

Partial Least Squares Regression (part II)

Predictive Modeling & Statistical Learning

Gaston Sanchez

CC BY-SA 4.0

PLS Regression

PLS Regression equation in terms of the original predictors:

$$\begin{aligned} \mathbf{y} &= d_1 \mathbf{z}_1 + d_2 \mathbf{z}_2 + \mathbf{e} \\ &= d_1 \mathbf{X} \mathbf{w}_1 + d_2 \mathbf{X} \mathbf{w}_2 + \mathbf{e} \\ &= \mathbf{X}(d_1 w_1^* + d_2 w_2^*) + \mathbf{e} \\ &= b_1 \mathbf{x}_1 + b_2 \mathbf{x}_2 + \cdots + b_p \mathbf{x}_p + \mathbf{e} \end{aligned}$$

Properties of PLS Regression

Properties

- ▶ $\mathbf{z}_h^\top \mathbf{z}_l = 0, \quad l > h$
- ▶ $\mathbf{w}_h^\top \mathbf{p}_h = 1$
- ▶ $\mathbf{w}_h^\top \mathbf{X}_l^\top = 0, \quad l \geq h$
- ▶ $\mathbf{w}_h^\top \mathbf{p}_l = 0, \quad l > h$
- ▶ $\mathbf{w}_h^\top \mathbf{w}_l = 0, \quad l > h$
- ▶ $\mathbf{z}_h^\top \mathbf{X}_l = 0, \quad l \geq h$
- ▶ $\mathbf{X}_h = \mathbf{X} \prod_{j=1}^p (\mathbf{I} - \mathbf{w}_j \mathbf{p}_j^\top), \quad h \geq 1$

Modified Weights \mathbf{w}_h^*

We know that $\mathbf{z}_h = \mathbf{X}_{h-1} \mathbf{w}_h$

\mathbf{z}_h can also be expressed as $\mathbf{z}_h = \mathbf{X} \mathbf{w}_h^*$

$$\mathbf{w}_h^* = \prod_{k=1}^{h-1} (\mathbf{I} - \mathbf{w}_k \mathbf{p}_k^T) \mathbf{w}_h$$

Modified Weights \mathbf{w}_h^*

In fact,

$$\mathbf{W}_h^* = \mathbf{W}_h(\mathbf{P}_h^T \mathbf{W}_h)^{-1}$$

$$\mathbf{Z}_h = \mathbf{X} \mathbf{W}_h(\mathbf{P}_h^T \mathbf{W}_h)^{-1}$$

Decomposition

The matrices of PLS components \mathbf{Z} and loadings \mathbf{P} can be used to decompose \mathbf{X} as:

$$\mathbf{X} = \mathbf{Z}\mathbf{P}^\top$$

It can be shown that:

$$\hat{\boldsymbol{\beta}}_{OLS} = \sum_{h=1}^p d_h \mathbf{w}_h^*$$

$$\hat{\mathbf{y}}_{OLS} = d_1 \mathbf{z}_1 + d_2 \mathbf{z}_2 + \cdots + d_p \mathbf{z}_p$$

What is PLSR doing?

Why PLS is worth it?

- ▶ The answer is stability of predictors.
- ▶ PLS keeps the number of variables as low as possible.
- ▶ In PLS, components are selected that give maximal reduction in the covariance $\mathbf{X}^T \mathbf{y}$ of the data.
- ▶ In that sense PLS will give the minimum number of variables that is necessary.
- ▶ The PLS regression is based on the SVD of $\mathbf{X}^T \mathbf{y}$

Some Insights

The first PLS component has the form $\mathbf{z} = \mathbf{X}\mathbf{w}$

Under the hood, the PLS regression involves **Tucker** criterion:

$$\arg \max_{\mathbf{w}} \{cov^2(\mathbf{y}, \mathbf{X}\mathbf{w})\}$$

What is this criterion doing?

