Penalized Regression in the Age of Big Data

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Abstract

With the prevalence of big data in the modern age, the importance of modeling high dimensional data and selecting influential features has increased greatly. High dimensional data is common in many fields such as genome decoding, rare disease identification, economic modeling, and environmental modeling. However, most traditional regression and classification machine learning models are not designed to handle high dimensional data or conduct variable selection. In this paper, we investigated the use of penalized regression methods instead of, or in conjunction with, the traditional machine learning methods. We focused on lasso, ridge, elastic net, SCAD, MCP, and adaptive versions of lasso, ridge, and elastic net models. For traditional machine learning models, we focused on random forest models, gradient boosting models in the form of XGBoost, and support vector machines. These models were evaluated using factorial design methods for Monte Carlo simulations under various data environments. Tests were conducted for 270 environments, with factors being the number of predictors, number of samples, signal to noise ratio, covariance matrix, and correlation strength. This served to identify the strengths and weaknesses of different penalization techniques in different environments. We also compared different models using empirical datasets to test their viability in real-world scenarios. Additionally, we considered penalization methods outlined earlier in logistic regression models for classifying data. These results were compared to random forest, gradient boosting, and support vector machine classification models using both Monte Carlo data generation methods and empirical data. For regression, we evaluated the models using the test mean squared error and variable selection accuracy; for classification, we considered test prediction accuracy and variable selection accuracy. We found that for both regression and classification, penalized regression models outperformed more traditional machine learning algorithms in most high-dimensional situations or in situations with a low number of data observations. By comparing traditional machine learning methods with penalized regression, we hope to expand the scope of machine learning methods for big data to include the various penalized regression techniques we tested. Additionally, we hope to create a greater understanding of the strengths and weaknesses of each model type and provide a reference for other researchers on which machine learning techniques they should use, depending on a range of factors.

Keywords: penalized regression, variable selection, classification, machine learning, large p little n problem, Monte Carlo simulations

1 Introduction

[Better intro paragraph]

Typically, data sets are represented as a table of values. Most columns represent **predictors** (also called variables, attributes, or features), while the rows represent **observations** (also called instances). The value

in the i-th row and j-th column represents the value for predictor j in observation i. At least one column is designated as a **response**, which is assumed to be related to some of the predictors in some way. Machine learning models attempt to predict the value of this response from the values of the predictors.

Let n be the number of observations for a data set, and let p be the number of predictors. In most situations, the number of observations greatly exceeds the number of predictors. However, as data collection becomes easier and as statistical modeling techniques are introduced to new disciplines, situations can arise where there are more predictors than observations. For instance, in the field of genomics, there may be thousands of genes that could cause a disease and only a few samples to train from.

In situations where there are more predictors than observations, many traditional machine learning techniques fail to give good predictions. The large number of predictors makes it easy for such models to **overfit**, meaning that the models make good predictors from the data used to train the model, but perform badly when given new data.

To resolve this "large p, little n" problem, many algorithms have been introduced to address situations where there are more predictors than observations. Many, but not all, of these techniques use **variable selection**, meaning that they select the predictors that are most correlated with the response. By ignoring predictors that are not strongly related to the response, the negative consequences of overfitting can be greatly reduced.

This paper investigates various methods used to handle the large p, small n problem. We considered subset selection methods such as forward selection, backward selection, stepwise forward selection and stepwise backward selection. In addition, we studied penalized regression models such as ridge regression, LASSO, elastic-net, adaptive LASSO, SCAD, and MCP. Models were trained and evaluated using both Monte Carlo simulations and empirical genomic data.

1.1 Background

Suppose that we have p predictor variables X_1, X_2, \ldots, X_p and one response variable Y that depends on some (or all) of the predictors. We assume that Y can be expressed as

$$Y = f(X_1, X_2, \dots, X_p) + \epsilon \tag{1}$$

where f is a function and ϵ is an independent random error with mean zero. The goal of supervised modeling is to find a function \hat{f} that is a suitable approximation for f. To find \hat{f} , we use a **training set**, a set of observations where the response variable Y is already known. Then, using the fitted model, we can predict the value of the response variable \hat{Y} for new observations, even if Y is unknown. Model performance can be evaluated using a **test set**, which is a set of observations that were not used to train the model.

There are two broad types of supervised models. **Regression modeling** is used when the response variable Y takes numerical values on a continuous interval. For example, a model that predicts the value of a home is a regression model. On the other hand, if Y can only take discrete values, then **classification modeling** is used. For instance, a model used to predict whether or not a patient has a disease is classification problem. This paper focuses on regression modeling.

1.2 Linear Regression and Ordinary Least Squares

In practice, the function f that relates the predictors to the response is complex. Most statistical models assume that f takes some particular form and estimates a function \hat{f} of that form. For example, many regression models assume that f is a linear function of the predictors; that is, linear models assume that

$$f(X_1, X_2, \dots, X_p) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots + \beta_p X_p$$
 (2)

where $\beta_0, \beta_1, \beta_2, \dots, \beta_p$ are coefficients. Notice that the coefficient β_0 is not multiplied with any predictor; it represents an intercept value. Fitting a linear model will give estimates for these coefficient values.

The most common method to approximate the coefficients in a linear model is by **ordinary least** squares. Suppose that we have n observations in our training set. Let x_{ij} represent the value of predictor j for observation i, and let y_i be the response for observation i. For some coefficient estimates $\beta_0, \beta_1, \beta_2, \ldots, \beta_p$, the expression

$$y_i - (\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_p x_{ip})$$
(3)

is called the **residual** for observation i; it is the difference between the true response value and the predicted response variable using the given coefficient values. Ordinary least squares chooses the coefficients $\beta_0, \beta_1, \beta_2, \ldots, \beta_p$ that minimize the **residual sum of squares**

$$RSS = \sum_{i=1}^{n} (y_i - (\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_p x_{ip}))^2$$
 (4)

Intuitively, if the residual sum of squares is low, then the differences between the response variable and its estimates is low. Thus, by minimizing the residual sum of squares, the function obtained from ordinary least squares is a relatively good approximation for f. Figure 1 demonstrates a model fitted with ordinary least squares when there is a single predictor variable.

