neq 0, j = 1, \ldots, p \}. \]

We assume that $\beta$ is sparse (most $\beta_j$’s are 0) and also that $\text{supp}(\beta)$ has structure. We base this structure on a distance measure $d(\cdot, \cdot)$ over the set of predictors:

\[ d(\cdot, \cdot) : \{1, \ldots, p\} \times \{1, \ldots, p\} \rightarrow \mathbb{R}. \]

Specifically, we assume that the nonzero elements of $\beta$ are spatially clustered with respect to $d(\cdot, \cdot)$. In other words, the nonzero entries of $\beta$ appear in some number of groups in which the members are “close” to each other—as defined by $d(\cdot, \cdot)$. Our goal is to accurately recover $\beta$, with particular emphasis on this sparsity structure.

We want to modify a sparse regression technique to produce solutions with clusters of nonzero coefficients. Penalized techniques such as the lasso are difficult to modify for this purpose. Recall that the lasso finds $\hat{\beta}$ that minimizes

\[ Q(\beta) = \sum_{i=1}^{n} (Y_i - X_i^T \beta)^2 + \lambda \sum_j |\beta_j|. \]

The lasso is computationally efficient because $Q(\beta)$ is convex. It is difficult to add a penalty to $Q(\beta)$ that encourages clustered solutions while maintaining convexity. Note that the fused lasso due to Tibshirani et al. (2005) adds a penalty of the form $\sum_j |\beta_j - \beta_{j-1}|$. This forces nearby coefficients to be close together in sign and magnitude. We want the support points to be close together, but we do not want
1. Input: \( A = \emptyset, X, Y, \varepsilon > 0 \).
2. Fit an OLS model: \( \hat{\beta} = \arg\min_{\beta} \|X\beta - y\|_2^2 \), s.t. \( \text{supp}(\beta) \subseteq A \).
3. Set \( i^* = \arg\max_{l \notin A} |(X\beta - y)^T x_l| \).
4. If \( |x_l^T (X\beta - y)| < \varepsilon \) then stop, else set \( A = A \cup i^* \) and go to step 2.

Table 1

**Forward stepwise regression**

Table 2

**CaSpaR: Clustered and Sparse Regression**

1. Input: \( A = \emptyset, X, Y, h > 0, \alpha \in (0, 1), \varepsilon > 0 \).
2. Fit an OLS model: \( \hat{\beta} = \arg\min_{\beta} \|X\beta - y\|_2^2 \), s.t. \( \text{supp}(\beta) \subseteq A \).
3. \( \forall l \notin A \), calculate: \( W_l = \frac{1}{|A|} \sum_{k \in A} K_h(d(l, k)) \). If this is the first iteration of the algorithm, set \( W_l = 1, \forall l \).
4. Set \( l^* = \arg\max_{l \notin A} W_l |(X\beta - y)^T x_l| \).
5. If \( |x_l^T (X\beta - y)| < \varepsilon \) then stop, else set \( A = A \cup l^* \) and go to step 2.
We then select the next predictor $j^*$ using a weighted criterion: $W_j (X\beta - y)^T x_j$. For most familiar kernels, such as a Gaussian kernel or an Epanechnikov kernel, this has the effect of boosting the criterion for predictors “near” those already included in the model, and diminishing the criterion for those “far away.” For practical application, we recommend a mixture of a familiar kernel, such as a boxcar or Epanechnikov, and a uniform distribution. This mixture, which we call the Stetson kernel, introduces an additional mixing parameter $\alpha$:

$$K_{h,\alpha}(x) = \alpha + (1 - \alpha)K_h(d(x)),$$

where $K_h$ is a kernel such as a boxcar, Epanechnikov or Gaussian. An example of this kernel appears in Figure 1. We particularly recommend the Epanechnikov or the boxcar kernel, because these kernels have no impact at all on predictors outside their bandwidth, and so $W_i = \alpha$ for these predictors. While this usually makes no difference in predictor selection, it simplifies precise computation and interpretation.

