# Dimension reduction, principal component analysis (PCA) and principal component regression (PCR)

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# 1 Principal component analysis (PCA)

- Principal component analysis (PCA) is a an exploratory technique which can sometimes help identifying low dimensional structures in high dimensional data.
- There are many such methods, often called dimension reduction methods. Presumably the most classical method (dating back to 1904) is called principal component analysis or PCA.
- PCA is a linear method and is based on the idea of finding a new set of variables which are linear combinations of the original variables and which are uncorrelated and explain as much of the variation in the data as possible.
- A related method is factor analysis (not discussed here)

# 2 Example: Crime rate in the US in 1977

The crime\_rate data contains the crime rate per 100.000 people in 50 US states in 1977.

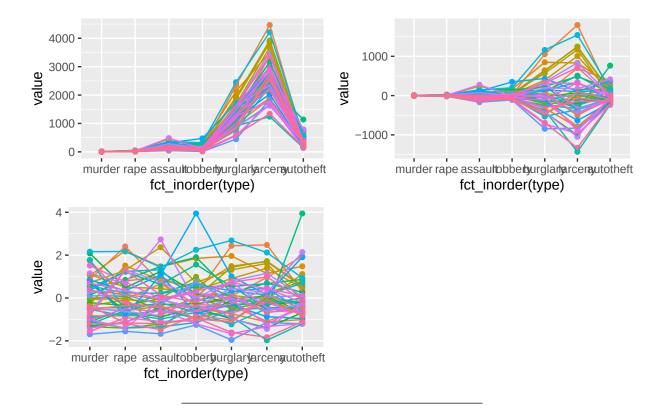
```
crime <- doBy::crime_rate</pre>
state <- rownames(crime)</pre>
crime_state <- cbind(crime, state=state)</pre>
rownames(crime_state) <- NULL</pre>
crime |> head(3)
           murder rape assault robbery burglary larceny autotheft
## Alabama
           14.2
                   25
                            278
                                     97
                                             1136
                                                     1882
                                                                281
## Alaska
             10.8
                    52
                            284
                                     97
                                             1332
                                                     3370
                                                                753
                                             2346
                                                     4467
## Arizona
             9.5
                    34
                            312
                                    138
                                                                440
crime_state |> head(3)
    murder rape assault robbery burglary larceny autotheft
                                                                state
## 1 14.2
             25
                     278
                              97
                                      1136
                                              1882
                                                          281 Alabama
## 2
       10.8
              52
                      284
                               97
                                      1332
                                               3370
                                                          753 Alaska
## 3
              34
                      312
                              138
                                      2346
                                               4467
                                                          440 Arizona
```

## 3 Summarizing multivariate data

## 3.1 Plotting multivariate data

```
## Two handy utilities:
make_long <- function(data., x){</pre>
    data. |>
        as.data.frame() |>
        pivot_longer(-!!sym(x), names_to = "type", values_to = "value")
}
do_plot1 <- function(data.){</pre>
    data. |>
        ggplot(aes(x=fct_inorder(type), y=value, group=state, color=state)) +
        geom_point() + geom_line() + theme(legend.position="none")
}
## Uncentered data
dat_long <- crime_state |> make_long("state")
## Centered but unscaled data
dat_c <- crime_state |> doBy::scale_df(center=T, scale=F)
dat_c_long <- dat_c |> make_long("state")
## Centered and scaled data
dat_cs <- crime_state |> doBy::scale_df(center=T, scale=T)
dat_cs_long <- dat_cs |> make_long("state")
```

```
p1 <- do_plot1(dat_long)
p2 <- do_plot1(dat_c_long)
p3 <- do_plot1(dat_cs_long)
cowplot::plot_grid(p1, p2, p3)
```