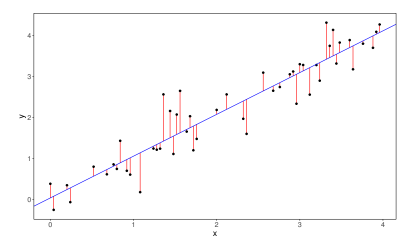


Figure 1: Ordinary least squares fitting with one predictor using simulated data. The blue line represents the line found by ordinary least squares, and the red line segments are the residuals.

One reason that ordinary least squares is popular is because it is very easy to compute. Let $\beta = [\beta_0, \beta_1, \beta_2, \dots, \beta_p]^{\top}$ be a $(p+1) \times 1$ vector of coefficient values and let \mathbf{X} be a $n \times (p+1)$ matrix where each row contains the predictor values for one observation (with an extra value of 1 in the first entry). Then $\mathbf{X}\beta$ is a vector of the estimated response values. Let \mathbf{y} represent the true response values. Then $\mathbf{y} - \mathbf{X}\beta$ is a vector of residuals. To choose coefficient estimates that minimize the residual sum of squares, we compute

$$\hat{\beta}^{\text{OLS}} = \arg\min_{\beta} \left\{ (\mathbf{y} - \mathbf{X}\beta)^{\top} (\mathbf{y} - \mathbf{X}\beta) \right\}$$
 (5)

where $(\mathbf{y} - \mathbf{X}\beta)^{\top}(\mathbf{y} - \mathbf{X}\beta)$ is the same residual sum of squares seen in Equation 4. From [3], this gives us the solution

$$\hat{\beta}^{\text{OLS}} = (\mathbf{X}^{\top} \mathbf{X})^{-1} \mathbf{X}^{\top} \mathbf{y}$$
 (6)

Another advantage of ordinary least squares is that it is an unbiased linear model. This means that if the relationship between the response variable and the predictors truly is linear (as given in Equation 2), then the expected value of the coefficient vector $\hat{\beta}^{\text{OLS}}$ is equal to the actual coefficient vector β . Furthermore, the **Gauss-Markov theorem** states that if the random error ϵ is independent and has constant variance, then ordinary least squares has the lowest variance among all linear unbiased estimators. In other words,

the coefficient estimates given by ordinary least squares are relatively close to the actual coefficient values when compared to other unbiased linear estimators. This makes ordinary least squares a relatively consistent model.

If ordinary least squares is unbiased and has the lowest variance among all unbiased models, why should we use any other type of linear model? Despite having a lower variance than other unbiased models, ordinary least squares can still have a high variance. This is especially an issue when the number of predictors p is large compared to the number of observations n. As p gets closer to n, a model fitted with ordinary least squares will typically **overfit** to the training set. This means that the fitted model makes very good predictions with the training data, but performs poorly when given test data that wasn't used to fit the model. Overfitting occurs because of the random error from Equation 1. Ordinary least squares is unable to distinguish between signal and noise, so it will tend to assume predictors are more strongly related to the response than they actually are.

In the extreme case where p exceeds n, the matrix $\mathbf{X}^{\top}\mathbf{X}$ from Equation 6 becomes non-invertible. This means that there are many coefficient estimates that minimize the residual sum of squares. In fact, any of these coefficient estimates creates a perfect fit to the training data, which will result in very bad predictions with test data.

By using models that have a small amount of bias, the high variance of ordinary least squares can be mitigated. Liu et. al. in [7] describe three types of variable selection algorithms. **Filter methods** work by evaluating the ability for each individual predictors to predict the response; then, a model is fit using the predictors selected. **Wrapper methods** fit models using different subsets of predictors and choose the model that has the best performance. Finally, **embedded methods** perform variable selection during the model training process. This paper focuses on the wrapper methods and embedded methods. In addition, we also used several non-linear machine learning methods to draw a comparison between linear regression models and non-linear models.

1.3 Subset Selection Methods

Subset selection methods are wrapper methods that attempt to find a subset of the predictors X_1, X_2, \ldots, X_p that are most correlated with the response variable Y. These algorithms usually fit models for many different subsets and choose the subset of predictors that results in the best model. Although subset selection techniques can be applied to many types of models, we will focus on subset selection with linear regression.

There are two main benefits to using subset selection methods. By reducing the set of available predictors to just those that are strongly related to the response, overfitting can be mitigated by ignoring predictors that provide little improvement to model performance. Another benefit of subset selection is that it creates a more interpretable model. If a data set includes thousands of predictors but only a few are related to the response, a model found using subset selection will be easy to understand than a model that relies on all of the parameters.

Best subset selection is a subset selection method that fits considers every possible combination of predictors. For every possible subset size k between 0 and p, best subset selection will fit the $\binom{p}{k}$ possible models using k predictors. Then, the best model for each value of k is chosen based on some performance metric. Finally, a final model can be selected from the (p+1) remaining models. Although best subset selection is guaranteed to find the subset of predictors that optimize the chosen metric, this method is computationally expensive. For a data set with p predictors, 2^p possible combinations must be considered. This makes best subset selection infeasible when the number of predictors is too large.