Some Insights

Recall that the covariance can be expressed as:

$$\text{cov}(\mathbf{y}, \mathbf{z}) = \text{cor}(\mathbf{y}, \mathbf{z}) \sqrt{\text{var}(\mathbf{y})} \sqrt{\text{var}(\mathbf{z})}$$

thus:

$$\text{cov}^2(\mathbf{y}, \mathbf{z}) = \text{cor}^2(\mathbf{y}, \mathbf{z}) \text{var}(\mathbf{y}) \text{var}(\mathbf{z})$$

Some Insights

What does PLSR optimize?

$$\arg \max_{\mathbf{w}} \{ \text{cov}^2(\mathbf{y}, \mathbf{X}\mathbf{w}) \}$$

is equivalent to:

$$\arg \max_{\mathbf{w}} \{ \text{cor}^2(\mathbf{y}, \mathbf{z}) \text{var}(\mathbf{y}) \text{var}(\mathbf{z}) \}$$

PLSR is a compromise between the multiple regression of \mathbf{y} on \mathbf{X} , and the PCA of \mathbf{X}

Some Insights

Tucker's criterion $cov^2(\mathbf{y}, \mathbf{X}\mathbf{w})$ is a compromise between:

- ▶ maximizing correlation $cor(\mathbf{z}, \mathbf{y})$ (OLS regression)
- ▶ maximizing variance of PLS components $var(\mathbf{X}\mathbf{w})$

Advantages

- ▶ PLSR is not based on any optimization criterion.
- ▶ Rather it is based on an iterative algorithm (which converges).
- ▶ However, it turns out that the PLS-solution is equivalent to the SVD of $\mathbf{X}^T \mathbf{y}$

Advantages of PLS Regression

Advantages

- ▶ Simplicity in its algorithm
- ▶ No need to invert any matrix
- ▶ No need to diagonalize any matrix
- ▶ You just need to compute simple regressions
- ▶ In other words, you just need inner products
- ▶ Missing data is allowed (but you need to modify the algorithm)
- ▶ Easily extendable to the multivariate case of various responses
- ▶ Handles cases where we have more predictors than observations ($p \gg n$)

Example: Gasoline Data

Gasoline Octane Ratings



https://commons.wikimedia.org/wiki/File:Gas_Station_Pump_Five_Octane_Ratings.jpg

Predicting Octane Number

- ▶ Predicting octane number of a gasoline from the NIR (Near Infra Red) spectrum of gasolines.
- ▶ The **octane number**, or octane rating, is a standard measure of the performance of an engine or aviation fuel.
- ▶ The higher the octane number, the more compression the fuel can withstand before detonating (igniting).
- ▶ Fuels with a higher octane rating are used in high performance gasoline engines that require higher compression ratios.

Research Octane Number (RON)



<http://www.waukeshacfr.com/f1-f2/>

The most common type of octane rating worldwide is the Research Octane Number (RON). RON is determined by running the fuel in a test engine with a variable compression ratio under controlled conditions, and comparing the results with those for mixtures of iso-octane and n-heptane.

Dataset gasoline.txt

- ▶ 60 gasolines, 402 variables
- ▶ Response Y : octane number
- ▶ Predictors X_1, \dots, X_{401} : NIR spectrum frequencies (900nm-1700nm)
- ▶ As you can tell: $p \gg n$
- ▶ We'll use the first 50 gasolines as the training set
- ▶ The remaining gasolines (last 10) will be used as test set

Dataset gasoline.txt

Data file gasoline.txt in the data/ folder of the github repo

```
gasoline <- read.table("gasoline.txt", header = TRUE)
```

```
dim(gasoline)
```

```
## [1] 60 402
```

Dataset gasoline.txt

First few rows of data:

	octane	NIR.900.nm	NIR.902.nm	NIR.904.nm	NIR.906.nm
1	85.30	-0.050193	-0.045903	-0.042187	-0.037177
2	85.25	-0.044227	-0.039602	-0.035673	-0.030911
3	88.45	-0.046867	-0.041260	-0.036979	-0.031458
4	83.40	-0.046705	-0.042240	-0.038561	-0.034513
5	87.90	-0.050859	-0.045145	-0.041025	-0.036357
6	85.50	-0.048094	-0.042739	-0.038812	-0.034017
7	88.90	-0.049906	-0.044558	-0.040543	-0.035716

