L_1 Regularization Path Algorithm for Generalized Linear Models

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Abstract

In this study, we introduce a path-following algorithm for L_1 regularized generalized linear models. The L_1 regularization procedure is useful especially because it, in effect, selects variables according to the amount of penalization on the L_1 norm of the coefficients, in a manner less greedy than forward selection/backward deletion. The GLM path algorithm efficiently computes solutions along the entire regularization path using the predictor-corrector method of convex-optimization. Selecting the step length of the regularization parameter is critical in controlling the overall accuracy of the paths; we suggest intuitive and flexible strategies for choosing appropriate values. We demonstrate the implementation with several simulated and real datasets.

1 Introduction

In this paper we propose a path-following algorithm for L_1 regularized generalized linear models (GLM). GLM models a random variable Y that follows a distribution in the exponential family using a linear combination of the predictors, $\mathbf{x}'\beta$, where \mathbf{x} and β denote vectors of the predictors and the coefficients, respectively. The random and the systematic components may be linked through a non-linear function; therefore, we estimate the coefficient β by solving a set of non-linear equations that satisfy the maximum likelihood criterion.

$$\hat{\beta} = \underset{\beta}{\operatorname{argmax}} L(\mathbf{y}; \beta), \tag{1}$$

where \mathcal{L} denotes the likelihood function with respect to the given data $\{(\mathbf{x}_i, y_i) : i = 1, ..., n\}$. When the number of predictors p exceeds the number of observations n, or when insignificant predictors are present, we can impose a penalization on the L_1 norm of the coefficients

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for an automatic variable selection effect. Analogous to Lasso (Tibshirani 1996) that added a penalty term to the squared error loss criterion, we modify criterion (1) with a regularization:

$$\hat{\beta}(\lambda) = \underset{\beta}{\operatorname{argmin}} \{ -\log L(\mathbf{y}; \beta) + \lambda \|\beta\|_1 \}, \tag{2}$$

where $\lambda > 0$ is the regularization parameter. Logistic regression with L_1 penalization has been introduced in Lokhorst (1999) and explored by several researchers, for example in Shevade & Keerthi (2003).

We introduce an algorithm that implements the predictor-corrector method to determine the entire path of the coefficient estimates as λ varies, i.e., to find $\{\hat{\beta}(\lambda) : 0 < \lambda < \infty\}$. Starting from $\lambda = \lambda_{max}$, where λ_{max} is the largest λ that makes $\hat{\beta}(\lambda)$ nonzero, our algorithm computes a series of solution sets, each time estimating the coefficients with a smaller λ based on the previous estimate. Each round of optimization consists of three steps: determining the step size in λ , predicting the corresponding change in the coefficients, and correcting the error in the previous prediction.

A traditional approach to variable selection is the forward selection/backward deletion method that adds/deletes variables in a greedy manner. L_1 regularization as in (2) can be viewed as a smoother and "more democratic" version of forward stepwise selection. The GLM path algorithm is not only less greedy than forward stepwise, but also provides models throughout the entire range of complexity, whereas forward stepwise often stops augmenting the model before reaching the most complex stage possible. By generating the regularization path rather than computing solutions at several fixed values of λ , we identify the order in which the variables enter or leave the model. Thus, we are able to find a regularized fit with any given number of parameters, as with the series of models from the forward stepwise procedure.

Efron, Hastie, Johnstone & Tibshirani (2004) suggested an efficient algorithm to determine the exact piecewise linear coefficient paths for Lasso; see Osborne, Presnell & Turlach (2000) for a closely related approach. The algorithm called Lars is also used for forward stagewise and least angle regression paths with slight modifications. Another example of a path-following procedure is SVM path (Hastie, Rosset, Tibshirani & Zhu 2004). They presented a method of drawing the entire regularization path for support vector machine simultaneously.

Unlike Lars or SVM paths, the GLM paths are not piecewise linear. We must select particular values of λ at which the coefficients are computed exactly; the granularity controls the overall accuracy of the paths. When the coefficients are computed on a fine grid of values for λ , the nonlinearity of the paths is more visible. We propose a way to compute the exact coefficients at the values of λ at which the set of nonzero coefficients changes. This strategy yields more accurate path in an efficient way than alternative methods and provides the exact order of the active set changes, which is important information in many application, such as gene selection.

Rosset (2004) suggested a general path-following algorithm that can be applied to any loss and penalty function with reasonable bounds on the domains and the derivatives. This

algorithm computes the coefficient paths in two steps: changing λ and updating the coefficient estimates through a Newton iteration. Zhao & Yu (2004) proposed Boosted Lasso that approximates the L_1 regularization path with respect to any convex loss function by allowing backward steps to forward stagewise fitting; whenever a step in forward stagewise fitting deviated from that of Lasso, Boosted Lasso would correct the step with a backward move. When this strategy is used with negative log-likelihood (of a distribution in the exponential family) loss function, it will approximate the L_1 regularized GLM path. As discussed by Zhao and Yu, the step sizes along the path are distributed such that Zhao and Yu's method finds the exact solutions at uniformly spaced values of $\|\beta\|_1$, while Rosset's method computes solutions at uniformly spaced λ . Our method is more flexible and efficient than these two approaches; we estimate the largest λ that will change the current active set of variables and solve for the new set of solutions at the estimated λ . Hence, the step lengths are not uniform for any single parameter but depend on the data; at the same time, we ensure that the solutions are exact at the locations where the active set changes. We demonstrate the accuracy and the efficiency of our strategy in Section 3.2.1.

Other researchers have implemented algorithms for L_1 regularized logistic regression for diverse applications. For example, Genkin, Lewis & Madigan (2004) proposed an algorithm for L_1 regularized logistic regression (for text categorization) in a Bayesian context, in which the parameter of the prior distribution was their regularization parameter. They chose the parameter based on the norm of the feature vectors or through cross-validation, performing a separate optimization for each potential value. Our method of using the solutions for a certain λ as the starting point for the next, smaller λ offers the critical advantage of reducing the number of computations.

In the following sections, we describe and support our approach in more detail with examples and justifications. We present the details of the GLM path algorithm in Section 2. In Section 3, our methods are illustrated with simulated and real datasets, including a microarray dataset consisting of over 7000 genes. We illustrate an extension of our path-following method to the Cox proportional hazards model in Section 4. We conclude with a summary and other possible extensions of our research in Section 5. Proofs for all the lemmas and theorems are provided in Appendix A.

2 GLM Path Algorithm

In this section, we describe the details of the GLM path algorithm. We compute the exact solution coefficients at particular values λ , and connect the coefficients in a piecewise linear manner for solutions corresponding to other values of λ .