Two alternative methods to best subset selection are **forward selection** and **backward selection**. Forward selection begins by fitting a model with no predictors and iteratively adds predictors into the model. The predictor added at each step is chosen to best increase the model fit. Conversely, backward selection starts from the full (ordinary least squares) model with all p predictors and repeatedly removes predictors. Then, like best subset selection, the final model is chosen from the candidate models fitted at each step. Note that backward selection can only be used when $p \le n$, since ordinary least squares cannot

be used when p > n. Although forward and backward selection are not guaranteed to encounter the best possible model, these methods avoid the exponential runtime of best subset selection. Consequently, forward and backward selection can be used for larger values of p.

The models produced by forward and backward selection can be improved by allowing for predictors to be added and removed in the same algorithm. Forward stepwise selection begins with an empty model and iteratively improves the model by either adding a new predictor or removing an obsolete one. Backward stepwise selection works in the same way but starts with the full model. Like backward selection, backward stepwise selection can only be used when $p \ge n$. These techniques take longer to run than ordinary forward and backward selection, but they are more likely to find the best possible model.

When fitting a model using any of the subset selection methods, the performance metric used when selecting the best model is very important. At first, it may seem reasonable to choose a common metric such as the residual sum of squares from Equation 4. However, many metrics, including the residual sum of squares, only describe a model's performance on training data. This is problematic because including more predictors will always decrease the residual sum of squares on the training data. If $p \leq n$, then the model fitted with all p predictors is exactly the same model produced ordinary least squares, which by definition minimizes the residual sum of squares! If p > n, then the full model cannot be fitted; instead, subset selection methods would choose a model with the maximum possible number of predictors.

If we wish to produce a model that makes reliable predictions on test data, we must use a different performance metric. Two of the most common metrics used for this purpose are the **Akaike Information Criterion** (AIC) and the **Bayesian Information Criterion** (BIC). These metrics can be expressed in terms of the log-likelihood function. In the special cases where we have a linear model where the random error ϵ is Gaussian, the Akaike Information Criterion is given by

$$AIC = \frac{1}{n\hat{\sigma}^2} (RSS + 2p\hat{\sigma}^2) \tag{7}$$

up to a constant, where $\hat{\sigma}^2$ is the estimated value of the variance of ϵ [5]. The Bayesian Information Criterion is

$$BIC = \frac{1}{n\hat{\sigma}^2} (RSS + \ln(n)\hat{\sigma}^2)$$
 (8)

up to a constant. These metrics work by using the residual sum of squares plus some additional penalty that increases when p is large. As a result, models that minimize AIC or BIC will have fewer predictors than models chosen just by minimizing the residual sum of squares. This can result in a model that has both a good training error and test error. If n > 7, then $\ln(n) > 2$ and so the penalty for BIC is larger than the penalty for AIC. Hence, a model selected using BIC will typically have fewer parameters than a model selected by AIC.

In addition to AIC and BIC, there are several other metrics that modify training error to estimate test error, such as C_p and adjusted R^2 [5]. However, this paper will focus on AIC and BIC.

1.4 Penalized Regression

In general, **penalized regression** works by fitting a model that punishes large coefficient estimates. By forcing coefficient values to shrink, the resulting model will have relatively low variance. Most, but not all, of these methods can perform variable selection.

All of the penalized regression methods in this paper solve an optimization problem of the form

$$\arg\min_{\beta} \left\{ \sum_{i=1}^{N} (y_i - (\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_p x_{ip})) + \sum_{j=1}^{p} P(\beta_j) \right\}$$
(9)

where the first summation is the usual residual sum of squares and $P(\beta_j)$ is a penalty function applied to each of the coefficients. In general, $P(\beta_i)$ is an even function that is non-decreasing as $|\beta_i|$ increases.

Ridge regression helps to solve multicollinearity in predictors while also minimizing insignificant predictors [4]. While it does not minimize these insignificant predictors completely to 0 and thus cannot be considered a variable selection method, it still proves very useful in large datasets.

Ridge regression works by minimizing Residual Sum Squared (RSS) plus a penalty as seen in Equation 10. λ is a tuning parameter and can be used to determine how much of an effect the penalty has on the regression. if $\lambda = 0$, then the regression acts exactly like ordinary least squares regression, but if $\lambda \to \infty$, then $\beta_i \to 0$ and the regression line will be a horizontal line at the intercept, β_0 .

$$\hat{\beta}^{\text{ridge}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{j=1}^{p} \beta_j x_{ij} \right)^2 + \lambda \sum_{j=1}^{p} \beta_j^2 \right\}$$
(10)

An alternative way to express ridge regression is with the equation

$$\hat{\beta}^{\text{ridge}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{j=1}^{p} \beta_j x_{ij} \right)^2 \right\} \quad \text{subject to} \quad \sum_{j=1}^{p} \beta_j^2 \le t$$
 (11)

for some tuning parameter t.

The **least absolute shrinkage and selection operation**, often referred to as LASSO, is a shrinkage method with a very similar form to lasso regression [8, 5, 6]. The coefficient estimates satisfy

$$\hat{\beta}^{\text{lasso}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{j=1}^{p} \beta_j x_{ij} \right) + \lambda \sum_{j=1}^{p} |\beta_j| \right\}$$
(12)

If $\lambda = 0$, then the lasso model is equivalent to the ordinary least squares model; if $\lambda \to \infty$, then the coefficients for all predictors will be set to 0. An equivalently way to define lasso regression is by

$$\hat{\beta}^{\text{lasso}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{j=1}^{p} \beta_j x_{ij} \right)^2 \right\} \quad \text{subject to} \quad \sum_{j=1}^{p} |\beta_j| \le t$$
 (13)

where t is a tuning parameter.