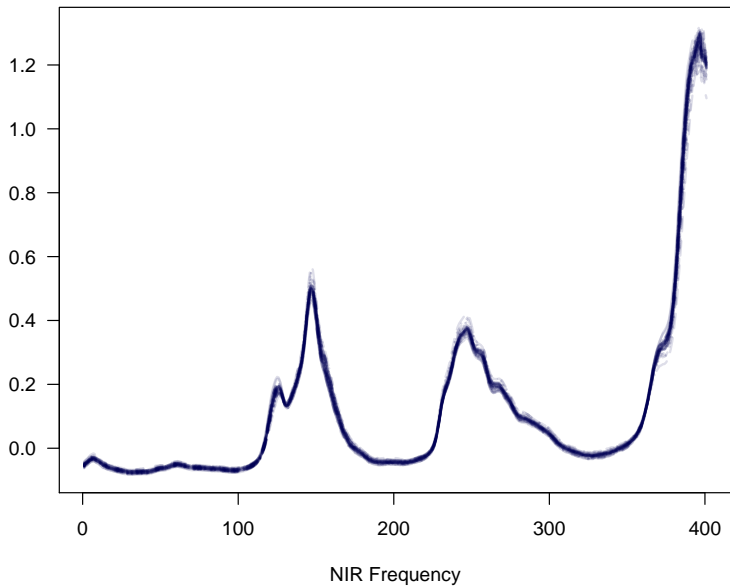
- ▶ First column octane is the response.
- ▶ Rest of columns are predictors.

PCA plot showing the distribution of NIR spectra data. The x-axis is labeled 'Dim 1 (71.57%)' and the y-axis is labeled 'Dim 2 (22.83%)'. A large cluster of points is on the right, and a smaller cluster is on the left. Vectors from the origin point to various NIR wavelengths, with labels like 'NIR.1696.nm' and 'NIR.1628.nm'.

Example: gasoline.txt

```
# response  
octane <- gasoline[,1]  
  
# predictors  
NIR <- gasoline[,2:ncol(gasoline)]  
  
# training and test sets  
train <- 1:50  
test  <- 51:60
```

NIR Spectrum



```
corrs <- cor(NIR, octane)
summary(corrs)
```

```
      V1
Min.   :-0.90362
1st Qu.: -0.38877
Median :-0.19437
Mean   :-0.18578
3rd Qu.: -0.05055
Max.   : 0.56396
```

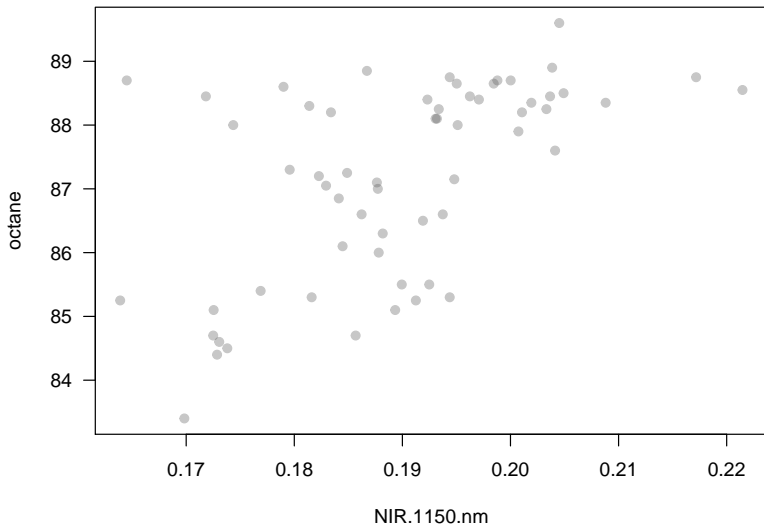
```
which.max(corrs)
```

```
[1] 126
```

```
corrs[which.max(corrs)]
```

```
[1] 0.5639595
```

Scatterplot of Octane with most correlated predictor



Out of curiosity let's try OLS with `lm()`

```
# OLS regression attempt
gas_train <- gasoline[1:50, ]
reg <- lm(octane ~ ., data = gas_train)
summary(reg)
```

Residuals:

ALL 50 residuals are 0: no residual degrees of freedom!

Coefficients: (352 not defined because of singularities)

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	136.74	NA	NA	NA
NIR.900.nm	-2276.35	NA	NA	NA
NIR.902.nm	144.74	NA	NA	NA
...				

