STATISTICS 452/652: Statistical Learning and Prediction

Lecture 7: Dimension Reduction

(Reading: ISLR 6.3)

1 Goals of lecture

- In order to reduce variance in prediction it helps to reduce the number of parameters to be estimated
- Sometimes explanatory variables are highly correlated
 - It is difficult to know which variable(s) to select from a group
 - Keeping them all adds variability needlessly when not all of them are needed
- It would help to have a way to "boil down" correlated variables to a smaller number measures that contain the main information within them.
 - E.G., maybe there are 10 variables all measuring some aspect of "size" of a person, and some notion of size is needed in a model, but not in 10 different versions.
- Can we use the data to engineer these new features for us automatically?

2 Principal Components

- Sometimes p is too large to work with (e.g. p > n)
- Even when it's not, it might be much larger than the dimensionality of the actual information contained with in the X-space.
 - For example, when X_1 and X_2 are strongly correlated, the information in (X_1, X_2) lies "almost" in a 1-dimensional space
 - See Fig 1.
- More complex relations may cause X to be nearly p_1 -dimensional, for $p_1 < p$.
- It would be nice to be able to reduce the dimension of X before using it for other things.

Figure 1: Correlated variables in p=2 dimensions can almost be represented by a single variable along the straight green line. (From ISLR)

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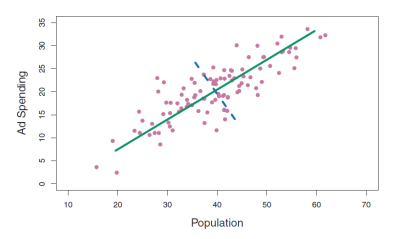


FIGURE 6.14. The population size (pop) and ad spending (ad) for 100 different cities are shown as purple circles. The green solid line indicates the first principal component, and the blue dashed line indicates the second principal component.

2.1 Principal Components Analysis (PCA)

- 1. Usually start by standardizing each X_j to have mean 0 and SD 1, but not strictly necessary
- 2. Find the direction within X that explains the most variability in X.
 - (a) "Project" points onto different lines through the middle of the data
 - (b) Compute variance of points on each line
 - (c) Choose the line where the variance is highest
 - i. This is the first principal component (PC1)
 - (d) Project all data onto this line (See Fig 2).
 - (e) The formula for this line is $Z_1 = \sum_{j=1}^p X_j \phi_{1_j}$, where the values $\phi_{11}, \ldots, \phi_{1p}$ are found from the data¹
 - (f) Each observation has a value of Z_1 that represents its position on a new coordinate axis.
 - i. Projecting p-dimensional points orthogonally onto this axis
 - (g) "Total variance" of data is sum on variances for each X_i
 - i. Variance of Z_1 points is larger than variance of any single X_j , but less than the total variance

¹Principal components are found using eigen decomposition, in case you were curious.

Figure 2: Principal components in p=2 dimensions. (From ISLR) 6. Linear Model Selection and Regularization

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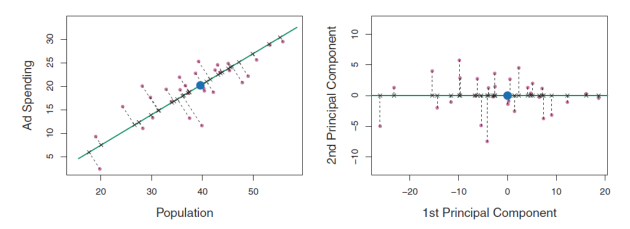


FIGURE 6.15. A subset of the advertising data. The mean pop and ad budgets are indicated with a blue circle. Left: The first principal component direction is shown in green. It is the dimension along which the data vary the most, and it also defines the line that is closest to all n of the observations. The distances from each observation to the principal component are represented using the black dashed line segments. The blue dot represents (pop, ad). Right: The left-hand panel has been rotated so that the first principal component direction coincides with the x-axis.