One useful property of the lasso method is that it can perform variable selection by setting some coefficients to zero. To understand why lasso regression can perform variable selection whereas ride regression cannot, consider Figure 2 below. This figure demonstrates the case when p=2 and t=1. The circle in 2a represents the condition $\beta_1^2 + \beta_2^2 < 1$ for ridge regression, while the red diamond in Figure 2b represents the condition $|\beta_1| + |\beta_2| < t$ for lasso regression. The blue curves represent contours of the residual sum of squares for values of β_1 and β_2 . The black point in the center of these curves is where the RSS is minimized; this represents the values of β_1 and β_2 that would be selected by ordinary least squares.

In 2a, the intersection of the black curve and the red diamond represents the parameter values chosen by ridge regression; this point minimizes the RSS under the condition $\beta_1^2 + \beta_2^2 \le 1$. Because the red region is a circle, this intersection cannot occur exactly at $\beta_1 = 0$; hence, the ridge method cannot remove the predictor β_1 . On the other hand, the square shape of the constrained region for lasso regression can perform variable selection because the intersection occurs at one of the axes.

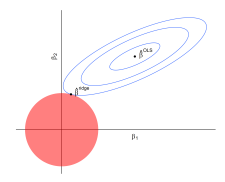
The lasso method is particularly useful in the case where p > n because of its ability to select variables; a model with fewer variables has less variance and is more interpretable. One major downside of lasso regression is that it does not handle multicollinearity as nicely as ridge regression. Another downside of lasso regression is that it does not have a closed-form solution, which can lead to instability in the model.

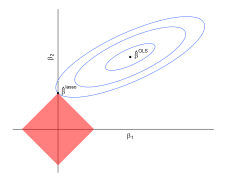
Elastic-net regression serves as a combination between ridge and lasso regression. It can handle multicollinearity as well as perform variable selection. The coefficients for elastic net regression can be determined

Figure 2: RSS contours and penalty bounds for the ridge and lasso models when p=2 and t=1.

(a) RSS contours and the ridge penalty boundary.







by

$$\hat{\beta}^{\text{ENet}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{i=1}^{p} \beta_j x_{ij} \right) + \lambda_2 \sum_{j=1}^{p} \beta_j^2 + \lambda_1 \sum_{j=1}^{p} |\beta_j| \right\}$$
(14)

where λ_1 and λ_2 are both tuning parameters to be determined later.

An important limitation to note is that elastic net performs best when it close to either ridge or lasso regression, meaning that either $\lambda_1 >> \lambda_2$ or vice versa [11]. Additionally, because elastic net requires two tuning parameters, this makes it much more difficult to determine the best combination of tuning parameters to minimize error in the regression. However, this problem has been largely solved through by the LARS-EN algorithm developed by Zou et. al. which efficiently solves for the tuning parameters.

Normally in lasso regression, each predictor is weighted the same in the penalty function. **Adaptive** lasso regression is different in that a weight, \hat{w}_j is multiplied to the penalty function. The coefficients for adaptive lasso regression as designed by Zou et. al. [10] can be defined by

$$\hat{\beta}^{\text{adaptive}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{i=1}^{p} \beta_j x_{ij} \right) + \lambda \sum_{j=1}^{p} \hat{w}_j |\beta_j| \right\}$$
(15)

where λ is a tuning parameter to be determined later and \hat{w}_j is defined as $\frac{1}{|\hat{\beta}|^{\gamma}}$ with γ being a chosen parameter greater than 0.

Because of the weight that is implemented in adaptive lasso regression, zero-coefficients have a weight that is inflated up to infinity, and thus are punished much more harshly than large coefficients whose weight is much smaller in comparison. This is a similar rationale to SCAD and helps to reduce some of the bias from lasso regression. Bridge regression is the general form of lasso regression from which adaptive lasso originates from. When $\gamma < 1$, bridge regression as shown in Equation 16 is not continuous, which results in model prediction instability. However, adaptive lasso regression is completely continuous and thus has much more consistent coefficients when fitted.

$$\hat{\beta}^{\text{lasso}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{i=1}^{p} \beta_j x_{ij} \right) + \lambda \sum_{j=1}^{p} |\beta_j|^{\gamma} \right\}$$
(16)

One major flaw of the lasso method is that the penalty punishes large coefficients, even if those coefficients should be large. One way to modify the lasso method is to use the **smoothly clipped absolute deviation** (SCAD) penalty [2]. The goal of this method is to punish large coefficients less severely, which can help

mitigate some of the bias introduced by the lasso method.

$$\hat{\beta}^{\text{SCAD}} = \arg\min_{\beta} \left\{ \sum_{i=1}^{n} \left(y_i - \beta_0 - \sum_{i=1}^{p} \beta_j x_{ij} \right) + \lambda \sum_{j=1}^{p} J_a(\beta_j, \lambda) \right\}$$
(17)

Here, $J_a(\beta, \lambda)$ is a penalty function that satisfies

$$\frac{dJ_a(\beta,\lambda)}{d\beta} = \lambda \cdot \operatorname{sign}(\beta) \left[I(|\beta| < \lambda) + \frac{(a\lambda - |\beta|)_+}{(a-1)\lambda} I(|\beta| > \lambda) \right]$$
(18)

where $\lambda \geq 0$ and $a \geq 2$ are tuning parameters. An equivalent way to write this is

$$\frac{dJ_a(\beta,\lambda)}{d\beta} \begin{cases} \lambda, & |\beta| \le \lambda \\ \frac{a\lambda - |\beta|}{a - 1}, & \lambda < |\beta| < a\lambda \\ 0, & \alpha\lambda < |\beta| \end{cases}$$
(19)

This penalty function does not punish coefficients with large magnitude as heavily as the lasso method. In fact, if the magnitude of a coefficient is larger than $a\lambda$, then the penalty becomes constant. See Figure 3a for a plot of the SCAD penalty as a function of the coefficient value.