The advantage of the Stetson kernel is that this mixture allows multiple groups of predictors to appear in the sparsity structure. If we were instead to only use a familiar kernel, then we would have $W_j = 0$ (or extremely small) for those $j$ far enough away from predictors already included in the model. This approach would lead to only a single group in the sparsity structure, built around the first selected predictor, whereas most applications call for multiple groups. The Stetson kernel avoids this problem. The uniform part of the Stetson kernel allows new predictor groups to enter the model. The kernel part of the mixture encourages clustering around predictors already included in the model.

Finally, note that CaSpaR is closely related to forward stepwise regression. Indeed, with $\alpha = 1$ CaSpaR reduces to forward stepwise regression. Therefore, as long we consider $\alpha = 1$ when picking parameters, we always consider the forward stepwise regression solution. Consequently, we have a loose guarantee that CaSpaR does no worse than forward stepwise regression. Moreover, we expect that some theoretical results relating to forward stepwise regression can be adapted to CaSpaR.
3.1. Tuning parameters. CaSpaR has three tuning parameters: $\epsilon$, $h$, and $\alpha$. The parameter $\epsilon$ controls the sparsity of the fitted model. The parameters $h$ and $\alpha$ control the amount of structure in the estimated support. For the Stetson kernel, as the bandwidth $h$ decreases, the predictors become more tightly grouped. As $\alpha$ increases, new clusters are allowed to form more easily. In the special case where $\alpha = 1$, the method reduces to the usual forward stepwise regression method. Let $CV(\epsilon, h, \alpha)$ denote the cross-validation score. We choose the parameters by minimizing $CV(\epsilon, h, \alpha)$. Note that since small changes in $h$ or $\alpha$ do not affect the order of predictor selection, this tuning can be accomplished using a simple grid search.

4. Results. We now return to our application to HIV drug resistance. Our data set consists of 553 amino acid sequences, all 99 amino acids in length. Each amino acid sequence corresponds to a different strain of HIV found within a patient. Each sequence has resistance measurements for up to seven HIV inhibiting drugs. Thus, the number of sequences available for our analysis varies depending on which drug we consider.

After we choose a drug and take the appropriate subset of our 553 sequences, we create our predictors. With twenty known amino acids, each position in these sequences thus takes twenty possible values. We thus define our mutation predictors as follows. At each of the 99 positions, we first search across all of the available sequences and record the set of amino acids that appear at that position in the data. This set is the collection of possible mutations at that particular position. If there is only one amino acid in this set, this corresponds to the case where that particular position displays no variation in amino acid over the data, and thus can be dropped from the analysis. We use mutations from positions with $M > 1$ possible amino acids to create $M - 1$ predictors. Each of these predictors is an indicator variable which, for a particular sequence, is equal to 1 if the particular amino acid appears at that particular position and 0 otherwise. We refer to these predictors as mutations. Since each mutation has an associated position in the primary sequence, we can define a distance between predictors as the absolute difference of their positions. Thus, the mutations that occur at the same position are distance 0 from each other.

Our design matrix $X$ is thus an $n_{\text{drug}} \times p_{\text{drug}}$ matrix. Here $n_{\text{drug}}$ is the number of sequences with measurements of the resistance score for the drug of interest. The number of sequences with resistance measurements for each drug are as follows: 453 for drug APV, 212 for ATV, 496 for IDV, 300 for LPV, 510 for NFV, 465 for RTV, and 493 for SQV. We then create the $p_{\text{drug}}$ mutation indicator predictors as described above. Since the number of samples varies with the drug, so does the number of mutation predictors. The number of predictors for each drug are as follows: 210 for drug APV, 180 for ATV, 215 for IDV, 199 for LPV, 219 for NFV, 215 for RTV, and 218 for SQV.