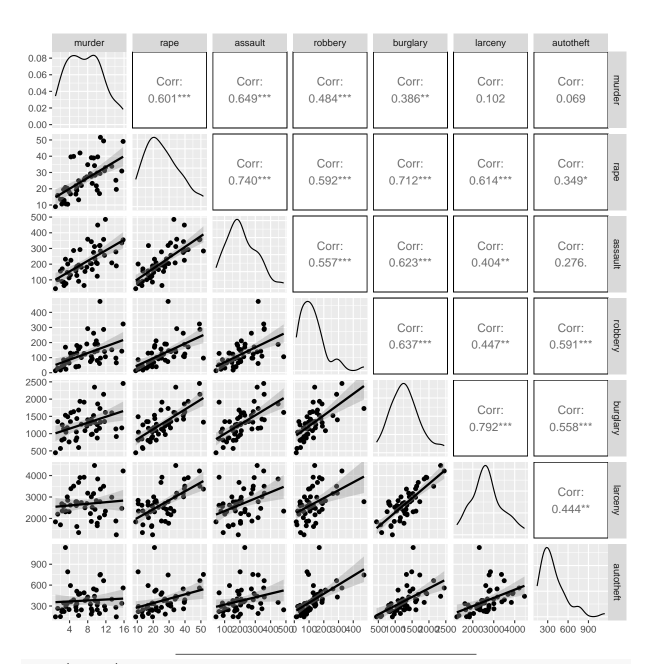
#### 3.2 Correlations

doBy::truncate0(cormat, tol=0.6)

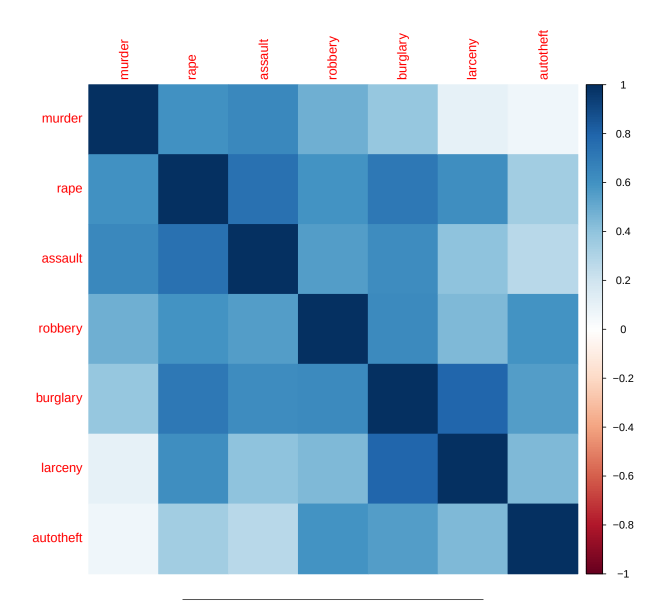
The relationship between some variables appear to be approximately linear, so we might want to calculate all pairwise correlations:

```
cormat <- cor(crime)</pre>
round(10*cormat)
             murder rape assault robbery burglary larceny autotheft
## murder
                       6
                                 6
                  10
                                         5
                                                   4
## rape
                   6
                       10
                                         6
                                                   7
                                                           6
                                                                      3
## assault
                        7
                                10
                                         6
                                                   6
                                                           4
                                                                      3
## robbery
                        6
                                6
                                        10
                                                   6
                                                           4
                                                                      6
## burglary
                                                  10
                                                           8
## larceny
                        6
                                 4
                                         4
                                                   8
                                                          10
                                                                      4
## autotheft
                                         6
                                                   6
                                                                     10
```

```
## 7 x 7 sparse Matrix of class "dgCMatrix"
             murder rape assault robbery burglary larceny autotheft
##
## murder
               1.00 0.60
                            0.65
## rape
               0.60 1.00
                            0.74
                                                      0.61
## assault
               0.65 0.74
                             1.00
                                              0.62
## robbery
                                     1.00
                                              0.64
## burglary
                    0.71
                             0.62
                                                      0.79
                                     0.64
                                              1.00
## larceny
                    0.61
                                              0.79
                                                      1.00
## autotheft
```



library(corrplot)
corrplot::corrplot(cormat, method="color")



# 4 Principal component analysis (PCA)

## 4.1 Basic idea

Easy to explain with only three variables

```
violent <- crime[,1:3]
violent |> head(3)
##
         murder rape assault
## Alabama 14.2 25
## Alaska
             10.8
                    52
                           284
## Arizona
                           312
violent |> cor()
##
           murder rape assault
## murder
           1.00 0.60
                         0.65
## rape
            0.60 1.00
                          0.74
## assault 0.65 0.74
                          1.00
```

3D-plot of the data  $\,$ 

Denote columns of observed variables by  $y_1$ ,  $y_2$  and  $y_3$ . The idea is to find a new set of variables  $x_1$ ,  $x_2$  and  $x_3$  such that the x variables 1) are uncorrelated and 2) explain as much of the variation in the data as possible.