Integrating with respect to β [1], we see that

$$J_{a}(\beta,\lambda) = \begin{cases} \lambda|\beta|, & |\beta| \leq \lambda\\ \frac{2a\lambda|\beta| - \beta^{2} - \lambda^{2}}{2(a-1)}, & \lambda < |\beta| < a\lambda\\ \frac{\lambda^{2}(a+1)}{2}, & a\lambda < |\beta| \end{cases}$$
(20)

The **minimax concave penalty** (MCP) method is very similar to smoothly clipped absolute deviation [9, 1]. Both methods are used to avoid the high bias caused by the lasso method. MCP uses a penalty function that satisfies

$$\frac{dJ_a(\beta,\lambda)}{d\beta} = \begin{cases} \operatorname{sign}(\beta) \left(\lambda - \frac{|\beta|}{a}\right), & |\beta| \le a\lambda \\ 0, & a\lambda < |\beta| \end{cases}$$
 (21)

where $\lambda \geq 0$ and a > 0 are tuning parameters. Integrating [1], we see that

$$J_a(\beta, \lambda) = \begin{cases} \lambda |\beta| - \frac{\beta^2}{2a}, & |\beta| \le a\lambda \\ \frac{1}{2}a\lambda^2, & a\lambda < |\beta| \end{cases}$$
 (22)

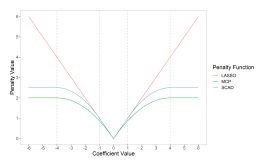
Figure 3 below shows the penalty functions (and their derivatives) for LASSO, SCAD, and MCP as a function of a coefficient value β . We see that LASSO applies a much stronger penalty to large coefficients than SCAD or MCP. Also, note that SCAD starts with a derivative equal to that of the lasso for small values of β ; on the other hand, the derivative of the penalty function for MCP starts decreasing immediately.

Now, consider the case where p=1 (there is only one predictor). Figure 4 shows the solutions given when using LASSO, SCAD, and MCP on such a model. The x-axis gives the actual coefficient for the single variable, and the y-axis represents the coefficient estimate produced using each of the algorithms. We used the particular values $\lambda=2$ and a=3. The gray line is the identity function, which also equals the solution obtained using ordinary least squares.

We see that all three methods set the predicted value to zero when the actual coefficient is small. Also, note that the LASSO method is always off from the identity function when the coefficient is large. On the other hand, SCAD and MCP merge with the identity function when the coefficient is sufficiently large. This shows that both SCAD and MCP can avoid the high bias that LASSO introduces.

Figure 3: Penalty functions for LASSO, SCAD, and MCP, as well as their derivatives. These plots use $\lambda = 2$ and a = 3.

(a) Penalty functions for LASSO, SCAD, and MCP



(b) Derivatives of the penalty functions for LASSO, SCAD, and MCP

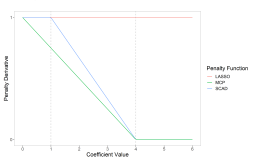
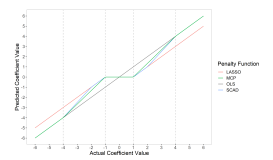


Figure 4: Solutions for LASSO, SCAD, and MCP for a single predictor when $\lambda=2,$ and a=3.



2 Methods

2.1 Models

2.2 Monte Carlo Simulations

Monte Carlo simulations use randomly generated data to fit and test our regression and classification models. There are several benefits to using simulated data rather than experimental data:

- The true relationship between the predictor variables and the response is known.
- Simulations can be iterated many times, giving sturdier results about the effectiveness of each model.
- We have full control over factors such as the number of predictors and the amount of correlation between predictors.

For the simulated data, we assumed that the relationship between the response variable y and the predictors x_1, x_2, \ldots, x_p was linear. That is, we assumed that

$$y = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p + \epsilon \tag{23}$$

where β_0 is some intercept, β_1, \ldots, β_p are coefficient values and $\epsilon \sim \mathcal{N}(0, \sigma^2)$ is a normally distributed random error with mean 0 and variance σ^2 .

To generate the data, we first defined $\beta = [\beta_0, \beta_1, \dots, \beta_p]^{\top}$, a $(p+1) \times 1$ vector of coefficient values. For our simulations, we used $\beta_0 = 1$, $\beta_1 = 2$, $\beta_2 = -2$, $\beta_5 = 0.5$ and $\beta_6 = 3$; the remaining coefficient values were set to 0.

Next, we generated \mathbf{X} , a $n \times (p+1)$ matrix of predictor variables. The first column contains 1 in all of its entries; this corresponds to the intercept of our linear model. Column i of \mathbf{X} contains the variable values for predictor x_{i-1} , for $1 \le i \le p$. These values were generated using the p-dimensional multivariate normal distribution $\mathcal{N}_p(0, \mathbf{\Sigma})$ with mean zero and covariance matrix $\mathbf{\Sigma}$. We assumed that every predictor had a standard deviation of 1, making the covariance matrix equivalent to a correlation matrix. Four different correlation matrix structures were considered in our study.

We then generated an $n \times 1$ error vector $\mathbf{e} \sim \mathcal{N}(0, \sigma^2)$ with mean zero and variance σ^2 . For regression models, the response \mathbf{y} can then be computed by

$$\mathbf{y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{e} \tag{24}$$

We used a **factorial design** for our simulations. This means that we considered several factors that affect the data generation process, each having multiple possible values. We then generated data using every possible combination of factor values, giving us a comprehensive assessment of model performance under various conditions.