- 3. Next look for the direction orthogonal (perpendicular) to the first that explains the next-most variability in X
 - (a) Direction is $Z_2 = \sum_{j=1}^p X_j \phi_{2j}$ for a new set of values $\phi_{21}, \dots, \phi_{2p}$
 - (b) Z_1 and Z_2 are uncorrelated
 - (c) A second coordinate axis
 - i. Projecting points onto this plane
 - (d) Variance of points around Z_2 is less than variance around Z_1
 - (e) Sum of these is smaller than or equal to total variance.
- 4. Keep repeating this until there is no more variability (or until you reach p components if there are p variables and they are not perfectly correlated.)
 - (a) Have Z_1, Z_2, \dots, Z_p new variables engineered from X
 - (b) These are called the PRINCIPAL COMPONENTS (PC's) of X.
- 5. Sum of variances of Z's equals the total variance
 - (a) It just splits it into new linear variables that each explain as much as they can in sequence
- 6. The hope is that just a few (M < p) PCs explain most of the variability in X, so you just use those PCs for further use.
 - (a) In Fig. 2, I might choose just to keep the first component and drop the second.
 - (b) Projecting the points onto the "reduced-dimensional subspace" loses information, but loss is minimized given M.
- 7. What we are really doing is rotating the axes to more directly align with the contents of the data. then dropping some axes $(Z_j$'s)

2.2 Choosing M

You can do this before the regression or after. See later for the "after" approach.

- Suppose we let T be the "total variance", the sum of the variances from each X_j , $j = 1, \ldots, p$
- Suppose we let v_j be the variance of the PC Z_j , j = 1, ..., p
- Then we know that $\sum_{j=1}^{p} v_j = T$
- The whole point of a PCA is hoping that you can find an M < p so that $\sum_{j=1}^{M} v_j \approx \sum_{j=1}^{p} v_j$
- So plot $\sum_{j=1}^{m} v_j$ vs. m for different values of m (cumulative variance plot)

- Increases rapidly, then levels off
- Choose M to be a value of m where the increase is no longer rapid and the leveling off has begun.
- Will see this in an example
- Or can plot v_j vs. j (SCREE PLOT) to see if there is a clear point at which v_j 's level off
- If you choose to standardize data so that each X_j has SD=1, interpretation is a little easier
 - Total variance $T = \sum_{j=1}^{p} v_j = p$, so average v_j is 1
 - Another possible way to select M is to choose all j with $v_i > 1$

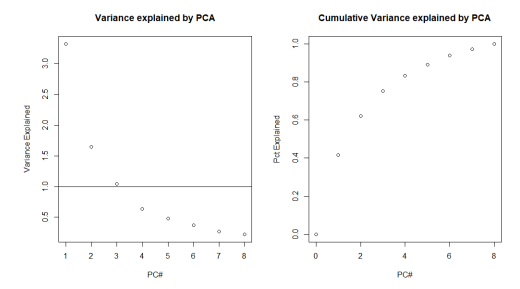
Example: Finding principal components on Prostate Data (L7 - Principal Comp and Partial LS.R) I use the full prostate data with 8 explanatory variables to perform PCA using the prcomp() function from the main stats package. We produce 8 PC's, Z_1, \ldots, Z_8 and see (1) how much of the total variance each one explains, and (2) what the coefficients look like. Adding scale.=TRUE does the standardizing. The output shows the Standard deviation $(\sqrt{v_j})$, the Proportion of Variance (v_j/T) , and the Cumulative Proportion $(\sum_{l=1}^j v_l/T)$ for PC Z_j , $j=1,\ldots,p$

The output shows

```
> pc <- prcomp(x=prostate[,2:9], scale.=TRUE)</pre>
> summary(pc)
Importance of components:
                           PC1
                                   PC2
                                          PC3
                                                  PC4
PC5
Standard deviation
                        1.8234 1.2846 1.0219 0.79990 0.69177
Proportion of Variance 0.4156 0.2063 0.1305 0.07998 0.05982
                        0.4156 0.6219 0.7524 0.83241 0.89223
Cumulative Proportion
                            PC6
                                     PC7
                                             PC8
Standard deviation
                        0.60959 0.51723 0.47227
Proportion of Variance 0.04645 0.03344 0.02788
Cumulative Proportion
                        0.93868 0.97212 1.00000
```

I can also make a scree plot or could make a cumulative variance plot. The scree plot is available from the plot() function directly on the prcomp object, but I have created a little nicer one (I think) with a tiny bit of programming. See Figure 3. It is usually hard to draw a line between "keep" and "drop", and this is no exception. The curve in the plots changes slope kind of slowly instead of suddenly. The simple $v_j > 1$ rule says to take M = 3. To me, the PCs from 5–8 seem not to add much variance, so I might choose 4 or 3 PCs. There is no "right" answer.

Figure 3: Scree plots for Prostate data. Left: the default produced by plot.prcomp; right: minet



Finally the program does print out the actual ϕ weights, but they don't really tell me anything interesting, so I don't show them here. Sometimes, you can see interesting relationships in the data from them.