2.1 Problem setup

Let $\{(\mathbf{x}_i, y_i) : \mathbf{x}_i \in \mathcal{R}^p, y_i \in \mathcal{R}, i = 1, ..., n\}$ be n pairs of p predictors and a response. Y follows a distribution in the exponential family with mean $\mu = E(Y)$ and variance V = Var(Y). Depending on its distribution, the domain of y_i could be a subset of \mathcal{R} . GLM models

the random component Y by equating its mean μ with the systematic component η through a link function g:

$$\eta = g(\mu) = \beta_0 + \mathbf{x}'\beta. \tag{3}$$

The density function of Y is expressed as follows (McCullagh & Nelder 1989):

$$L(y;\theta,\phi) = \exp\{(y\theta - b(\theta))/a(\phi) + c(y,\phi)\}. \tag{4}$$

 $a(\cdot)$, $b(\cdot)$, and $c(\cdot)$ are functions that vary according to the distributions. Assuming that the dispersion parameter ϕ is known, we are interested in finding the maximum likelihood solution for the natural parameter θ , and thus $(\beta_0, \beta')'$, with a penalization on the size of the L_1 norm of the coefficients ($\|\beta\|_1$). Therefore, our criterion with a fixed λ is reduced to finding $\boldsymbol{\beta} = (\beta_0, \beta')'$, which minimizes the following:

$$(\beta, \lambda) = -\sum_{i=1}^{n} \{y_i \theta(\beta)_i - b(\theta(\beta)_i)\} + \lambda \|\beta\|_1.$$
 (5)

Assuming that none of the components of β is zero and differentiating $l(\beta, \lambda)$ with respect to β , we define a function \mathbf{H} :

$$H(\boldsymbol{\beta}, \lambda) = \frac{\partial l}{\partial \boldsymbol{\beta}} = -\mathbf{X}'\mathbf{W}(\mathbf{y} - \boldsymbol{\mu})\frac{\partial \eta}{\partial \mu} + \lambda Sgn\begin{pmatrix} 0\\ \beta \end{pmatrix}, \tag{6}$$

where \mathbf{X} is an n by (p+1) matrix including the column of 1's, \mathbf{W} is a diagonal matrix with n diagonal elements $V_i^{-1}(\frac{\partial \mu}{\partial \eta})_i^2$, and $(\mathbf{y}-\boldsymbol{\mu})\frac{\partial \eta}{\partial \mu}$ is a vector with n elements $(y_i-\mu_i)(\frac{\partial \eta}{\partial \mu})_i$. Although we have assumed that none of the elements of $\boldsymbol{\beta}$ is zero, the set of nonzero components of $\boldsymbol{\beta}$ changes with λ , and $H(\boldsymbol{\beta}, \lambda)$ must be redefined accordingly.

Our goal is to compute the entire solution path for the coefficients β , with λ varying from λ_{max} to 0. We achieve this by drawing the curve $H(\beta, \lambda) = 0$ in (p + 2) dimensional space $(\beta \in \mathcal{R}^{p+1})$ and $\lambda \in \mathcal{R}_+$. Rosset, Zhu & Hastie (2004) provided sufficient conditions for the existence of a unique solution $\beta(\lambda)$ that minimizes the convex function $l(\beta, \lambda)$ for each $\lambda \in \mathcal{R}_+$. We first restrict our attention to the cases where the conditions are satisfied and present the algorithm; we suggest a strategy to extend the algorithm to the cases where the conditions do not hold in Section 2.4. In the former situation, a unique continuous and differentiable function $\beta(\lambda)$, such that $H(\beta(\lambda), \lambda) = 0$ exists within each open range of λ that yields a certain active set of variables; the existence of such mappings $(\lambda \to \beta(\lambda))$ can be shown using the implicit function theorem (Munkres 1991). We find the mapping $\beta(\lambda)$ sequentially with decreasing λ .

2.2 Predictor - Corrector algorithm

The predictor-corrector algorithm is one of the fundamental strategies for implementing numerical continuation (introduced and applied in various publications, for example, in

Allgower & Georg (1990) and Garcia & Zangwill (1981)). Numerical continuation has long been used in mathematics to identify the set of solutions to nonlinear equations that are traced through a 1-dimensional parameter. Among many approaches, the predictor-corrector method explicitly finds a series of solutions by using the initial conditions (solutions at one extreme value of the parameter) and continuing to find the adjacent solutions based on the current solutions. We elaborate on how the predictor-corrector method is used to trace the curve $H(\beta, \lambda) = 0$ through λ in our problem setting.

The following lemma provides the initialization of the coefficient paths:

Lemma 2.1. When λ exceeds a certain threshold, the intercept is the only nonzero coefficient: $\hat{\beta}_0 = g(\bar{y})$ and

$$H((\hat{\beta}_0, 0, \dots, 0)', \lambda) = 0 \text{ for } \lambda > \max_{j \in \{1, \dots, p\}} |\mathbf{x}_j' \hat{\mathbf{W}}(\mathbf{y} - \bar{y}\mathbf{1}) g'(\bar{y})|.$$
 (7)

We denote this threshold of λ as λ_{max} . As λ is decreased further, other variables join the active set, beginning with the variable $j_0 = \operatorname{argmax}_j |\mathbf{x}_j'(\mathbf{y} - \bar{y}\mathbf{1})|$. Reducing λ , we alternate between a predictor and a corrector step; the steps of the k-th iteration are as follows:

- 1. Step length: determine the decrement in λ . Given λ_k , we approximate the next largest λ , at which the active set changes, namely λ_{k+1} .
- 2. Predictor step: linearly approximate the corresponding change in β with the decrease in λ ; call it $\hat{\beta}^{k+}$.
- 3. Corrector step: find the exact solution of $\boldsymbol{\beta}$ that pairs with λ_{k+1} (i.e., $\boldsymbol{\beta}(\lambda_{k+1})$), using $\hat{\boldsymbol{\beta}}^{k+1}$ as the starting value; call it $\hat{\boldsymbol{\beta}}^{k+1}$.
- 4. Active set: test to see if the current active set must be modified; if so, repeat the corrector step with the updated active set.

2.2.1 Predictor step

In the k-th predictor step, $\beta(\lambda_{k+1})$ is approximated by

$$\hat{\boldsymbol{\beta}}^{k+} = \hat{\boldsymbol{\beta}}^k + (\lambda_{k+1} - \lambda_k) \frac{\partial \boldsymbol{\beta}}{\partial \lambda}$$
 (8)

$$= \hat{\boldsymbol{\beta}}^{k} - (\lambda_{k+1} - \lambda_{k})(\mathbf{X}_{A}'\mathbf{W}_{k}\mathbf{X}_{A})^{-1}Sgn\begin{pmatrix}0\\\hat{\boldsymbol{\beta}}^{k}\end{pmatrix}.$$
(9)

 \mathbf{W}_k and \mathbf{X}_A denote the current weight matrix and the columns of \mathbf{X} for the factors in the current active set, respectively. $\boldsymbol{\beta}$ in the above equations are composed only of current nonzero coefficients. This linearization is equivalent to making a quadratic approximation of the log-likelihood and extending the current solution $\hat{\boldsymbol{\beta}}^k$ by taking a weighted *Lasso* step (as in LARS).

Define $f(\lambda) = H(\beta(\lambda), \lambda)$; in the domain that yields the current active set, $f(\lambda)$ is zero for all λ . By differentiating f with respect to λ , we obtain

$$f'(\lambda) = \frac{\partial H}{\partial \lambda} + \frac{\partial H}{\partial \beta} \frac{\partial \beta}{\partial \lambda} = 0, \tag{10}$$

from which we compute $\partial \beta/\partial \lambda$.

The following theorem shows that the predictor step approximation can be arbitrarily close to the real solution by making $\lambda_k - \lambda_{k+1}$ small.

Theorem 2.2. Denote $h_k = \lambda_k - \lambda_{k+1}$, and assume that h_k is small enough that the active sets at $\lambda = \lambda_k$ and $\lambda = \lambda_{k+1}$ are the same. Then the approximated solution $\hat{\boldsymbol{\beta}}^{k+1}$ differs from the real solution $\hat{\boldsymbol{\beta}}^{k+1}$ by $O(h_k^2)$.

