Chapter 6: Linear Model Selection & Regularization

In the regression setting, the standard linear model is commonly used to describe the relationship between a response Y and a set of variables X_1, \ldots, X_p .

The linear model has distinct advantages in terms of inference and is often surprisingly competitive for prediction. How can it be improved?

We can yield both better *prediction accuracy* and *model interpretability*:

prediction accuracy: if the true relationship is ~ linear, least squares will have low bias
if
$$n >> p \Rightarrow$$
 also have low voriance \Rightarrow perform well on test data.
But if n is not much larger than $p \Rightarrow$ high variability \Rightarrow poor performance.
If $p > n$: no longer a unique solution \Rightarrow variance \Rightarrow cannot be used at all!
god: reduce variance without adding too much bias.

model interpretability: often many variables used in a regression are not in fact associated
with the response.
By removing them (setting
$$\hat{\beta}_i = 0$$
) we could obtain a more easily interpretable model.
Note: least squares will hardly ever result in $\hat{\beta}_i = 0$.
 \sum need variable selection.

Same ideas apply to logistic regression.

1 Subset Selection

We consider methods for selecting subsets of predictors.

1.1 Best Subset Selection.

To perform *best subset selection*, we fit a separate least squares regression for each possible combination of the p predictors. ~ (12) = p(p-1) models w/ exactly 1 predictor, etc.

Algorithm:

1. lit Mo denite null model - no predictors. 2. For K=11-1P (a) Fit all (PK) models that contruct & predictors. (b) Pide the best of Those (MK). Best is defined by JRSS (TR2). 3. Solut a single best model from Mo,..., Mp using CV error, Cp, AIC/BIC, or adjusted R² Why can't we use R^2 for step 3? as p^{\uparrow} , $R^2 \uparrow$ always. We can perform something similar with logistic regression.

P=10 => 1000 middles!

1.2 Stepwise Selection

For computational reasons, best subset selection cannot be performed for very large p. \neg impossible with $p \ge 40$. Dest subsit may also suffer When plarge because v/ a large search space be can find spuribus good models that work on training data but perform poerly w/test data.

<u>Stepwise selection is a computationally efficient procedure that considers a much smaller</u> subset of models.

Forward Stepwise Selection:

start w/ no predictors and add predictors one at a fine until all predictors are in the madel. choose The "best" from these. 1. Let Mo denote the well model - no predictors, 2. For k= 0, --, p-1 (a) Consider all p-k models that angreat the predictors in M_K with I additional predictors.
(b) Choose The best among those p-k and call it M_{KH} (*TR*², *LRSS*). 3. Select a single best model from Mo, ..., Mp using CV error, Cp, ALC/BIC, adj R². -Now we fit It $\Sigma_{K=0}^{p-1}(p-k) = 1 + \frac{p(p+1)}{2}$ moduls.

4)

Backward Stepwise Selection: Begin W/ full model and take predictors away one at a time until we get to null model. 1. Let Mp denote the full modul - contains all p predictors. 2. For K= p,p-y-1: (a) consider all kmodels that contain all predictors except one of predictors in ME (K-1 predictors). (6) choose bust among new and call it Mpc1 (7R², #RSS). 3. Scleet the single best model from Mo, -, Mp using CV error, Cp, AIC/BIC, adj R? * Neither forward nor backwards stepwise selection are guaranteed to find the best model containing a subset of the p predictors. for ward selection can be used when p=n (but only up to n-1 predictors, not p!). **1.3** Choosing the Optimal Model Need away way to fick "best" model that depends on test error (training error net a good be either estimate estimate of This) Les either asstrate mis directly or a djust traching errors for model size. $7C_p = \frac{1}{n} \left(RSS + \frac{1}{2} d \hat{6}^2 \right)$ estimate of variance four fill model. papetind # of predidors in subst model add penalty to training error (RSS) to adjust for underestinate of test error. as $d\uparrow$, Cp1 (droose model of lowest value). AIC & BIC (cr use for maximum likelihood E+s $\rightarrow AIC = \frac{1}{nG^2} (RSS + 2dG^2)$ $BIC = \frac{1}{h^{22}} \left(RSS + \log(h) d^{22} \right)$ choose model w/ low value, fince $\log(n) \ge 2$ for $n \ge 7 = 7$ heavier penalty on models by many variables sted R^2 (least squeres moduls) $\ge 7 esults$ in smaller moduls. Adjusted R^2 (least squares moduls) $A^2 = 1 - \frac{RSS}{TSS}$ always 1 as d1 $Adj R^{2} = 1 - \frac{Rss/(n-d-1)}{Tss/(n-1)}$ thouse model w/ highest a of R². Validation and Cross-Validation - Directly estimate test error of validation or CV and choose model w/ lowest est. - Verg general, can be used w/ any modul, even when it's not dear how may eror. upredictors " we have. Now have fast computers => prese are preferred.

2 Shrinkage Methods

The subset selection methods involve using least squares to fit a linear model that contains a subset of the predictors. As an alternative, we can fit a model with all p predictors using a technique that constrains (*regularizes*) the estimates.

Shrinking the coefficient estimates can significantly reduce their variance!