3 Principal Components Regression

- Principal components regression (PCR) is just...(wait for it)...regression on the principal components!
- Instead of fitting $f_X(X) = \beta_0 + \beta_1 X_1 + \ldots + \beta_p X_p$ with p+1 parameters, fit

$$f_{ZM}(Z) = \theta_0 + \theta_1 Z_1 + \ldots + \theta_M Z_M$$

which has just M+1 regression parameters.

Details

- Once the PCs Z_1, \ldots, Z_p are created, and the choice of M < p is made, there is nothing new.
 - It is just linear regression on the engineered features instead of the original variables.
- Because each Z is made up of different amounts of all X_j 's, this is not a variable selection process
 - Every original X_j contributes to the predicted values, but in different ways

- Can function sort of like shrinkage
- Factoid: If you let M=p (i.e., use all of the PCs as long as n>p) you get back exactly the same predicted values with Z_1, \ldots, Z_p as you would from a linear regression on X_1, \ldots, X_p

3.1 Treating M as a tuning parameter

- Previous selection of M in Section 2.2 used only information about variance within X
 - Process does not use errors in Y to balance bias and variance
 - It is merely trying to restructure information in X to retain "as much of X as possible" while reducing the number of parameters we will later ask regression to estimate.
 - It is totally based on judgment and guesswork.
- Alternatively, you could defer choice of M and let Y help you decide later
- Specifically, you can think of PCR is a family of regression models
 - Kind of like polynomials, you can try fitting regressions with $M=1,2,3,\ldots,p$
 - Then choose the optimal value of M using CV or some other measure of validation error.
- This method uses Y, and is subject to usual concerns about selecting one model from many

4 Partial Least Squares

- PCR suffers from the fact that the newly engineered features are made without knowledge of Y
 - Just because a PC explains the largest amount of variance within X doesn't mean that that particular combination explains much variability in Y
 - Even if you treat M as a tuning parameter, you still don't use Y to influence the ϕ weights in the PC's
 - * They are entirely determined by the correlation structure among the X_j 's
- Partial Least Squares (PLS) similarly creates weighted combinations of the X_j 's, but it lets the responses choose the ϕ weights
- The algorithm is more complicated than I want to explain, but the essence is this:
 - 1. Start by computing the correlation between Y and each X_i

2. Use rescaled versions of the correlations them as the ϕ weights for the first component:

$$Z_1^{PLS} = \phi_{11}^{PLS} X_1 + \phi_{12}^{PLS} X_2 + \ldots + \phi_{1p}^{PLS} X_p$$

- (a) The stronger the correlation is between X_j and Y compared to other variables, the more it counts toward Z_1^{PLS}
- 3. Repeat the process on residuals from regressions of Y vs. Z_1^{PLS} and each X_j vs. Z_1^{PLS} to get a new set of correlations and new weights
 - (a) These create Z_2^{PLS}
- 4. Repeat for a total of M times for some chosen M
- 5. Do multiple linear regression of $Y = \theta_0 + \theta_1 Z_1^{PLS} + \ldots + \theta_M Z_M^{PLS}$
- Can treat M as a tuning parameter and choose it using CV
 - In both PLS and PCR, must perform *entire process* (creating components, performing regressions for different M) on training set
 - Produce a set of predictions on validation set for each regression model
 - DO NOT use full data to create components, and then run only the regression part on training set.
- PLS has similar properties to PCR, but may need smaller M because components do
 use information about Y
 - For this reason, PLS is probably the better tool to learn
 - Performance reportedly tends to be similar
 - In both cases, you can rearrange the coefficients algebraically to rewrite the prediction function as a linear regression in terms of all 8 variables, but with coefficients that are different from LS or Ridge.
 - * If you really want to. I usually don't...

Example: Partial Least Squares on Prostate Data (L7 - Principal Comp and Partial LS.R) This is really not the right data set for showing off what PLS can do. PLS is really meant for cases where p is larger than we are willing to make a model out of, so we really wan to reduce the dimension somehow. In this case, p = 8, so it's kind of a dull demonstration. I'll do it anyway, rather than introduce more data sets.

The pls package does both PLS and PCR regression. I use the plsr() function for PLS regression. The code is pretty similar to lm() except that I can tell it

- 1. The maximum number of components I want it to consider, which is useful when p is very large, and
- 2. The option to use validation to choose number of components adding the validation= argument
 - (a) Can use "none", "CV", or "LOO" (leave one out CV).

- (b) If "CV" is used, V=10 unless changed with segments=
- 3. LOTS more options and helper functions to better understand the model results, which I am not specifically interested in at the moment.