Residual standard error: NaN on 0 degrees of freedom

Multiple R-squared: 1, Adjusted R-squared: NaN

F-statistic: NaN on 49 and 0 DF, p-value: NA

Partial Least Squares Regression

```
library(pls)

set.seed(1)
pls1 <- plsr(octane ~ ., ncomp = 10, data = gasoline, subset = train,
             scale = TRUE, validation = "LOO")

pls1

## Partial least squares regression , fitted with the kernel algorithm.
## Cross-validated using 50 leave-one-out segments.
## Call:
## plsr(formula = octane ~ ., ncomp = 10, data = gasoline, subset = train,
```

Summarized Output from `pls1`

```
summary(pls1)
```

```
## Data: X dimension: 50 401
```

```
## Y dimension: 50 1
```

```
## Fit method: kernelpls
```

```
## Number of components considered: 10
```

```
##
```

```
## VALIDATION: RMSEP
```

```
## Cross-validated using 50 leave-one-out segments.
```

```
##      (Intercept)  1 comps  2 comps  3 comps  4 comps  5 comps  6 comps  7 comps
```

```
## CV           1.545   1.321   0.7857   0.2869   0.2254   0.2295   0.2145   0.2287
```

```
## adjCV         1.545   1.322   0.7848   0.2866   0.2251   0.2287   0.2141   0.2279
```

```
##      8 comps  9 comps 10 comps
```

```
## CV           0.2586   0.2710   0.2695
```

```
## adjCV        0.2567   0.2692   0.2676
```

```
##
```

```
## TRAINING: % variance explained
```

```
##      1 comps  2 comps  3 comps  4 comps  5 comps  6 comps  7 comps  8 comps
```

```
## X           64.31   85.24   95.79   97.22   97.59   98.19   98.61   98.74
```

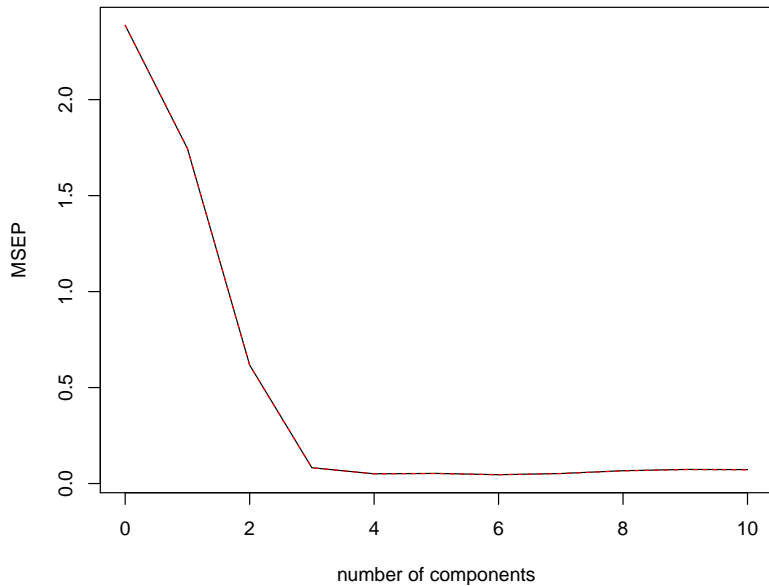
```
## octane      31.59   79.29   97.13   98.49   98.91   99.01   99.10   99.37
```

```
##      9 comps 10 comps
```

```
## X           99.10   99.25
```

```
## octane      99.46   99.57
```