2.2.2 Corrector step

In the following corrector step, we use $\hat{\boldsymbol{\beta}}^{k+}$ as the initial value to find the $\boldsymbol{\beta}$ that minimizes $l(\boldsymbol{\beta}, \lambda_{k+1})$, as defined in (5) (i.e., that solves $H(\boldsymbol{\beta}, \lambda_{k+1}) = 0$ for $\boldsymbol{\beta}$). Any (convex) optimization method that applies to the minimization of a differentiable objective function with linear constraints may be implemented. The previous predictor step has provided a warm start; because $\hat{\boldsymbol{\beta}}^{k+}$ is usually close to the exact solution $\hat{\boldsymbol{\beta}}^{k+1}$, the cost of solving for the exact solution is low. The corrector steps not only find the exact solutions at a given λ but also yield the directions of $\boldsymbol{\beta}$ for the subsequent predictor steps.

We connect $\hat{\boldsymbol{\beta}}^{k+1}$ with $\hat{\boldsymbol{\beta}}^k$, forming the k-th linear segment of the path. We justify this approach by showing that if $\lambda_k - \lambda_{k+1}$ is small, then any point along the linear segment is close to the true path in some sense.

Theorem 2.3. If the solutions at λ_k and $\lambda_{k+1} = \lambda_k - h_k$, namely $\hat{\boldsymbol{\beta}}^k$ and $\hat{\boldsymbol{\beta}}^{k+1}$, are connected such that our estimate at $\lambda = \lambda_k - \alpha h_k$ for some $\alpha \in [0, 1]$ is

$$\hat{\boldsymbol{\beta}}(\lambda - \alpha h_k) = \hat{\boldsymbol{\beta}}^k + \alpha (\hat{\boldsymbol{\beta}}^{k+1} - \hat{\boldsymbol{\beta}}^k), \tag{11}$$

then $\hat{\beta}(\lambda - \alpha h_k)$ differs from the real solution $\beta(\lambda - \alpha h_k)$ by $O(h_k^2)$.

2.2.3 Active set

The active set \mathcal{A} begins from the intercept as in Lemma 2.1; after each corrector step, we check to see if \mathcal{A} should have been augmented. The following procedure for checking is justified and used by Rosset & Zhu (2003) and Rosset (2004):

$$\left|\mathbf{x}_{j}^{\prime}\mathbf{W}(\mathbf{y}-\boldsymbol{\mu})\frac{\partial\eta}{\partial\mu}\right| > \lambda \text{ for any } j \in \mathcal{A}^{c} \Longrightarrow \mathcal{A} \leftarrow \mathcal{A} \cup \{j\}.$$
 (12)

We repeat the corrector step with the modified active set until the active set is not augmented further. We then remove the variables with zero coefficients from the active set. That is,

$$|\hat{\beta}_j| = 0 \text{ for any } j \in \mathcal{A} \Longrightarrow \mathcal{A} \leftarrow \mathcal{A} \setminus \{j\}.$$
 (13)

2.2.4 Step length

Two natural choices for the step length $\delta_k = \lambda_k - \lambda_{k+1}$ are:

- $\Delta_k = \Delta$, fixed for every k, or
- a fixed change L in L_1 arc-length, achieved by setting $\Delta_k = L/\|\partial \beta/\partial \lambda\|_1$.

As we decrease the step size, the exact solutions are computed on a finer grid of λ values, and the coefficient path becomes more accurate.

We propose a more efficient and useful strategy:

• select the smallest Δ_k that will change the active set of variables.

We give an intuitive explanation of how we achieve this, by drawing on analogies with the Lars algorithm (Efron et al. 2004). At the end of the k-th iteration, the corrector step can be characterized as finding a weighted Lasso solution that satisfies $-\mathbf{X}_A'\mathbf{W}_k(\mathbf{y}-\boldsymbol{\mu})\frac{\partial \eta}{\partial \mu}+\lambda_k Sgn\binom{0}{\beta}=0$. This weighted Lasso also produces the direction for the next predictor step. If the weights \mathbf{W}_k were fixed, the weighted Lars algorithm would be able to compute the exact step length to the next active-set change point. We use this step length, even though in practice the weights change as the path progresses.

Lemma 2.4. Let $\hat{\boldsymbol{\mu}}$ be the estimates of \mathbf{y} from a corrector step, and denote the corresponding weighted correlations as

$$\hat{\mathbf{c}} = \mathbf{X}' \hat{\mathbf{W}} (\mathbf{y} - \hat{\boldsymbol{\mu}}) \frac{\partial \eta}{\partial u}. \tag{14}$$

The absolute correlations of the factors in \mathcal{A} (except for the intercept) are λ , while the values are smaller than λ for the factors in \mathcal{A}^c .

The next predictor step extends $\hat{\beta}$ as in (9), and, thus, the current correlations change. Denoting the vector of changes in correlation for a unit decrease in λ as \mathbf{a} ,

$$\mathbf{c}(h) = \hat{\mathbf{c}} - h\mathbf{a} \tag{15}$$

$$= \hat{\mathbf{c}} - h\mathbf{X}'\hat{\mathbf{W}}\mathbf{X}_A(\mathbf{X}_A'\hat{\mathbf{W}}\mathbf{X}_A)^{-1}Sgn\begin{pmatrix}0\\\hat{\beta}\end{pmatrix}, \tag{16}$$

where h > 0 is a given decrease in λ . For the factors in \mathcal{A} , the values of \mathbf{a} are those of $Sgn\begin{pmatrix} 0 \\ \hat{\beta} \end{pmatrix}$. To find the h with which any factor in \mathcal{A}^c yields the same absolute correlation as the ones in \mathcal{A} , we solve the following equations:

$$|c_j(h)| = |\hat{c}_j - ha_j| = \lambda - h \quad \text{for any } j \in \mathcal{A}^c.$$
 (17)

The equations suggest an estimate of the step length in λ as

$$h = \min_{j \in \mathcal{A}^c} \left\{ \frac{\lambda - \hat{c}_j}{1 - a_j}, \frac{\lambda + \hat{c}_j}{1 + a_j} \right\}. \tag{18}$$

In addition, to check if any variable in the active set reaches 0 before λ decreases by h, we solve the equations

$$\beta_j(\tilde{h}) = \hat{\beta}_j + \tilde{h}(\mathbf{X}_A'\hat{\mathbf{W}}\mathbf{X}_A)^{-1}Sgn\begin{pmatrix}0\\\hat{\beta}\end{pmatrix} = 0 \quad \text{for any } j \in \mathcal{A}.$$
 (19)

If $0 < \tilde{h} < h$ for any $j \in \mathcal{A}$, we expect that the corresponding variable will be eliminated from the active set before any other variable joins it; therefore, \tilde{h} rather than h is used as the next step length.

As a by-product of this step length approximation strategy, $\partial \beta/\partial \lambda$ in (8) is computed. When fitting with high-dimensional data, the active set changes with a small decrease in λ , and thus, the role of predictor step as in (8)-(9) is not critical. However, we would still include predictor steps for the unusual cases of a large decrement in λ , and with the predictor step direction automatically computed, the remaining computations are trivial.

Letting the coefficient paths be piecewise linear with the knots placed where the active set changes is a reasonable simplification of the truth based on our experience (using both simulated and real datasets). If the smallest step length that modifies the active set were to be larger than the value we have estimated, the active set remains the same, even after the corrector step. If the true step length were smaller than expected, and, thus, we missed the entering point of a new active variable by far, we would repeat a corrector step with an increased λ . (We estimate the increase in a manner analogous to (19).) Therefore, our path algorithm almost precisely detects the values of λ at which the active set changes, in the sense that we compute the exact coefficients at least once before their absolute values grow larger than δ (a small fixed quantity). δ can be set to be any small constant; one can evaluate how small it is using the standard error estimates for the coefficients from the bootstrap analysis, which is illustrated later in Section 3.2.2.