We compare CaSpaR to forward stepwise regression and lasso models. For all methods, we use ten-fold cross-validation to choose all the tuning parameters. We
Table 3
Summary of results across all models and drugs. For each model, we give the mean-square-error, as well as the number of mutations (predictors) selected in parentheses. We see that CaSpaR is comparable to forward stepwise regression in terms of MSE, with about the same number of predictors included in the model. The lasso does better in MSE, but includes many more mutations than either stepwise method. As we previously noted, neither forward stepwise regression nor the lasso allows for a structured sparse solution.

| Drug name | Stepwise | CaSpaR | Lasso |
|-----------|----------|--------|-------|
| APV       | 0.514 (7)| 0.477 (14) | 0.422 (51) |
| ATV       | 0.588 (6) | 0.494 (11) | 0.477 (39) |
| IDV       | 0.541 (13)| 0.580 (10) | 0.449 (77) |
| LPV       | 0.614 (5) | 0.507 (15) | 0.518 (35) |
| NFV       | 0.650 (19)| 0.637 (22) | 0.661 (40) |
| RTV       | 0.659 (8) | 0.714 (5)  | 0.570 (58) |
| SQV       | 0.426 (31)| 0.508 (21) | 0.447 (63) |

We present a summary of our results in Table 3. Compared to stepwise regression, CaSpaR has comparable mean-squared-error (MSE) and number of mutations selected. In most cases, CaSpaR selects a few more mutations and has a slightly lower MSE. The lasso generally does better in terms of MSE, but includes many more mutations. These results are complicated and cumbersome to interpret as a model of resistance. Overall, CaSpaR gives relatively sparse models, as desired.

Figure 2 compares the sparsity structure in the CaSpaR and stepwise solutions in four of the drugs. If we compare the sparsity patterns of the stepwise and CaSpaR solutions, we see that CaSpaR gives more clustered solutions, as expected. As mentioned before, CaSpaR and stepwise regression select about the same number of mutations. The clustered CaSpaR solutions, however, select mutations from fewer positions than stepwise regression. The CaSpaR models therefore give a comparable level of prediction accuracy and sparsity, while also having a better biological interpretation: these clusters may correspond to a functional area of the protein.

5. Simulation study. We next report the results of a simulation study. We show that CaSpaR recovers a structured sparsity pattern more effectively than forward stepwise regression and lasso. For CaSpaR, we use a Stetson kernel, and tune the parameters with a grid search over $\alpha = \{0, 0.1, 0.2, \ldots, 1\}$, and over $h = \{1, 2, 3, 4\}$ to find the optimal tuning parameters. For each method, we use 10-fold cross-validation to choose all tuning parameters and
FIG. 2. Comparison of stepwise and CaSpaR models across four drugs: APV, ATV, RTV and SQV. Each plot gives the coefficients for the selected mutation predictors, versus the locations of these mutations in the protein sequence. Each vertical line represents the magnitude of the coefficient for a mutation predictor. Note that some sequence locations can have multiple mutations.

stopping times. To measure the performance of each method, we use

\[
\text{Recovery Error} = \frac{\| \hat{\beta} - \beta \|_2^2}{\| \beta \|_2^2},
\]

where \( \hat{\beta} \) is the coefficient estimated by the method and \( \beta \) is the true coefficient vector. This metric appeals to us since it captures both selection and estimation performance. We also compare the true positive rate and false positive rate in order to directly measure selection performance. Here, a true positive is when a nonzero entry of \( \hat{\beta} \) is also nonzero in \( \beta \). A false positive is when a nonzero entry of \( \hat{\beta} \) is zero in \( \beta \).