octane



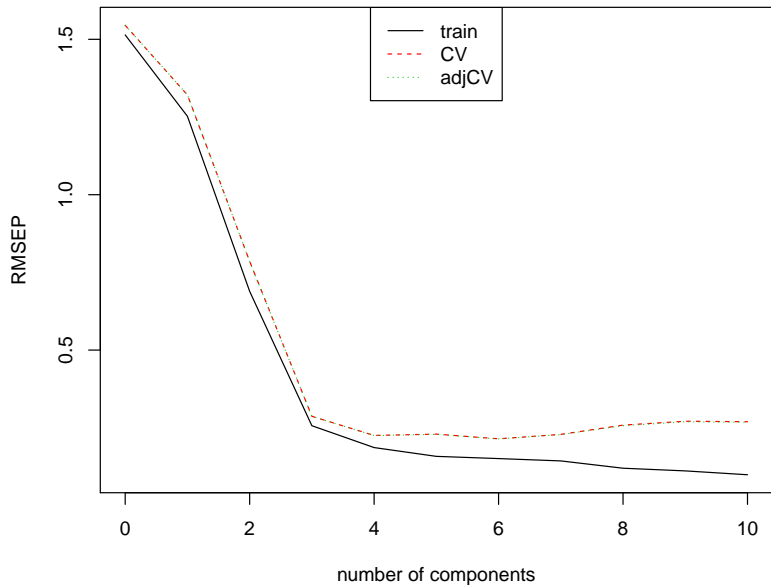
Test MSEs

##	ncomp	MSE_test
## [1,]	1	1.61005872
## [2,]	2	0.56881957
## [3,]	3	0.19325159
## [4,]	4	0.03332153
## [5,]	5	0.19678268
## [6,]	6	0.08161284
## [7,]	7	0.10074208
## [8,]	8	0.26969233

which minimum test MSE?

```
## [1] 4
```

octane



Final PLS Regression

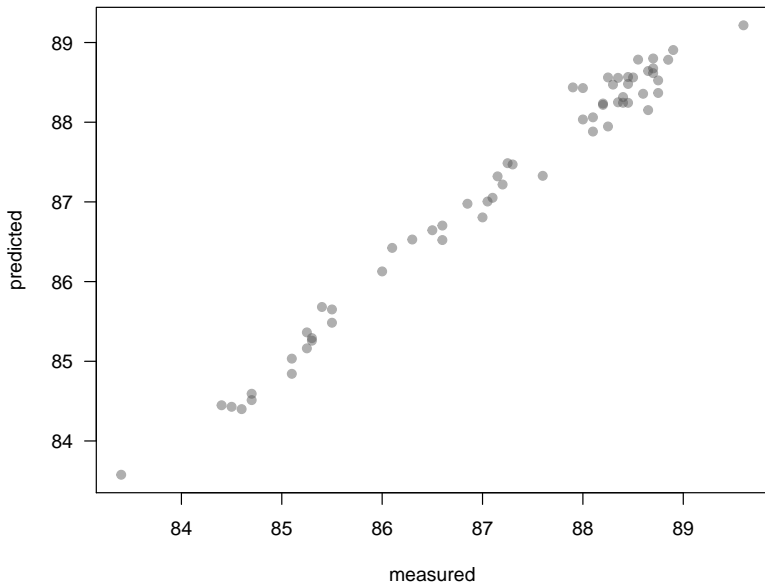
```
pls_fit <- plsr(octane ~ ., ncomp = 4, data = gasoline,  
               scale = TRUE)  
summary(pls_fit)
```



```
## Data:  X dimension: 60 401  
##  Y dimension: 60 1  
## Fit method: kernelpls  
## Number of components considered: 4  
## TRAINING: % variance explained
```

	1 comps	2 comps	3 comps	4 comps
## X	64.97	83.51	93.72	96.33
## octane	30.54	79.79	97.73	98.27

Observed and predicted values (4 PLS comps)



References

- ▶ **Modern Multivariate Statistical Techniques** by Julian Izenman (2008). *Chapter 5, sec 6: Biased Regression Methods*. Springer.
- ▶ **Linear Models with R** by Julian Faraway (2015). *Chapter 11: Shrinkage Methods*. CRC Press.
- ▶ **Some theoretical aspects of partial least squares regression** by Inge Helland (2001). *Chemometrics and Intelligent Laboratory Systems*, 58, 97-107.
- ▶ **Partial Least Squares Regression and Statistical Models** by Inge Helland (1990). *Scandinavian Journal of Statistics*. Vol. 17, No. 2. p. 97-114.

References (French Literature)

- ▶ **La Regression PLS: Theorie et Pratique** by Michel Tenenhaus (1998). Editions, Technip.
- ▶ **Probabilites, analyse des donnees et statistique** by Gilbert Saporta (2011). *Chapter 17: La regression multiple et le modele lineaire general*. Editions Technip, Paris.