We can easily show that in the case of Gaussian distribution with the identity link, the piecewise linear paths are exact. Because $\hat{\boldsymbol{\mu}} = \mathbf{X}\hat{\boldsymbol{\beta}}$, and $V_i = Var(y_i)$ is constant for $i = 1, \ldots, n, H(\boldsymbol{\beta}, \lambda)$ simplifies to $-\mathbf{X}'(\mathbf{y}-\boldsymbol{\mu}) + \lambda Sgn\binom{0}{\beta}$. The step lengths are computed with no error; in addition, since the predictor steps yield the exact coefficient values, corrector steps are not necessary. In fact, the paths are identical to those of Lasso.

2.3 Degrees of freedom

We use the size of the active set as a measure of the degrees of freedom, which changes, not necessarily monotonically, along the solution paths. That is,

$$df(\lambda) = |\mathcal{A}(\lambda)|, \tag{20}$$

where $|\mathcal{A}(\lambda)|$ denotes the size of the active set corresponding to λ . This is based on $df(\lambda) = E|\mathcal{A}(\lambda)|$, which holds in the case of Lasso. This remarkable formula was discovered by Efron et al. (2004) and improved by Zou & Hastie (2004); the effect of shrinking cancels the price paid in variance for the aggressive searching for the best variable to include in the model.

Here we present a heuristic justification for using (20) for GLM in general, based on the results developed in Zou & Hastie (2004).

One can show that the estimates of $\boldsymbol{\beta}$ at the end of a corrector step solve a weighted Lasso problem,

$$\min_{\boldsymbol{\beta}} (\mathbf{z} - \mathbf{X}\boldsymbol{\beta})^T \mathbf{W} (\mathbf{z} - \mathbf{X}\boldsymbol{\beta}) + \lambda ||\boldsymbol{\beta}||_1, \tag{21}$$

where the working response vector is defined as

$$\mathbf{z} = \boldsymbol{\eta} + (\mathbf{y} - \boldsymbol{\mu}) \frac{\partial \boldsymbol{\eta}}{\partial \boldsymbol{\mu}}.$$
 (22)

The solution to (21) would be an appropriate fit for a linear model

$$\mathbf{z} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\epsilon} \text{ where } \boldsymbol{\epsilon} \sim (\mathbf{0}, \mathbf{W}^{-1}).$$
 (23)

This covariance is correct at the true values of η and μ , and can be defended asymptotically if appropriate assumptions are made. In fact, when $\lambda = 0$, and assuming ϵ has a Gaussian distribution, (23) leads directly to the standard asymptotic formulas and Gaussianity for the maximum-likelihood estimates in the exponential family.¹

Under these heuristics, we apply the *Stein's Lemma* (Stein 1981) to the transformed response ($\mathbf{W}^{1/2}\mathbf{z}$) so that its errors are homoskedastic. We refer readers to Zou & Hastie (2004) for the details of the application of the lemma. Simulations show that (20) approximates the degrees of freedom reasonably closely, although we omit the details here.

2.4 Adding a quadratic penalty

When some columns of X are strongly correlated, the coefficient estimates are highly unstable; the solutions might not be unique if some column are linearly dependent or redundant in the sense that they do not satisfy the conditions for Theorem 5 of Rosset et al. (2004). To overcome these situations, we propose adding a quadratic penalty term to the criterion, following the *elasticnet* proposal of Zou & Hastie (2005). That is, we compute the solution paths that satisfy the following:

$$\hat{\boldsymbol{\beta}}(\lambda_1) = \underset{\boldsymbol{\beta}}{\operatorname{argmin}} \{ -\log L(\mathbf{y}; \boldsymbol{\beta}) + \lambda_1 \|\boldsymbol{\beta}\|_1 + \frac{\lambda_2}{2} \|\boldsymbol{\beta}\|_2^2 \},$$
 (24)

where $\lambda_1 \in (0, \infty)$, and λ_2 is a fixed, small, positive constant. As a result, strong correlations among the features do not affect the stability of the fit. When the correlations are not strong, the effect of the quadratic penalty with a small λ_2 is negligible.

Assuming that all the elements of $\boldsymbol{\beta}$ are nonzero, if \mathbf{X} does not have a full column rank, $\partial H(\boldsymbol{\beta}, \lambda)/\partial \boldsymbol{\beta} = \mathbf{X}'\mathbf{W}\mathbf{X}$ is singular, where H is defined as in (6). By adding a quadratic

This assumption is clearly not true for Bernoulli responses; however, if \mathbf{y} represents grouped binomial proportions, then under the correct asymptotic assumptions $\boldsymbol{\epsilon}$ is Gaussian.

penalty term, as in (24), we redefine H:

$$\tilde{H}(\boldsymbol{\beta}, \lambda_1, \lambda_2) = -\mathbf{X}'\mathbf{W}(\mathbf{y} - \boldsymbol{\mu})\frac{\partial \eta}{\partial \mu} + \lambda_1 Sgn\begin{pmatrix} 0 \\ \beta \end{pmatrix} + \lambda_2 \begin{pmatrix} 0 \\ \beta \end{pmatrix}.$$
 (25)

Accordingly, the following $\partial \tilde{H}/\partial \boldsymbol{\beta}$ is non-singular, in general, with any $\lambda_2 > 0$:

$$\frac{\partial \tilde{H}}{\partial \boldsymbol{\beta}} = \mathbf{X}' \mathbf{W} \mathbf{X} + \lambda_2 \begin{pmatrix} 0 & \mathbf{0}' \\ \mathbf{0} & I \end{pmatrix}. \tag{26}$$

Therefore, when λ_2 is fixed at a constant, and λ_1 varies in an open set, such that the current active set remains the same, a unique, continuous, and differentiable function $\boldsymbol{\beta}(\lambda_1)$ satisfies $\tilde{H}(\boldsymbol{\beta}(\lambda_1), \lambda_1, \lambda_2) = 0$. This connection between the non-singularity and existence of a unique, continuous and differentiable coefficient path is based on the implicit function theorem (Munkres 1991).

Zou & Hastie (2005) proposed elastic regression, which added an L_2 norm penalty term to the criterion for Lasso. Zou and Hastie adjusted the values of both λ_1 and λ_2 so that variable selection and grouping effects were achieved simultaneously. For our purpose of handling inputs with strong correlations, we fixed λ_2 at a very small number, while changing the value of λ_1 for different amounts of regularization.

In the case of logistic regression, adding an L_2 penalty term is also helpful as it elegantly handles the separable data. Without the L_2 penalization, and if the data are separable by the predictors, $\|\hat{\beta}\|_1$ grows to infinity as λ_1 approaches zero. Rosset et al. (2004) showed that the normalized coefficients $\hat{\beta}/\|\hat{\beta}\|_1$ converge to the L_1 margin-maximizing separating hyperplane as λ_1 decreases to zero. In such cases, the fitted probabilities approach 0/1, and thus, the maximum likelihood solutions are undefined. However, by restricting $\|\beta\|_2$ with any small amount of quadratic penalization, we let the coefficients converge to the L_2 penalized logistic regression solutions instead of infinity as λ_1 approaches zero. As an alternative solution to the separation in logistic regression, one can apply the Jeffreys prior to the likelihood function as suggested in Firth (1993) and further demonstrated in Heinze & Schemper (2002).

3 Data Analysis

In this section, we demonstrate our algorithm through a simulation and two real datasets: South African heart disease data and leukemia cancer gene expression data. Our examples focus on binary data, hence the logistic regression GLM.