We simulate 100 \( n \times p \) data matrices \( X \) with \( p = 250 \) columns. Each entry of these \( X \) is an i.i.d. draw from a standard normal distribution. We generate 100 corresponding true coefficient vectors \( \beta \) so that each has 7 groups of 5 nonzero coefficients, randomly placed. Thus, there are 35 nonzero entries in each \( \beta \). Within each nonzero group, we set one entry of \( \beta \) equal to 6, and the rest equal to 3 (see the top panel of Figure 3 for a display of a sample coefficient vector). We then randomize the signs of the nonzero entries. We add independent Gaussian noise with variance 1 to the simulated response.
FIG. 3. Recovery of coefficients for a single simulated data set ($n = 100$). The top panel displays the target coefficient vector. The next three panels show the estimated coefficients for Stepwise, CaSpaR and LASSO, having recovery errors 0.848, 0.059, 0.542, respectively.

To compare the three methods, we increase $n$ from 50 to 150 ($n = 50, 75, 100, 125, 150$) and compare the average recovery errors of the three methods; cf. Figure 4. CaSpaR gives near-optimal performance with fewer data points than the other methods. An example of the differences in performance between the three methods on a single simulated data set ($n = 100$) is given in Figure 3. CaSpaR recovers the signal well, while the other two methods do not. Figures 5 and 6 display a comparison of the true positive rates and false positive rates of the three methods. We see that CaSpaR achieves the best balance of these two properties, with near optimal performance when $n = 150$—a property not seen with stepwise regression or the lasso. We therefore conclude that CaSpaR can reconstruct sparse signals more effectively than stepwise regression or the lasso.

6. Theoretical properties. In this section we discuss the theoretical properties of CaSpaR. We begin by explaining how CaSpaR relates to other methods.

6.1. Related work. Several existing regression methods take into account structure as well as sparsity. Yuan and Lin (2006) introduced the grouped lasso, which allows only groups of predictors to be selected at once. This is desirable when the groups represent closely linked predictors—such as a set of predictors...
Recovery error ($\|\hat{\beta} - \beta\|_2^2 / \|\beta\|_2^2$) on simulated data with 1-dimensional structured sparsity. Black points: stepwise regression; green points: lasso; red points: CaSpaR. We can see that with less data CaSpaR achieves a much better recovery rate than either of the other two methods.

True positive rate (number of correctly identified nonzero entries of $\beta$ in $\hat{\beta}$/total number of nonzero entries of $\beta$) on simulated data with 1-dimensional structured sparsity. Black points: stepwise regression; green points: lasso; red points: CaSpaR. CaSpaR is competitive with the other two methods. Note that the superior true positive rate of the lasso comes at the cost of a high rate of false positives.
that code the levels of a multilevel factor predictor. Since this method modifies the lasso, it can be cast as a convex minimization problem. However, the groups have to be predefined, and the method does not allow for overlap between groups, making this method somewhat inflexible.

Huang, Zhang and Metaxas (2009) introduced an algorithm called StructOMP that modifies forward stepwise regression (also known as orthogonal matching pursuit or OMP). Here, the desired sparsity structure is encoded as a set of blocks, each of which is assigned a cost. The algorithm proceeds by greedily adding blocks one at a time to reduce the loss, scaled by the cost of the added block. StructOMP allows for very flexible sparsity structures. In particular, it can approximate a general class of sparsity structures the authors term graph sparsity, which we discuss in Section 6.2.

Recent work by Jacob, Obozinski and Vert (2009) relating to the grouped lasso extends the possible group structures to include overlapping groups. Like StructOMP, the overlapping group penalty can produce models that approximately follow graph sparsity. This approach has the advantage of being a convex minimization problem. As we discuss in the next section, for graph sparsity, this method, like StructOMP, gives only an approximation to graph sparsity because of computational considerations.

6.2. Graph sparsity. Graph sparsity is a specific type of structured sparsity introduced by Huang, Zhang and Metaxas (2009). Consider a graph $G$ whose nodes
include the set $\mathcal{I} = \{1, 2, \ldots, p\}$. Thus, each predictor is a node of $G$, but for
generality we allow other nodes to be in the graph as well. We then define the
neighborhood of a node $v$ as the set of nodes with an edge connecting it to $v$. More
generally, we could allow for $k$-neighborhoods—the set of all nodes with a path
of at most $k$ edges connecting it to $v$. We then consider a sparsity structure where
the important predictors appear within neighborhoods, or a series of connected
neighborhoods.