Feel free to explore the package vignette at https://cran.r-project.org/web/packages/pls/vignettes/pls-manual.pdf

In the output below, the validation error is the root-MSPE, and there are lines for the CV error as we learned it and for a "bias-adjusted" version that I haven't read about. I am fine with the regular version. Under TRAINING, the lpsa rows show the model R^2 fitting 1,2,...8 PLS components. I am not sure what the X rows mean.

```
> library(pls)
> mod.pls = plsr(lpsa~., data=prostate, ncomp=8, validation="CV")
> summary(mod.pls)
        X dimension: 97 8
Data:
        Y dimension: 97 1
Fit method: kernelpls
Number of components considered: 8
VALIDATION: RMSEP
Cross-validated using 10 random segments.
       (Intercept)
                      1 comps
                                2 comps
                                         3 comps
                                                   4 comps
CV
               1.16
                         1.05
                                  1.002
                                           0.8206
                                                    0.7805
               1.16
                         1.05
                                  1.038
                                           0.8171
                                                    0.7767
adjCV
                 6 comps
                           7 comps
       5 comps
                                     8 comps
                  0.7510
CV
        0.7599
                            0.7471
                                      0.7462
        0.7573
                  0.7465
                            0.7428
                                      0.7420
adjCV
TRAINING: % variance explained
                2 comps
                          3 comps
                                              5 comps
                                                        6 comps
      1 comps
                                    4 comps
Χ
        93.48
                  98.52
                            99.67
                                      99.76
                                                99.94
                                                          99.97
lpsa
         18.19
                  26.68
                            56.31
                                      62.15
                                                63.65
                                                          66.06
      7 comps
                8 comps
Χ
         99.99
                 100.00
         66.31
                  66.34
lpsa
```

Strangely, CV has selected a model with all 8 components, although the root-MSPE's are not changing much after about 4. This is clear in the R^2 values as well—there is little gain after about 4 components. I would be tempted to use just 4.

5 What to take away from this

- Dimension reduction uses feature engineering on ALL X_j to create new variables that are linear combinations of the originals
- Run regression on small number of these components
 - Considers information from each X_j
 - Uses fewer parameters than running p-dimensional regression
- Drawback is that there is no guarantee that the components capture all relevant information about complicated relationships between Y and the X_j 's
- Also no reduction in the number of variables that must be measured to create the predictions

6 Exercises

Application

Motor Insurance Data:

Refer to the motor insurance data that we studied in the exercises in Lecture 4. Recall that there are two factors in the data.

- 1. Convert the two factors, make and zone into full sets of indicator variables, add them to the data frame, and drop the factors. An easy way to do this is using the dummy.vars() function from the package caret. For example, my code looks like this:
 - > library (caret) > ins.dv = data.frame(predict(dummyVars("~.", data=ins), newdata=ins)) > head(ins.dv) zone.2 zone.3 zone.4 zone.5 per km zone.1 11.649460 9.615939 8.028129 11.22264011.290310 10.163730 zone.7 bonus make.1 make.2make.3 make.4 make.5make.6make.7make.8 make.9 insured claims 669.34 55.70 81.78 200.68 64.73176.70
 - (a) To make sure that you have done this correctly, **report the dimensions of the new data frame.**
- 2. Run a principal components analysis on these data, excluding the response variable per, using scale.=TRUE to standardize the data.
 - (a) **How many PC's get chosen** if you use the guideline of choosing all PC's with higher than average contributions to the explained variance?
 - (b) Make a scree plot and a cumulative variance plot. Show these plots.

(c) Based on these plots, **indicate how many PC's** you think explain "enough" variability while achieving dimension reduction? **Explain very briefly why you chose that number.**

OZONE DATA

Refer Lecture 6, Problem 3, where you used 10-fold CV for comparing LASSO models

- 1. Using the same 10 folds estimate the MSPE for PLS.
 - (a) On each training set, run the PLS function, choosing the optimum number of components using 10-fold CV. You can figure out how many components is optimal according to CV using code like this, assuming that I have named my model, mod.pls

```
mp.cv = mod.pls$validation
Opt.Comps = which.min(sqrt(mp.cv$PRESS/nrow(ins.dv)))
```

Report the optimal number of components for each of the 10 folds.

- (b) Use the predict() function to predict responses on the test set. You need to specify ncomps for this function.
- (c) Report the separate MSPEs from each fold, $MSPE_v$, v = 1, ..., 10 and the MSPE for the full data.
- (d) ADD a boxplot for PLS to the boxplots you made for other models. Comment on how well PLS compares to the other models.
- (e) Remake the plot using relative MSPE and comment.