3.1 Simulated data example

We simulated a dataset of 100 observations with 5 variables and a binary response. Figure 1 shows three sets of coefficient paths with respect to λ , with different selection of step sizes. In the first plot, the exact solutions were computed at the values of λ where the active set

changed, and the solutions were connected in a piecewise linear manner. The second plot shows the paths with exact solutions on a much finer grid of λ values; we controlled the arc length to be less than 0.1 between any two adjacent values of λ . We observe the true curvature of the paths. The first plot is a reasonable approximation of the second, especially because the active set is correctly specified at any value of λ . The third plot shows the solution paths we generated using the Boosted Lasso algorithm by Zhao & Yu (2004). To compare with the previous plot, we used the negative (binomial) log-likelihood loss and the step size constant $\epsilon = 0.1$; 60 and 58 steps were taken by the GLM path algorithm (the second panel) and the Boosted Lasso algorithm (the third panel), respectively. The Boosted Lasso solution paths are less smooth as the solutions oscillate around the real path as λ decreases.

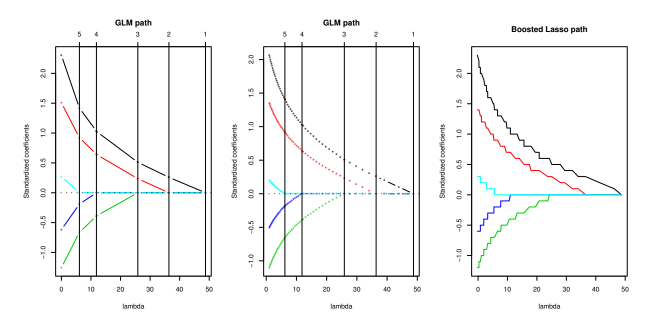


Figure 1: Simulated data from Section 3.1: In the first plot, the exact solutions were computed at the values of λ where the active set changed, and the solutions were connected in a piecewise linear manner. The second plot shows the paths with exact solutions at much finer grids of λ ; we controlled the arc length to be less than 0.1 between any two adjacent values of λ . The third plot shows the solution paths we generated using the Boosted Lasso algorithm, with the step size constant $\epsilon = 0.1$.

3.2 South African heart disease data

This dataset consists of 9 different features of 462 samples as well as the responses indicating the presence of heart disease. The dataset has also been used in Hastie, Tibshirani & Friedman (2001) with a detailed description of the data. Using the disease/non-disease response variable, we can fit a logistic regression path.

3.2.1 Selecting the step length

The first plot of Figure 2 shows the exact set of paths; the coefficients were precisely computed at 300 different values of λ ranging from 81.9 to 0, with the constraint that every arc length be less than 0.01. The L_1 norm of the coefficients forms the x-axis, and the vertical breaks indicate where the active set is modified. Comparing this plot to the second panel, which we achieved in 13 steps rather than 300, we find that the two are almost identical. Our strategy to find the λ values at which the active set changes resulted in an estimate of the values with reasonable accuracy. In addition, the exact paths are curvy but are almost indistinguishable from the piecewise linear version, justifying our simplification scheme. For both plots, the right-most solutions corresponding to $\lambda = 0$ are the maximum likelihood estimates.

The bottom panel of Figure 2 illustrates the paths with respect to the steps. Two extra steps were needed between the knots at which the sixth and the seventh variable joined the active set. However, the step lengths in λ are tiny in this region; since the first approximation of λ that would change the active set was larger than the true value by only a small amount, λ decreased again by extremely small amounts. For most other steps, the subsequent $\lambda's$ that would modify the active set were accurately estimated on their first attempts.

We have proposed three different strategies for selecting the step sizes in λ in Section 2.2.4:

- 1. Fixing the step size Δ : $\Delta_k = \Delta$
- 2. Fixing the arc length L: $\Delta_k = L/\|\partial \beta/\partial \lambda\|_1$
- 3. Estimating where the active set changes

To verify that Method 3 yields more accurate paths with a smaller number of steps and, thus, a smaller number of computations, we present the following comparison. For the three methods, we counted the number of steps taken and computed the corresponding sum of squared errors in β , $\sum_{m=1}^{200} \|\hat{\beta}_{(m)} - \beta_{(m)}\|^2$. $\hat{\beta}_{(m)}$ and $\beta_{(m)}$ denote the coefficient estimates at the m-th (out of 200 evenly spaced grid values in $\|\beta\|_1$) grid along the path, from the path generated using a certain step length computation method and the exact path, respectively.

Method 1			Method 2			Method 3	
Δ	num. steps	error	L	num. steps	error	num. steps	error
8	12	2.56e-1	0.23	11	1.01e-1	13	7.11e-4
1	83	2.04e-1	0.1	26	7.78e-2		
0.3	274	6.75e-3	0.02	142	2.28e-2		
0.15	547	7.16e-5	0.01	300	4.25e-5		

Table 1: As shown in the first row, the first two strategies of selecting the step lengths, with a comparable number of steps, achieved much lower accuracy than the third. The first two methods needed a few hundred steps to yield the same accuracy that the third method achieved in only 13 steps.

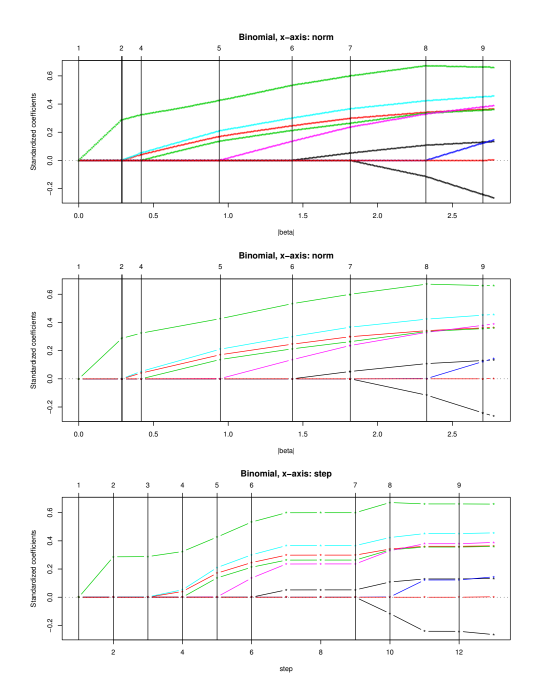


Figure 2: Heart disease data from Section 3.2: The first plot shows the exact set of paths; the coefficients were precisely computed at 300 different grids of λ ranging from 81.9 to 0, with the constraint that every arc length be less than 0.01. The vertical breaks indicate where the active set is modified, and the L_1 norm of the coefficients forms the x-axis. Comparing this plot to the second panel, which we achieved in 13 steps rather than 300, we find that the two are almost identical. The bottom panel represents the paths as a function of step-number, to illustrate how minor the corrections are.

As shown in the first row of Table 1, the first two strategies of selecting the step lengths, with a comparable number of steps, achieved much lower accuracy than the third. Furthermore, the first two methods needed a few hundred steps to yield the same accuracy that the third method achieved in only 13 steps. Thus, Method 3 provides accuracy and efficiency in addition to the information about where the junction points are located.

3.2.2 Bootstrap analysis of the coefficients

Given the series of solution sets with a varying size of the active set, we select an appropriate value of λ and, thus, a set of coefficients. We may then validate the chosen coefficient estimates through a bootstrap analysis (Efron & Tibshirani 1993).

We illustrate the validation procedure, choosing the λ that yields the smallest BIC (Bayesian information criteria). For each of the B=1000 bootstrap samples, we fit a logistic regression path and selected λ that minimized the BIC score, thereby generating a bootstrap distribution of each coefficient estimate. Table 2 summarizes the coefficient estimates computed from the whole data, the mean and the standard error of the estimates computed from the B bootstrap samples, and the percentage of the bootstrap coefficients at zero. For the variables with zero coefficients (adiposity, obesity, and alcohol), over 60% of the bootstrap estimates were zero.