- n, the number of observations: 50, 200, 1000.
- p, the number of predictors: 10, 100, 2000.
- σ , the standard deviation of the random error: 1, 3, 6.
- The correlation matrix structure: independent, symmetric compound, autoregressive, blockwise.
- ρ , the correlation between predictors: 0.2, 0.5, 0.9.

By taking every possible combination of these factors, we obtain $3 \times 3 \times 3 \times 4 \times 3 = 324$ different settings for the simulations. However, because an independent correlation matrix does not use the correlation value ρ , we actually only used 270 combinations. For each combination of factors, we ran 100 simulations. In each simulation, we generated two data sets: one to train the various models, and one to test the models and evaluate performance. Both data sets contained n observations, meaning that a total of 2n observations were generated for each simulation.

As mentioned earlier, we considered four different covariance matrix structures. These structures determine the correlation between different predictors. If Σ is a correlation matrix, then Σ_{ij} , the entry at the i-th row and j-th column, represents the correlation between predictors i and j. If $\Sigma_{ij} = 0$, there is no correlation; but if $\Sigma_{ij} = 1$, then predictors i and j are perfectly correlated. Note that a correlation matrix is always symmetric, so $\Sigma_{ij} = \Sigma_{ji}$ for all indices i and j. This correlation can severely impact the performance of statistical models; if several predictors are highly correlated, then machine learning algorithms are less able to determine which predictors are actually related to the response.

The first correlation structure we considered is **independent correlation**. This means that the correlation matrix Σ has the form

$$\Sigma = \begin{bmatrix} 1 & 0 & \cdots & 0 \\ 0 & 1 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 1 \end{bmatrix}$$
 (25)

In other words, there is no correlation between different predictors, since $\Sigma_{ij} = 0$ whenever $i \neq j$.

The next covariance structure is called **symmetric compound**. This structure has the form

$$\Sigma = \begin{bmatrix} 1 & \rho & \cdots & \rho \\ \rho & 1 & \cdots & \rho \\ \vdots & \vdots & \ddots & \vdots \\ \rho & \rho & \cdots & 1 \end{bmatrix}$$
 (26)

where $\rho \in [0, 1]$ is some correlation value. A symmetric compound covariance structure assumes that $\Sigma_{ij} = \rho$ whenever $i \neq j$, meaning that all predictors are equally correlated with one another.

An autoregressive covariance structure assumes that

$$\Sigma = \begin{bmatrix} 1 & \rho & \cdots & \rho^{p-1} \\ \rho & 1 & \cdots & \rho^{p-2} \\ \vdots & \vdots & \ddots & \vdots \\ \rho^{p-1} & \rho^{p-2} & \cdots & 1 \end{bmatrix}$$

$$(27)$$

For any indices i and j, we have $\Sigma_{ij} = \rho^{|i-j|}$. Consequently, each predictor is strongly correlated with nearby predictors and weakly correlated with more distant predictors. This form of covariance is commonly seen when using time series, since observed values at nearby times are likely to be highly correlated with one another.

Finally, a blockwise correlation matrix has the block-diagonal form

$$\Sigma = \begin{vmatrix} \mathbf{B}_1 & 0 & \cdots & 0 \\ 0 & \mathbf{B}_2 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & \mathbf{B}_k \end{vmatrix}$$
 (28)

where 0 represents a block containing all zeroes, and each block \mathbf{B}_i has a form identical to the symmetric compound matrix in Equation 26. This implies that predictors within the same block have correlation $\rho \in [0, 1]$, whereas predictors in different blocks have zero correlation. One important consideration when using blockwise correlation is the size of each block. For our simulations, we used a block size of 5 when p = 10, a block size of 25 when p = 100, and a block size of 100 when p = 2000.

All of our simulations were run using version 4.1.0 of R. Several different libraries were used to fit machine learning models using our simulated data. Table 1 summarizes the libraries used to fit models.

Library	Models used	Version
stats	Ordinary least squares	4.1.0
MASS	Forward/Backward selection	7.3 - 54
glmnet	Ridge, lasso, elastic-net	4.1-1
gcdnet	Adaptive ridge, adaptive lasso, adaptive elastic-net	1.0.5
ncvreg	SCAD and MCP	3.13.0
xgboost	Gradient boosting	1.4.1.1
ranger	Random forest	0.12.1
e1071	Support vector machine	1.7-7

Table 1: R Libraries used and the models used from each library

For ridge, lasso, and elastic-net regression using glmnet, we used the cv.glmnet function. This function uses cross-validation to determine the value of λ that minimizes the cross-validation error. We used ten folds with cv.glmnet. Using cross-validation can help generate a model that has a good test performance. For elastic-net regression, we used the hyperparameter $\alpha=0.8$. This means that the elastic-net model is more similar to lasso (where $\alpha=1$) than ridge (where $\alpha=0$). The remaining hyperparameters were given their default values.

We used the cv.gcdnet function from the gcdnet library for the adaptive versions of ridge, lasso, and elastic-net. Again, ten folds were used for the cross-validation, and all hyperparameters were given their default values.

For SCAD and MCP models, we used the cv.ncvreg function from the ncvreg library. We used the default values of a for both models: 3 for MCP and 3.7 for SCAD (note that the ncvreg documentation calls this hyperparameter γ instead of a).

For the three non-linear models (gradient boosting, random forests, and support vector machines), we used cross-validation and grid search to find suitable hyperparameters, and then fit a model using the full training set using those hyperparameters. For gradient boosting with xgboost, we used different values for the learning rate (0.1, 0.3, and 0.5) and maximum tree depth (1, 3, and 7). A maximum of 1000 trees were generated, with an early stopping condition if the model failed to improve for several iterations in a row. The cross-validation function used five folds.