2.1 Ridge Regression

Recall that the least squares fitting procedure estimates β_1, \ldots, β_p using values that minimize

Ridge Regression is similar to least squares, except that the coefficients are estimated by minimizing

The tuning parameter λ serves to control the impact on the regression parameters.

2.1 Ridge Regression

The standard least squares coefficient estimates are scale invariant.

In contrast, the ridge regression coefficients $\hat{\beta}_{\lambda}^{R}$ can change substantially when multiplying a given predictor by a constant.

Therefore, it is best to apply ridge regression *after standardizing the predictors* so that they are on the same scale:

Why does ridge regression work?

2.2 The Lasso

Ridge regression does have one obvious disadvantage.

This may not be a problem for prediction accuracy, but it could be a challenge for model interpretation when p is very large.

The *lasso* is an alternative that overcomes this disadvantage. The lasso coefficients $\hat{\beta}_{\lambda}^{L}$ minimize

As with ridge regression, the lasso shrinks the coefficient estimates towards zero.

As a result, lasso models are generally easier to interpret.

Why does the lasso result in estimates that are exactly equal to zero but ridge regression does not? One can show that the lasso and ridge regression coefficient estimates solve the following problems

In other words, when we perform the lasso we are trying to find the set of coefficient estimates that lead to the smalled RSS, subject to the contraint that there is a budget s for how large $\sum_{j=1}^{p} |\beta_j|$ can be.

2.3 Tuning

We still need a mechanism by which we can determine which of the models under consideration is "best".

For both the lasso and ridge regression, we need to select λ (or the budget s).

How?

3 Dimension Reduction Methods

So far we have controlled variance in two ways:

We now explore a class of approaches that

We refer to these techniques as *dimension reduction* methods.

The term dimension reduction comes from the fact that this approach reduces the problem of estimating p + 1 coefficients to the problem of estimating M + 1 coefficients where M < p.

Dimension reduction serves to constrain β_j , since now they must take a particular form.

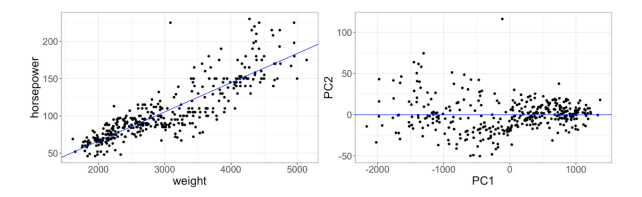
All dimension reduction methods work in two steps.

3.1 Principle Component Regression

Principal Components Analysis (PCA) is a popular approach for deriving a lowdimensional set of features from a large set of variables.

The *first principal component* directions of the data is that along which the obervations vary the most.

We can construct up to p principal components, where the 2nd principal component is a linear combination of the variables that are uncorrelated to the first principal component and has the largest variance subject to this constraint.



3.1 Principle Component Regres...

The Principal Components Regression approach (PCR) involves

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Key idea:

In other words, we assume that the directions in which X_1, \ldots, X_p show the most variation are the directions that are associated with Y.

How to choose M, the number of components?

Note: PCR is not feature selection!

3.2 Partial Least Squares

The PCR approach involved identifying linear combinations that best represent the predictors X_1, \ldots, X_p .

Consequently, PCR suffers from a drawback

Alternatively, partial least squares (PLS) is a supervised version.

Roughly speaking, the PLS approach attempts to find directions that help explain both the reponse and the predictors.

The first PLS direction is computed,

To identify the second PLS direction,

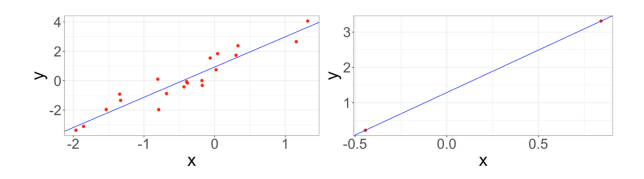
As with PCR, the number of partial least squares directions is chosen as a tuning parameter.

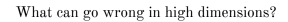
4 Considerations in High Dimensions

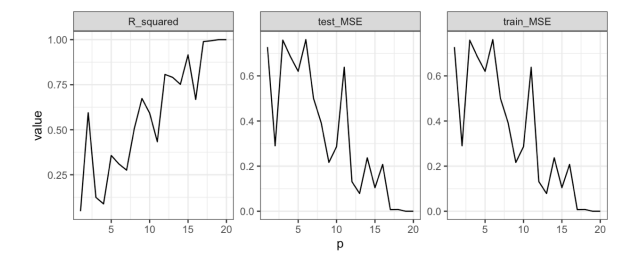
Most traditional statistical techniques for regression and classification are intendend for the low-dimensional setting.

In the past 25 years, new technologies have changed the way that data are collected in many fields. It is not commonplace to collect an almost unlimited number of feature measurements.

Data sets containing more features than observations are often referred to as *high-dimensional*.







Many of the methods that we've seen for fitting *less flexible* models work well in the high-dimension setting.

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When we perform the lasso, ridge regression, or other regression procedures in the highdimensional setting, we must be careful how we report our results.