Feature	\hat{eta}	$\operatorname{Mean}(\hat{\beta}^b)$	$SE(\hat{\beta}^b)$	Num. zero/B
sbp	0.0521	0.0857	0.1048	0.397
tobacco	0.2988	0.3018	0.1269	0.015
ldl	0.2636	0.2925	0.1381	0.040
adiposity	0	0.0367	0.1192	0.700
famhist	0.3633	0.3755	0.1218	0.006
typea	0.2363	0.2672	0.1420	0.054
obesity	0	-0.0875	0.1510	0.633
alcohol	0	0.0078	0.0676	0.646
age	0.5997	0.6109	0.1478	0.000

Table 2: Heart disease data from Section 3.2: The coefficient estimates computed from the whole data, the mean and the standard error of the estimates computed from the B bootstrap samples, and the percentage of the bootstrap coefficients at zero. For the variables with zero coefficients, over 60% of the bootstrap estimates were zero.

Figure 3 shows the bootstrap distributions of the standardized coefficients. Under the assumption that the original data were randomly sampled from the population, the histograms display the distributions of the coefficient estimates chosen by BIC criterion. As marked by the red vertical bars, coefficient estimates from the whole data that are nonzero fall near the center of the bootstrap distributions. For the predictors whose coefficients are zero, the histograms peak at zero. The thick vertical bars show the frequencies of zero coefficients.

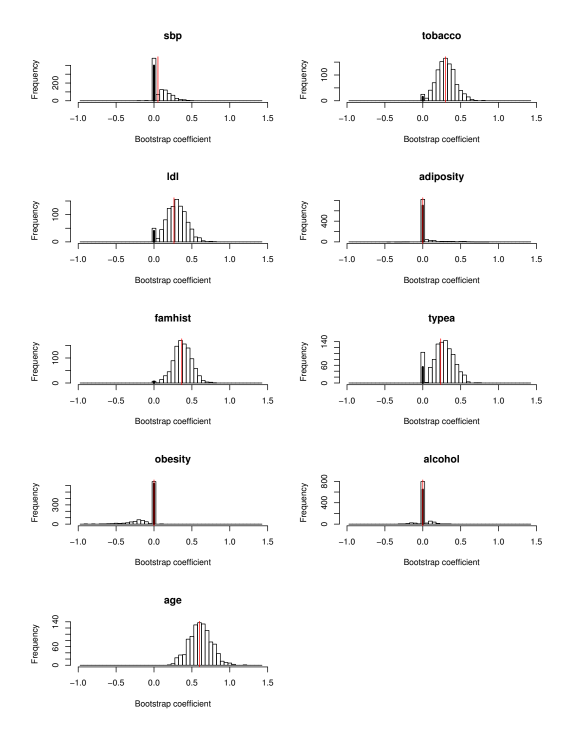


Figure 3: The bootstrap distributions of the standardized coefficients: Under the assumption that the original data were randomly sampled from the population, the histograms display the distributions of the coefficient estimates chosen by BIC criterion. As marked by the red vertical bars, coefficient estimates from the whole data that are nonzero fall near the mean of the bootstrap distributions. For the predictors whose coefficients are zero, the histograms peak at zero. The thick vertical bars show the frequencies of zero coefficients.

3.3 Leukemia cancer gene expression data

The GLM path algorithm is suitable for data consisting of far more variables than the samples (so-called $p \gg n$ scenarios) because it successfully selects up to n variables along the regularization path regardless of the number of input variables. We demonstrate this use of our algorithm through a logistic regression applied to the leukemia cancer gene expression dataset by Golub, Slonim, Tamayo, Huard, Gaasenbeek, Mesirov, Coller, Loh, Downing, Caligiuri, Bloomfield & Lander (1999). The dataset contains the training and the test samples of sizes 38 and 34, respectively. For each sample, 7129 gene expression measurements and a label indicating the cancer type (AML: acute myeloid leukemia or ALL: acute lymphoblastic leukemia) are available.

The first panel of Figure 4 shows the coefficient paths we achieved using the training data; the size of the active set cannot exceed the sample size at any segment of the paths (This fact is proved in Rosset et al. (2004)). The vertical line marks the chosen level of regularization (based on cross-validation), where 23 variables had nonzero coefficients. The second panel of Figure 4 illustrates the patterns of ten-fold cross-validation and test errors. As indicated by the vertical line, we selected λ where the cross-validation error achieved the minimum.

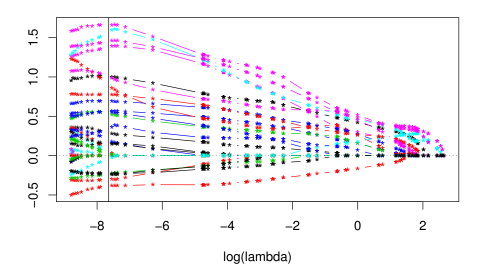
Table 1 shows the errors and the number of variables used in the prediction. We also compared the performance to other methods that used the same dataset in their literature. With a cross-validation error of 1/38 and a test error of 2/34, L_1 penalized logistic regression is comparable to or more accurate than other competing methods for analysis of this microarray dataset.

Method	CV error	Test error	No. of genes used
L_1 PLR	1/38	2/34	23
L_2 PLR (UR): (Zhu & Hastie 2004)	2/38	3/34	16
L_2 PLR (RFE): (Zhu & Hastie 2004)	2/38	1/34	26
SVM (UR): (Zhu & Hastie 2004)	2/38	3/34	22
SVM (RFE): (Zhu & Hastie 2004)	2/38	1/34	31
NSC classification: (Tibshirani et al. 2002)	1/38	2/34	21

Table 3: With a cross-validation error of 1/38 and a test error of 2/34, L_1 penalized logistic regression is comparable to or more accurate than other competing methods for analysis of this microarray dataset. Although we did not perform any pre-processing to filter from the original 7129 genes, automatic gene selection reduced the number of effective genes to 23. UR refers to univariate ranking; RFE refers to recursive feature elimination.

Although we did not perform any pre-processing to filter from the original 7129 genes, automatic gene selection reduced the number of effective genes to 23. This set of genes included 7, 14, 8, 15, and 5 of the genes selected through L_2 PLR (UR), L_2 PLR (RFE), SVM (UR), SVM (RFE), and NSC classification, respectively. Each of these methods selected only a small proportion of the available genes, which were highly correlated within sub-groups.

Coefficient path



Cross-validation & Test errors

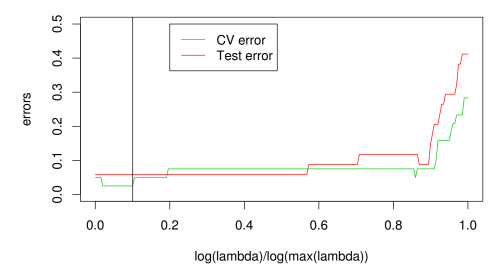


Figure 4: Leukemia data from Section 3.3: The first panel shows the coefficient paths we achieved using the training data; the size of the active set cannot exceed the sample size at any segment of the paths. The vertical line marks the chosen level of regularization (based on cross-validation), where 23 variables had nonzero coefficients. The second panel illustrates the patterns of ten-fold cross-validation and test errors. As indicated by the vertical line, we selected λ where the cross-validation error achieved the minimum.

As a result, the gene groups from different methods did not entirely overlap, but some of the genes were commonly significant across different models.