For example, consider a grid graph, such as in the case of a pixelated image.
Each pixel is connected to four neighbors, one to each cardinal direction. The
sparsity structure for this graph connects visually related components in the image.

CaSpaR can approximate graph sparsity if we employ an appropriate distance
function and bandwidth. Given a graph $G$, the distance function can be defined in
terms of the graph:

$$d(l, m) = \min\{\text{Length of paths from } l \text{ to } m, \text{ as defined by } G\}. \tag{10}$$

More generally, each edge can be weighted, and $d(\cdot, \cdot)$ can be the minimal
weighted path length. We then can define neighborhood size via the bandwidth $h$.
For the Stetson kernel, the mixing parameter $\alpha$ controls the number of connected
neighborhoods, where $\alpha = 0$ allows only one. In the image example, we can define
$d(\cdot, \cdot)$ as above. Then, with $h \in (1, 2)$, CaSpaR would find a sparsity structure of
connected pixels.

CaSpaR is a very flexible way to approximate graph sparsity. First, it allows
for neighborhoods to be locally defined through the bandwidth while still allowing
neighborhoods to grow arbitrarily large as the method proceeds. Second, when
used with the Stetson kernel, CaSpaR allows the user to control the degree to
which graph sparsity is adhered via the mixing parameter $\alpha$.

In comparison, the algorithms for the StructOMP of Huang, Zhang and
Metaxas (2009) and graph lasso of Jacob, Obozinski and Vert (2009) approxi-
mate graph sparsity by constructing a set of node neighborhoods, based on the
graph structure. These generate a set of blocks or groups, that are then used in the
OMP or group lasso framework, respectively. However, to control the computa-
tional cost, they limit the neighborhood size used to make these blocks or groups.
Because CaSpaR grows neighborhoods instead of seeking to add them all at once
as a group or block, this is not necessary. These algorithms can handle large groups
or blocks, but only at significant computational cost.

Further, in StructOMP, there is no clear way to control the degree to which
graph sparsity is followed in the solution. The blocks are each assigned a cost,
but this cost is relatively restrictive. In graph lasso, the group penalty is controlled
by a parameter $\lambda$, just as with the $\ell_1$ lasso penalty. However, the group penalty
controls sparsity as well as the structure, so as $\lambda$ decreases, the model becomes
less sparse as well as less structured. A separate $\ell_1$ penalty could allow the model
to be controlled for sparsity and structure separately.
6.3. Consistency. We now explain how a result in Zhang (2009) on stepwise regression can be adapted to CaSpaR. We summarize the result from the literature as follows: under assumptions about the data matrix and the response, it can be shown that with high probability, when the forward stepwise procedure stops, it stops with all correctly selected predictors—that is, all the nonzero entries of the final $\hat{\beta}$ are also nonzero in the true target $\beta$. Note that there may be additional “false negatives.” Moreover, if all of the target coefficients are above a threshold set by the noise level, then the entire sparsity pattern is captured exactly.

We closely follow the proof in Zhang (2009). This result requires more conditions than the similar result for stepwise regression. However, since we assume that we have a certain set of tuning parameters $\{\alpha, h\}$, the assumptions are not too harsh. For ease of reference, we use notation similar to Zhang (2009).