For random forests using ranger, we tuned the number of predictors used per decision tree $(\lfloor \sqrt{p} \rfloor, \lfloor p/3 \rfloor,$ and $\lfloor p/2 \rfloor)$ and the number of trees (300, 400, 500 and 600). The cross-validation function used five folds.

Finally, for support vector machines using e1071, we varied ϵ (TODO: explain this) and the cost function (0.5, 1, and 2).

2.3 Empirical Data

For empirical data, we used the Breast Cancer database from The Cancer Genome Atlas (bcTCGA). This contains the gene expression data of 17323 genes from 536 patients. One of these genes is the BRCA1 gene which is among the first genes discovered that can increase the risk of breast cancer. Because the BRCA1 gene interacts with other genes, it is useful to find genes that interact with BRCA1 to test in further studies. The distribution of the BRCA1 gene expression levels in the bcTCGA database can be seen in Figure ??. The BRCA1 gene expression level will act as the output value in our regression analysis and the other 17322 genes will serve as predictor values.

This data is a prime example of the p >> n problem where there are many more predictors than data samples.

3 Results

- 3.1 Regression Models
- 3.2 Classification Models
- 4 Discussion
- 4.1 Findings
- 4.2 Contributions
- 4.3 Future Work

		type independent	ent	symmetric					-	antoregressive	sive					blockwise					
		corr		corr					-	corr						corr					
		0		0.2		0.5 +os+ mso		0.9 +os+ mss	- +	0.2 tost mes	•	0.5		0.9	- +	0.2		0.5	- '	0.9 tost mes	
st.dev	model.name	Mean	SD	Mean	SD	Mean Mean	SD	Mean	SD	Mean	SD		SD		SD		SD	Mean	SD	Mean	SD
1	OLS AIC back	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251
	BIC back.	1.164	0.243	1.168	0.261	1.196	0.259	1.170	0.249	1.162	0.245	1.174	0.232	1.186	0.247	1.191	0.267	1.199	0.248	1.180	0.248
	AIC step. back.	1.218	0.249	1.207	0.239	1.231	0.262	1.207	0.254	1.216	0.235	1.209	0.237	1.238	0.256	1.221	0.256	1.223	0.249	1.216	0.243
	BIC step. back.	1.164	0.243	1.168	0.261	1.196	0.259	1.169	0.249	1.201	0.245	1.174	0.232	1.248	0.248	1.191	0.267	1.199	0.248	1.180	0.248
	BIC for.	1.159	0.252	1.163	0.261	1.195	0.255	1.196	0.308	1.156	0.241	1.171	0.236	1.396	0.402	1.188	0.263	1.199	0.244	1.230	0.368
	AIC step. for.	1.209	0.247	1.197	0.243	1.231	0.263	1.196	0.246	1.201	0.233	1.202	0.243	1.244	0.321	1.223	0.260	1.221	0.248	1.225	0.292
	BIC step. for.	1.159	0.252	1.163	0.261	1.195	0.255	1.196	0.308	1.156	0.241	1.171	0.236	1.396	0.402	1.188	0.263	1.199	0.244	1.230	0.368
	Ridge	1.586	0.343	1.593	0.408	1.695	0.499	1.843	0.385	1.643	0.333	1.749	0.398	1.866	0.504	1.663	0.351	1.665	0.424	1.801	0.406
	Lasso E-net	1.000 2000 1.000 1.000	0.322	1.390	0.397	1.402	0.349	1.395	0.362	1.399	0.308	1.410	0.319	1.387	0.418	1.418	0.326	1.360	0.301	1.395	0.383
	Adap, ridge	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.276	0.251	1.275	0.251	1.276	0.251	1.276	0.251	1.276	0.251
	Adap. lasso	1.382	0.334	1.395	0.406	1.422	0.380	1.399	0.333	1.399	0.315	1.417	0.327	1.382	0.417	1.424	0.327	1.373	0.324	1.389	0.387
	Adap e-net	1.372	0.320	1.390	0.405	1.408	0.373	1.397	0.345	1.390	0.300	1.414	0.312	1.380	0.403	1.427	0.334	1.369	0.316	1.385	0.401
	SCAD	1.200	0.249	1.207	0.262	1.230	0.256	1.185	0.257	1.201	0.274	1.196	0.249	1.201	0.249	1.205	0.254	1.219	0.254	1.174	0.262
	GB	3.728	1.140	3.696	1.221	3.823	1.140	2.926	0.696	4.023	1.169	3.848	1.154	2.934	0.815	3.693	1.197	3.747	1.015	2.907	0.727
	RF	6.899	1.779	6.581	1.856	5.279	1.275	2.592	0.534	6.952	1.671	5.883	1.390	2.779	0.658	6.807	1.848	5.704	1.279	2.867	0.683
¢	SVM	5.770	1.710	5.321	1.816	4.345	1.686	2.991	1.473	5.567	1.619	5.084	1.606	3.257	1.327	5.438	1.818	4.451	1.332	3.194	1.313
n	OLS ATC back	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258	11.483	2.258
	BIC back.	10.472	2.190	10.576	2.501	10.877	2.114	10.648	2.065	10.450	2.087	10.562	2.092	10.762	2.189	10.737	2.573	10.916	2.390	10.713	2.142
	AIC step. back.	10.964	2.239	10.932	2.448	11.095	2.257	10.911	2.117	10.918	2.153	10.885	2.137	11.072	2.249	11.050	2.499	11.067	2.375	11.131	2.176