4 L_1 Regularized Cox Proportional Hazards Models

The path-following method that we applied to the L_1 regularized GLM may also be used to generate other nonlinear regularization paths. We illustrate an analogous implementation of the predictor-corrector method for drawing the L_1 regularization path for the Cox proportional hazards model (Cox 1972). Tibshirani (1997) proposed fitting the Cox model with a penalty on the size of the L_1 norm of the coefficients. This shrinkage method computes the coefficients with a criterion similar to (2):

$$\hat{\beta}(\lambda) = \underset{\beta}{\operatorname{argmin}} \{ -\log L(\mathbf{y}; \beta) + \lambda \|\beta\|_1 \}, \tag{27}$$

where L denotes the partial likelihood. We formulate the entire coefficient paths $\{\hat{\beta}(\lambda): 0 < \lambda < \lambda_{max}\}$, where λ_{max} is the largest λ that makes $\hat{\beta}(\lambda)$ nonzero, through the predictor-corrector scheme. As a result of L_1 penalization, the solutions are sparse; thus, the active set changes along with λ .

4.1 Method

Let $\{(\mathbf{x}_i, y_i, \delta_i) : \mathbf{x}_i \in \mathcal{R}^p, y_i \in \mathcal{R}_+, \delta_i \in \{0, 1\}, i = 1, ..., n\}$ be n triples of p factors, a response indicating the survival time, and a binary variable, $\delta_i = 1$ for complete (died) observations, while $\delta_i = 0$ for right-censored patients. Based on the criterion (27), we find the coefficients that minimize the following objective function for each λ :

$$l(\beta, \lambda) = -\sum_{i=1}^{n} \delta_{i} \beta' x_{i} + \sum_{i=1}^{n} \delta_{i} \log(\sum_{i \in R_{i}} e^{\beta' x_{j}}) + \lambda \|\beta\|_{1},$$
 (28)

where R_i is the set of indices for the patients at risk at time y_i -0. To compute the coefficients, we solve $H(\beta, \lambda) = 0$ for β , where H is defined as follows using only the current nonzero components of β :

$$H(\beta, \lambda) = \frac{\partial l}{\partial \beta} = -\sum_{i=1}^{n} \delta_{i} x_{i} + \sum_{i=1}^{n} \delta_{i} \sum_{j \in R_{i}} w_{ij} x_{j} + \lambda Sgn(\beta), \tag{29}$$

where $w_{ij} = e^{\beta' x_j} / \sum_{j \in R_i} e^{\beta' x_j}$.

We refer the readers to Appendix B for further details of the procedure.

4.2 Real data example

We demonstrate the L_1 regularization path algorithm for the Cox model using the heart transplant survival data introduced in Crowley & Hu (1977). The dataset consists of 172 samples with the following four features, as well as their survival time and censor information:

• age: age -48 years

• year: year of acceptance, in years after 11/1/1967

• surgery: prior bypass surgery, 1, if yes

• transplant: received transplant, 1, if yes

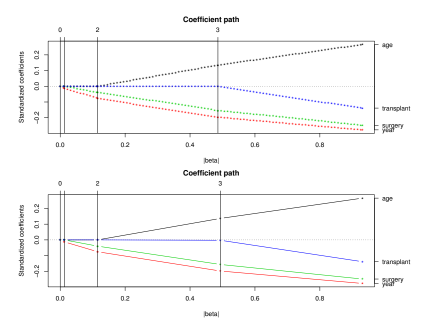


Figure 5: Survival data from Section 4.2: In the top panel, the coefficients were computed at fine grids of λ , whereas in the bottom panel, the solutions were computed only when the active set was expected to change. Similar to the GLM path examples, the exact coefficient paths shown on the top plot are almost piecewise linear; it is difficult to distinguish the two versions generated by different step sizes of λ .

In the top panel of Figure 5, the coefficients were computed at fine grids of λ , whereas in the bottom panel, the solutions were computed only when the active set was expected to change. Similar to the GLM path examples, the exact coefficient paths shown on the top plot are almost piecewise linear; it is difficult to distinguish the two versions generated by different step sizes in λ .

5 Discussion

In this paper, we have introduced a path-following algorithm to fit generalized linear models with L_1 regularization. As applied to regression (Tibshirani 1996, Tibshirani 1997) and classification methods (Genkin et al. 2004, Shevade & Keerthi 2003, Zhu, Rosset, Hastie & Tibshirani 2003), penalizing the size of the L_1 norm of the coefficients is useful because it accompanies variable selection. This strategy has provided us with a much smoother feature selection mechanism than the forward stepwise process.

Although the regularization parameter (λ in our case) influences the prediction performance in the aforementioned models considerably, determining the parameter can be troublesome or demand heavy computation. The GLM path algorithm facilitates model selection by implementing the predictor-corrector method and finding the entire regularization path sequentially, thereby avoiding independent optimization at different values of λ . Even with large intervals in λ , the predictor steps provide the subsequent corrector steps with reasonable estimates (starting values); therefore, the intervals can be wide without increasing the computations by a large number, as long as the paths can be assumed to be approximately linear within the intervals. Having generated the path, we estimate the globally optimal shrinkage level by cross-validating on the ratio λ/λ_{max} or $\log(\lambda)/\log(\lambda_{max})$. We compute the cross-validated loss at fixed values of the ratio rather than the values of λ because λ_{max} varies for every fold, and thus, a certain value of λ may correspond to different magnitude of shrinkage within each fold.

In Section 3.2.1, we proposed three different methods to determine the step lengths in λ and emphasized the efficiency and accuracy of the strategy of finding the transition points. One may suggest a more naive approach of pre-selecting certain values of λ and generating the coefficient paths by connecting the solutions to those grids. However, as shown in the comparison of the methods summarized in Table 1, such a strategy will generate paths that are either inaccurate or demand computations. In addition, computing the exact solutions at the values of λ where the active set changes ensures that we correctly specify the order in which variables are selected.

We can extend the use of the predictor-corrector scheme by generalizing the loss+penalty function to any convex and almost differentiable functions. For example, we can find the entire regularization path for the Cox proportional hazards model with L_1 penalization, as described in Section 4. Rosset & Zhu (2003) illustrated sufficient conditions for the regularized solution paths to be piecewise linear. Just as the solution paths for Gaussian distribution were computed with no error through the predictor-corrector method, so any other piecewise linear solution paths can be computed exactly by applying the same strategy.

The path-following algorithms for GLM and Cox proportional hazards model have been implemented in the contributed R package glmpath available from CRAN.

Appendix

A Proofs

Proof. Lemma 2.1

Minimizing (5) is equivalent to minimizing the following:

$$-\sum_{i=1}^{n} \{y_i \theta(\beta)_i - b(\theta(\beta)_i)\} + \sum_{j=1}^{p} \{\lambda(\beta_j^+ + \beta_j^-) - \lambda_j^+ \beta_j^+ - \lambda_j^- \beta_j^-\},$$
(30)

where $\beta = \beta^+ + \beta^-$, $\beta_j^+ \beta_j^- = 0$, $\beta_j^+, \beta_j^- \ge 0$ and $\lambda_j^+, \lambda_j^- \ge 0 \ \forall j = 1, \dots, p$. The Karush-Kuhn-Tucker (KKT) optimality conditions for this equivalent criterion are

$$-\mathbf{x}_{j}'\hat{\mathbf{W}}(\mathbf{y} - \hat{\boldsymbol{\mu}})\frac{\partial \eta}{\partial \mu} + \lambda - \lambda_{j}^{+} = 0, \tag{31}$$

$$-\mathbf{x}_{j}'\hat{\mathbf{W}}(\mathbf{y} - \hat{\boldsymbol{\mu}})\frac{\partial \eta}{\partial \mu} + \lambda - \lambda_{j}^{-} = 0, \tag{32}$$

$$\lambda_i^+ \hat{\beta}_i^+ = 0, \tag{33}$$

$$\lambda_j^- \hat{\beta}_j^- = 0, \quad \forall j = 1, \dots, p. \tag{34}$$

The KKT conditions imply

$$\left|\mathbf{x}_{j}'\hat{\mathbf{W}}(\mathbf{y} - \hat{\boldsymbol{\mu}})\frac{\partial \eta}{\partial \mu}\right| < \lambda \implies \hat{\beta}_{j} = 0 \text{ for } j = 1,\dots, p.$$
 (35)

When $\hat{\beta}_j = 0$ for all j = 1, ..., p, the KKT conditions again imply

$$\mathbf{1}'\hat{\mathbf{W}}(\mathbf{y} - \hat{\boldsymbol{\mu}})\frac{\partial \eta}{\partial \mu} = 0, \tag{36}$$

which, in turn, yields $\hat{\boldsymbol{\mu}} = \bar{y}\mathbf{1} = g^{-1}(\hat{\beta}_0)\mathbf{1}$.