We have an $n \times p$ matrix $X$ consisting of $p$ $n$-vectors $\{x_1, x_2, \ldots, x_p\}$, and an $n$-vector $y$. We assume that there is a target $\beta \in \mathbb{R}^p$, such that

$$E y = X \beta. \quad (11)$$

This assumption means that the linear model is correct. It also roughly means there is a target coefficient vector $\bar{\beta}$ that estimates $y$ well, relative to the noise level. For both stepwise and CaSpaR methods, we define $\beta^{(k)}$ as the coefficient vector after the $k$th step. Recall the definition of the support of a vector:

$$\text{supp}(\beta) = \{j : \beta_j \neq 0\}. \quad (12)$$

We then define $F^{(k)} = \text{supp}(\beta^{(k)})$, $F = \text{supp}(\beta)$. Let

$$\hat{\beta}_{X}(F, y) = \arg\min_{\beta \in \mathbb{R}^p} \|X\beta - y\|_2^2 \text{ subject to } \text{supp}(\beta) \subseteq F. \quad (13)$$

Finally, we define two technical quantities:

$$\mu_X(F) = \max_{j \notin F} \|(X^T X)^{-1} X^T x_j\|_1 \quad (14)$$

and

$$\rho_X(F) = \inf_{\beta} \left\{ \frac{1}{n} \|X\beta\|_2^2 / \|\beta\|_2^2 : \text{supp}(\beta) \subseteq F \right\}. \quad (15)$$

For CaSpaR, we define a distance measure on our predictor index $1, 2, \ldots, p$: $d(\cdot, \cdot)$. We assume that we are using a boxcar kernel, or a Stetson kernel with a boxcar kernel: $K_{h, m}(l) = I_{d(md(k,l)<h}$. We then define the following set, which represents the candidate predictors—predictors not already included in the model—“underneath” the kernel:

$$A^{(k)} = \{m : d(l, m) < h, m \notin F^{(k)}\}. \quad (16)$$

It follows that

$$W_j = \begin{cases} \alpha + (1 - \alpha) / k & : j \in A^{(k)}, \\ \alpha & : j \notin A^{(k)}. \end{cases} \quad (17)$$
Finally, recall that we have $\varepsilon$ as the stopping criterion for CaSpaR. If at step $k$ we select $x_{i(k)}$ as the next predictor to be included in the model, then if
\begin{equation}
|x^T_{i(k)}(X\beta^{(k-1)} - y)| \leq \varepsilon,
\end{equation}
CaSpaR stops at step $k - 1$.

**Theorem 1.** Suppose that:
1. $\frac{1}{n}\|x_j\|_2^2 = 1 \forall j \in 1, 2, \ldots, p$.
2. $\exists \beta \in \mathbb{R}^p$, with $\mathcal{F} = \text{supp}(\beta)$ s.t. $y = X\beta$.
3. $\mu_X(\mathcal{F}) < 1$.
4. $\rho_X(\mathcal{F}) > 0$.
5. The elements of $y$; $\{y_i\}_{i=1, 2, \ldots, n}$ are independent sub-Gaussian random variables: $\exists \sigma > 0$ s.t. $\forall i, \forall t \in \mathbb{R}, \mathbb{E}[e^{t(y_i - E[y_i])}] \leq e^{\sigma t^2/2}$.
6. Given $\eta \in (0, 1)$, let the stopping criterion satisfy $\varepsilon > \frac{1}{1 - \mu_X(\mathcal{F})} \sigma \sqrt{2 \log(2p/\eta)}$.
7. There are $\{\alpha, h\}$ such that for each $k$, at least one of the following conditions holds:
   - (a) $\max_{j \notin \mathcal{F}} |x^T_j(X\beta^{(k-1)} - y)| < \alpha$,
   - (b) $A^{(k-1)} \subseteq \mathcal{F}$,
   - (c) $A^{(k-1)} \supseteq \mathcal{F}$.

Then, when the procedure stops at step $k - 1$, with probability greater than $1 - 2\eta$, the following hold:
1. $F^{(k-1)} \subset \mathcal{F}$,
2. $|\mathcal{F} - F^{(k-1)}| \leq 2||\{j \in \mathcal{F}: |\beta_j| < 3\varepsilon\rho_X(\mathcal{F})^{-1}/\sqrt{n}\}||$,
3. $\|\beta^{(k-1)} - \hat{\beta}_X(\mathcal{F}, y)\|_2 \leq \varepsilon\rho_X(\mathcal{F})^{-1} \sqrt{|\mathcal{F} - F^{(k-1)}|/n}$,
4. $\|\hat{\beta}_X(\mathcal{F}, y) - \beta\|_\infty \leq \sigma \sqrt{2 \log(2|\mathcal{F}/\eta|/(n\rho_X(\mathcal{F}))}$.