	BIC step. back.	10.472	2.190	10.576	2.501	10.877	2.114	10.649	2.066	10.450	2.087	10.568	2.090	10.762	2.189	10.737	2.573	10.916	2.390	10.713	2.142
	AIC for.	10.882	2.223	10.874	2.487	11.018	2.153	10.903	2.093	10.831	2.217	10.833	2.183	11.154	2.631	11.010	2.523	10.998	2.333	11.025	2.312
	AIC sten. for.	10.882	2.223	10.369	2.487	11.015	2.151	10.903	2.093	10.831	2.217	10.816	2.120	11.148	2.678	11.010	2.523	11.022	2.355	11.032	2.309
	BIC step. for.	10.430	2.272	10.569	2.438	10.822	2.084	10.674	2.187	10.434	2.102	10.537	2.125	12.520	3.690	10.759	2.600	10.812	2.296	11.144	2.978
	Ridge	14.277	3.091	15.172	4.146	14.822	3.577	16.744	3.571	14.733	2.883	15.737	3.583	16.693	4.239	14.737	4.027	15.851	3.455	16.429	4.332
	Lasso Fract	12.491	2.899	12.648	3.262	12.484	2.851	12.557	3.285	12.500	2.852	12.693	2.869	12.366	3.500	12.618	3.418	12.834	2.919	12.849	3.744
	Adap, ridge	11.482	2.257	11.483	2.258	11.482	2.257	11.482	2.256	11.482	2.257	11.483	2.257	11.477	2.256	11.482	2.258	11.482	2.257	11.481	2.257
	Adap. lasso	12.442	3.003	12.793	3.416	12.469	2.929	12.659	3.286	12.592	2.977	12.753	2.945	12.457	3.749	12.743	3.599	12.906	2.843	12.787	3.655
	Adap e-net	12.344	2.884	12.689	3.276	12.423	2.875	12.742	3.309	12.578	2.802	12.728	2.804	12.487	3.549	12.708	3.515	12.822	2.821	12.883	3.698
	MCP	10.804	2.240	10.827	2.485	10.982	2.163	10.872	2.215	10.680	2.301	10.838	2.245	10.848	2.297	10.804	2.313	10.912	2.156	10.713	2.255
	GB	35.185	12.026	35.724	10.740	34.796	11.994	25.465	6.593	34.832	10.704	33.856	8.918	26.327	7.057	34.641	10.940	34.518	10.047	26.503	7.561
	RF	61.937	15.929	58.397	14.389	47.302	11.784	23.032	4.826	61.755	15.868	52.935	12.555	24.747	6.186	58.857	15.627	52.980	13.097	25.289	6.085
9	OLS	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030	45.930	9.030
	AIC back.	43.855	8.957	43.726	9.793	44.453	8.971	43.642	8.470	43.673	8.613	43.511	8.525	44.354	9.128	44.198	9.994	44.186	9.546	44.524	8.705
	AIC step back		8.759	42.304	0.004	43.509	8.456 9.026	42.592	8.239	41.802	8.547	42.247	8.307	43.046	007.00	42.950	0 994	43.003	9.500	42.854	8.570
	BIC step. back.		8.759	42.304	10.004	43.509	8.456	42.596	8.265	41.802	8.347	42.272	8.359	43.046	8.756	42.950	10.291	43.663	9.560	42.854	8.570
	AIC for.		8.892	43.497	9.947	44.071	8.611	43.611	8.374	43.325	8.867	43.334	8.731	44.618	10.525	44.039	10.092	43.992	9.332	44.101	9.247
	BIC for.	41.719	9.087	42.276	9.754	43.288	8.336	42.638	8.640	41.735	8.409	42.168	8.481	50.079	14.758	43.036	10.400	43.298	9.251	44.588	11.894
	BIC step. for.	41.719	9.087	42.276	9.754	43.288	8.336	42.697	8.747	41.735	8.409	42.147	8.498	50.079	14.761	43.036	10.400	43.246	9.183	44.576	11.910
	Ridge	57.109	12.364	60.687	16.583	59.289	14.307	66.974	14.284	58.930	11.533	62.949	14.330	66.771	16.957	58.947	16.108	63.405	13.819	65.715	17.329
	Lasso Frot	49.962	11.595	50.593	13.047	49.936	11.403	50.230	13.138	50.002	11.410	50.772	11.476	50.464	13.999	50.472	13.673	51.335	11.675	51.397	14.976
	Adap, ridge	45.929	9.029	45.931	9.031	45.927	9.028	45.928	9.024	45.929	9.029	45.931	9.030	45.910	9.026	45.930	9.030	45.929	9.029	45.924	9.027
	Adap. lasso	49.766	12.013	51.171	13.663	49.877	11.717	50.634	13.144	50.368	11.908	51.011	11.779	49.827	14.997	50.971	14.396	51.624	11.372	51.147	14.622
	Adap e-net	49.376	11.536	50.756	13.102	49.693	11.501	50.969	13.234	50.312	11.208	50.913	11.215	49.949	14.195	50.830	14.061	51.289	11.285	51.532	14.793
	SCAD	43.214	8.769	43.307	9.860	44.382	8.653	43.487	9.365	42.873	9.203	43.071	8.978	42.996	9.187	43.217	9.251	43.650	8.623	42.850	8.998
	GB	135.812	40.816	140.921	40.573	135.510	43.275	102.763	23.849	137.560		137.808	40.068	105.821	29.447	141.906	45.697	137.212	37.811	105.834	28.599
	RF	248.001	64.295	233.826	57.679	188.905	46.879	92.071	٠	246.706	٠. ·	211.647	50.441	99.022	01.0	235.415	62.415	211.856	52.405	101.307	24.452
	SVM	207.957	61.720	196.815	61.630	159.388	56.783	106.807	53.412	200.958	60.842	182.386	56.868	116.606	48.592	188.419	60.121	166.756	57.132	18.048	50.595

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