Proof. Theorem 2.2

Since $\frac{\partial \boldsymbol{\beta}}{\partial \lambda} = -(\mathbf{X}_A' \mathbf{W}_k \mathbf{X}_A)^{-1} Sgn \begin{pmatrix} 0 \\ \hat{\beta}^k \end{pmatrix}$ is continuously differentiable with respect to $\lambda \in (\lambda_{k+1}, \lambda_k]$,

$$\hat{\boldsymbol{\beta}}^{k+1} = \hat{\boldsymbol{\beta}}^k - h_k \frac{\partial \boldsymbol{\beta}}{\partial \lambda} \Big|_{\lambda_k} + O(h_k^2)$$
 (37)

$$= \hat{\boldsymbol{\beta}}^{k+} + O(h_k^2), \tag{38}$$

from which the conclusion follows.

Proof. Theorem 2.3

Since $\partial \beta/\partial \lambda$ is continuously differentiable with respect to $\lambda \in (\lambda_{k+1}, \lambda_k]$, the following equations hold:

$$\hat{\boldsymbol{\beta}}(\lambda - \alpha h_k) = \hat{\boldsymbol{\beta}}^k - \alpha h_k \left(\frac{\hat{\boldsymbol{\beta}}^{k+1} - \hat{\boldsymbol{\beta}}^k}{-h_k}\right)$$
 (39)

$$= \left. \hat{\boldsymbol{\beta}}^{k} - \alpha h_{k} \frac{\partial \boldsymbol{\beta}}{\partial \lambda} \right|_{\lambda_{k}} + O(h_{k}^{2}), \tag{40}$$

and similarly the true solution at $\lambda = \lambda_k - \alpha h_k$ is

$$\beta(\lambda - \alpha h_k) = \hat{\beta}^k - \alpha h_k \frac{\partial \beta}{\partial \lambda} \Big|_{\lambda_k} + O(h_k^2). \tag{41}$$

The conclusion follows directly from the above equations.

Proof. Lemma 2.4

The Karush-Kuhn-Tucker (KKT) optimality conditions (31)-(34) imply

$$\hat{\beta}_j \neq 0 \implies \left| \mathbf{x}_j' \hat{\mathbf{W}} (\mathbf{y} - \hat{\boldsymbol{\mu}}) \frac{\partial \eta}{\partial \mu} \right| = \lambda.$$
 (42)

This condition, combined with (7) and (35), proves the argument.

B L_1 Regularization Path Algorithm for the Cox Model

Here we describe details of the L_1 regularization path algorithm for the Cox model, briefly introduced in Section 4. We use the same notations as presented in Section 4.

The Cox model with L_1 penalization finds the coefficients that minimize the following objective function:

$$l(\beta, \lambda) = -\sum_{i=1}^{n} \delta_{i} \beta' x_{i} + \sum_{i=1}^{n} \delta_{i} \log(\sum_{j \in R_{i}} e^{\beta' x_{j}}) + \lambda \|\beta\|_{1}.$$
 (43)

The first and the second derivatives of l with respect to β are as follows:

$$\frac{\partial l}{\partial \beta} = H(\beta, \lambda) = -\sum_{i=1}^{n} \delta_{i} x_{i} + \sum_{i=1}^{n} \delta_{i} \sum_{j \in R_{i}} w_{ij} x_{j} + \lambda Sgn(\beta)$$

$$(44)$$

$$\frac{\partial^2 l}{\partial \beta \partial \beta'} = \frac{\partial H}{\partial \beta} = \sum_{i=1}^n \partial_i \{ \sum_{j \in R_i} x_j x_j' w_{ij} - (\sum_{j \in R_i} x_j w_{ij}) (\sum_{j \in R_i} x_j' w_{ij}) \}$$
(45)

$$= \mathbf{X}'\mathbf{A}\mathbf{X},\tag{46}$$

where $w_{ij} = e^{\beta' x_j} / \sum_{j \in R_i} e^{\beta' x_j}$, and $\mathbf{A} = \frac{\partial l^2}{\partial \eta \partial \eta'}$ with $\eta = \mathbf{X}\beta$.

If $\beta_j = 0$ for j = 1, ..., p, then $w_{ij} = 1/|R_i|$ for i = 1, ..., n and j = 1, ..., p, and

$$\frac{\partial l}{\partial \beta} = -\sum_{i=1}^{n} \delta_i (x_i - \frac{1}{|R_i|} \sum_{j \in R_i} x_j). \tag{47}$$

 $\hat{\beta}_j = 0$ for all j if $\lambda > \max_{j \in \{1,...,p\}} |\partial l/\partial \beta_j|$. As λ is decreased further, an iterative procedure begins; variables enter the active set, beginning with $j_0 = argmax_j |\partial l/\partial \beta_j|$. The four steps of an iteration are as follows:

1. Predictor step

In the k-th predictor step, $\beta(\lambda_{k+1})$ is approximated as in (8), with

$$\frac{\partial \beta}{\partial \lambda} = -\left(\frac{\partial H}{\partial \beta}\right)^{-1} \frac{\partial H}{\partial \lambda} = -(\mathbf{X}_A' \mathbf{A} \mathbf{X}_A)^{-1} Sgn(\beta). \tag{48}$$

 \mathbf{X}_A contains the columns of \mathbf{X} for the current active variables.

2. Corrector step

In the k-th corrector step, we compute the exact solution $\beta(\lambda_{k+1})$ using the approximation from the previous predictor step as the initial value.

3. Active set

Denoting the correlation between the factors and the current residual as $\hat{\mathbf{c}}$,

$$\hat{\mathbf{c}} = \sum_{i=1}^{n} \delta_i x_i - \sum_{i=1}^{n} \delta_i \sum_{j \in R_i} w_{ij} x_j. \tag{49}$$

After each corrector step, if $|\hat{\mathbf{c}}_l| > \lambda$ for any $l \in \mathcal{A}^c$, we augment the active set by adding x_l . Corrector steps are repeated until the active set is not augmented further. If $\hat{\beta}_l = 0$ for any $l \in \mathcal{A}$, we eliminate x_l from the active set.

4. Step length

If $\lambda = 0$, the algorithm stops. If $\lambda > 0$, we approximate the smallest decrement in λ with which the active set will be modified. As λ is decreased by h, the approximated change in the current correlation (49) is as follows:

$$\mathbf{c}(h) = \hat{\mathbf{c}} - h\mathbf{X}'\mathbf{A}\mathbf{X}_A(\mathbf{X}_A\mathbf{A}\mathbf{X}_A)^{-1}Sgn(\beta). \tag{50}$$

Based on (50), we approximate the next largest λ at which the active set will be augmented/reduced.

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