We omit the proof as it is very similar to the proof in Zhang (2009).

**6.3.1. Discussion of the result.** The theorem states that when the procedure stops: (1) the selected predictors have truly nonzero $\beta_i$; (2) the number of false negatives is bounded by the number of small truly nonzero $\beta_j$—relative to the noise level; (3) the estimator is close to the best possible $\beta$, which is estimated in the presence of noise using all the truly nonzero predictors; and (4) the difference between the best estimate in the presence of noise and that of the true $\beta$ is bounded.

The proof of this result is based on induction at each step of the procedure. The extra conditions are motivated by the following analysis. We denote any predictor
for which $\beta_j = 0$ as a noise predictor and any predictor for which $\beta_j \neq 0$ as a signal predictor. When we consider adding a predictor in a step of forward stepwise regression, we consider two quantities:

$$\max_{j \in F} |x_j^T (X\hat{\beta}^{(k-1)} - y)|,$$

(19)

$$\max_{i \in F} |x_i^T (X\hat{\beta}^{(k-1)} - y)|.$$

(20)

These are, respectively, proportional to the maximum correlation between the current residuals and a noise predictor and the maximum correlation between the current residuals and a signal predictor. We refer to these two predictors as the “best” signal predictor and the “best” noise predictor.

For CaSpaR, we must consider how the weights applied to these quantities affect the analysis. We therefore consider the cases where: (a) the best signal predictor and the best noise predictor are in $A_{(k)}$, (b) neither the best signal predictor nor the best noise predictor are in $A_{(k)}$, or (c) the best signal predictor is in $A_{(k)}$ but the best noise predictor is not, or (d) the best noise predictor is in $A_{(k)}$ but the best signal predictor is not. Except for scenario (d), the original result for stepwise regression holds. We therefore make additional assumptions to ensure that case (d) does not occur. Those conditions are as follows:

1. The ratio of the criterion for the best noise predictor to the best signal predictor is less than $\alpha$.
2. All of the predictors under the kernel are signal predictors.
3. All of the signal predictors are under the kernel.

The first ensures that in case (d) the correlation between the signal predictor is large enough to be selected even in this case. Because the weights $W_j$ only depend on membership in $A_{(k)}$, the second and third conditions ensure that case (d) never occurs: the second means there are only signal predictors in $A_{(k)}$, and the third means that there are no signal predictors not in $A_{(k)}$.

These assumptions are fairly mild, especially if we have a strong belief that $\text{supp}(\beta)$ is truly structured. We propose that the first condition holds for early steps of CaSpaR. We can reasonably assume that it is possible for an oracle $\alpha$ to be such that the signal is sufficiently dominant over noise. The last two conditions should hold for later steps of the algorithm: enough points within each cluster have already been discovered so that it only remains to fill in the clusters.

7. Conclusion. We introduced a new method, CaSpaR, that allows us to build sparse regression models where we have some additional information about the structure of the sparsity pattern. We presented an application as well as a simulation study that show the method performs differently than the most popular sparse regression techniques. We discussed the general concept of graph sparsity, and
showed that, under high “signal-to-noise” conditions $\|\beta\|^2/\sigma \approx 500$, our method provides a flexible way to approximate graph sparsity. Our simulation study suggests that under structured sparsity conditions, CaSpaR can recover the true target with less data than standard techniques. This motivates future work to show that this property has a theoretical basis. Other topics of interest include adding backward steps to the CaSpaR algorithm as well as an extension to a convex minimization procedure, which may have some computational advantages over the stepwise procedure.

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