3 Dimension Reduction Methods

So far we have controlled variance in two ways:

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

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(1) Let
$$Z_{1,1-1} Z_{M}$$
 represent $M < \rho$ linear combinations of our original predictors
 $Z_{m} = \sum_{j=1}^{P} \phi_{jm} X_{j}$
for constants $\phi_{1m,1-1}, \phi_{pm}$ $m = 1, ..., M$
(2) Fit the linear regression model using least squares
 $\gamma_{i} = \Theta_{0} + \sum_{m=1}^{M} \Theta_{m} Z_{im} + \Sigma_{i}$ $i = k ..., n$
 $\gamma_{i} = \sigma_{0} + \sum_{m=1}^{M} \Theta_{m} Z_{im} + \Sigma_{i}$ $i = k ..., n$

2.3 Tuning

The term <u>dimension reduction</u> 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.

$$\frac{1}{1} \langle p, \frac{1}{1} \rangle = \frac{1}{1} \left[\sum_{m=1}^{N} \theta_m \sum_{m=1}^{N} \theta_m \sum_{m=1}^{P} \theta_m \sum_{j=1}^{P} \theta_{jm} \sum_{j=1}^{P} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{P} \theta_{jm} \sum_{j=1}^{N} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{P} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{P} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{P} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{P} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{N} \theta_m \sum_{j=1}^{N} \theta_m \sum_{j=1}^{N} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{N} \theta_m \sum_{m=1}^{N} \theta_m \sum_{j=1}^{N} \theta_m \sum_$$

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

$$B_{j} = \sum_{m=1}^{M} \Theta_{m} \phi_{im}$$

$$= \sum_{m=1$$

All dimension reduction methods work in two steps.

() transformed predictors are obtained (get
$$\{ g \}_{jm} \}_{j=1,..,p}$$
)
Lime
(a) Model is fit using M transformed predictors from (1) -
We done in multiple trays.
We will talk about 2.

ONE UNY Z()--)ZM

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3.1 Principle Component Regression

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

PCA is an unsupervised approach for reducing the dimension of an map data The first principal component directions of the data is that along which the obervations X. st principal direction vary the most. The 1st principal components are obtained by projectily the data onto the 1st principal Tri Component direction. a point is projected onto a line by finding the point \mathbf{X}_{1} on the line closest to the original point. out of every possible linear combination of X_1 and X_2 such that $\phi_{11}^2 + \phi_{21}^2 = 1$, choose ϕ_{11} s.t. point × projected onto the line. Var $(\phi_{11}(x,-\overline{x}_1) + \phi_{21}(x_2-\overline{x}_2))^{is}$ maximized. $\mathbf{z}_{i} = \mathbf{p}_{i} (\mathbf{x}_{i} - \mathbf{\bar{x}}_{i}) +$ We can construct up to p principal components, where the 2nd principal component is a $\phi_{2}(z_{1}-\overline{z})$ linear combination of the variables that are uncorrelated to the first principal component and has the largest variance subject to this constraint. => 2nd PC direction is for isla-in perpendicular to 1st PC direction. 2 PC direction are called 1st PC direction. Iprincipal component scores 2100 17 200 1St PC ł horsepower 50 PC2 150 direction dimension along which data 100 te tary most -50-4000 -2000 2000 5000 -1000 1000 3000 weight Zi, i=1,...,n projected onto principal component directions. 1 St PC contains the most information -> ptm pC contains the least.

The Principal Components Regression approach (PCR) involves

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!

NOTE: recommended standardizing predictors X1,..., Xp to each have st. dev. = I before gutting the PCs.

3.2 Partial Least Squares

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

dire ctims

We identified this directions is an unsupervised US? (response ? is not and to be productions of the production of the directions is a guarantee that the direction that be st explained the predictors will also be the best direction is the explain the relaxington that be st explained the predictors will also be the best direction is the explained the relaxington of directions in reduction.
(1) identify new fail least squares (PLS) is a supervised version. of dimension reduction (1) identify new fail vers
$$Z_{12-2} \geq m$$
 liner combination of original predictors (2) is a supervised version. of dimension reduction (1) identify new fail the squares (PLS) is a supervised version of dimension is directions (2) it is a product square predictors.
(2) it is nodel (least squares) using transformed predictors.
(3) it liner model (least squares) using transformed predictors.
(4) the predictors of the predictors of the predictors of the specific predictors.
(2) it is a grant to use y (not just x) to find directions that help explain both the reports and the predictors.
The first PLS direction is computed,
(1) standardize the p predictors (still have st. dow = 2).
(3) set each β_{11} equal to the coefficient from a simple liner regression $\gamma \sim \chi_1$
Since the coefficient from SLR of $\gamma \sim \chi_1$ of $Cor (Y, \chi_1)$ PLS places highest weight be a visibles that are strangly related (linerly) to the regionse.
To identify the second PLS direction, $\chi_1 \sim Z_1$
(1) regress each waveles χ_{12-2} χ_2 on Z_1 and get residuals ($G_1 = \chi_{11} - \chi_{11} - \chi_{11} - \chi_{12} - \chi_{13} - \chi_$

In practice, PLS usually performs no better than ridge or PCR.

L> supervised nature of problem does reduce, but often increases variance. => not always better.

4 Considerations in High Dimensions

Most traditional statistical techniques for regression and classification are intendend for the low-dimensional setting. $n >> \rho$

This is because throughout the history of the field, the bulk of scientific problems requiring statistics have been low dimensional.

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. (p very lenge).

But n can still be limited due to cost, scappling availability, etc.

But we may only only have a few thousand people who have commented to use that history-

for a given use the fonteres would be absense (D) or presence (I) of each population starth term. $\Rightarrow p | large but n \approx 2000$ Data sets containing more features than observations are often referred to as <u>high-dimensional</u>. $p \geq n$

classical approaches (like least squares) are not appropriate in this outhing.
(why? think bias-variance trade-off and overfitting).
=> we need to be careful when
$$n \approx \rho$$
 or $n < \rho$.
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What can go wrong in high dimensions? Going the table about least squares but same issues arise v/ logistic regression or 2DA

If prim or pon regardless of if there is a relationship W/ response pont. least squeres will yield a bet of wefficients that is an (almost if n =>residuals =0.



Simulated data w/ N=20 and regression performed with seturen 1 and 20 testures. features generated m/ no relationship to response



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

1. 2.

3.

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