The first principal component  $x_1$  is a linear combination (weighted sum) of the y variables:

$$x_1 = r_{11} \cdot y_1 + r_{21} \cdot y_2 + \dots + r_{31} \cdot y_3$$

where  $r_{11}, r_{21}, r_{31}$  are the weights. The weights are chosen so that  $x_1$  explains as much of the variation in the data as possible.

The second principal component  $x_2$  is also a linear combination of the y variables

$$x_2 = r_{12} \cdot y_1 + r_{22} \cdot y_2 + \dots + r_{32} \cdot y_3$$

where  $r_{12}, r_{22}, r_{32}$  are the weights. The weights are chosen so that 1)  $x_2$  explains second most variation in the data as possible and 2) is uncorrelated with  $x_1$ .

The third principal component  $x_3$  is a linear combination of the y variables:

$$x_3 = r_{13} \cdot y_1 + r_{23} \cdot y_2 + \dots + r_{33} \cdot y_3$$

where  $r_{13}, r_{23}, r_{33}$  are the weights. The weights are chosen so that 1)  $x_3$  explains third most variation in the data as possible and 2) is uncorrelated with  $x_1$  and  $x_2$ .

The weights  $r_{ij}$  are chosen so that (there are a couple of extra details)

- 1. Each  $x_j$  has length 1
- 2.  $var(x_1) \ge var(x_2) \ge var(x_1)$
- 3. All principal components  $x_1, x_2, x_3$  are uncorrelated.

It can be shown that

$$v_{total} = var(y_1) + var(y_2) + var(y_3) = var(x_1) + var(x_2) + var(x_3) \label{eq:var}$$

So

$$var(x_1)/v_{total}, (var(x_1) + var(x_2))/v_{total}$$

is the variance explained by the first and the first two principal components, respectively.

```
pca <- prcomp(violent, center=T, scale.=T)</pre>
summary(pca)
## Importance of components:
##
                           PC1
                                 PC2
                                         PC3
## Standard deviation
                         1.526 0.646 0.5046
## Proportion of Variance 0.776 0.139 0.0849
## Cumulative Proportion 0.776 0.915 1.0000
pca$x |> head(3) ## Principal components / scores
           PC1 PC2
##
## Alabama -1.3 -1.28 -0.28
## Alaska -2.3 0.71 1.10
## Arizona -1.4 0.23 -0.19
```

```
pca$x |> cov()
                    PC2
           PC1
## PC1 2.3e+00 -4.6e-16 1.8e-16
## PC2 -4.6e-16 4.2e-01 6.4e-17
## PC3 1.8e-16 6.4e-17 2.5e-01
pca$rotation ## Loadings / weights
            PC1
                  PC2
                        PC3
## murder
          -0.55 - 0.82
                       0.15
## rape
          -0.58 0.51 0.63
## assault -0.59 0.26 -0.76
```

#### Interpretation:

```
cor(violent, pca$x[,1:2])

## PC1 PC2

## murder -0.85 -0.53

## rape -0.89 0.33
```

#### Interpretation:

## assault -0.91 0.17

- $x_1$  is a weighted sum of the three variables. The weights are approximately equal. Hence  $x_1$  is a measure of the average crime rate.
- $x_2$  gets negative contributions from murder and positive contributions from assault and rape. Hence  $x_2$  is a measure of the violent crime rate.

### 4.2 The full dataset

Principal components can be obtained with prcomp()

Default is that variables are centered to have mean zero (center=T), but when the variation of the variables are very different it is often a good idea to standardize variables to have variance one (scale.=T).

The first PC explains about 60% of the variation and with the first three components about 85% of the variation is explained.

## 4.3 Interpretation - loadings / rotations

```
cor(crime, pca$x[,1:3])
##
              PC1
                     PC2
                             PC3
## murder
             -0.61 -0.700 -0.152
## rape
             -0.88 -0.189 0.208
## assault
             -0.80 -0.382 0.059
## robbery
             -0.81 0.047 -0.422
## burglary
            -0.89 0.226 0.179
## larceny
             -0.72 0.448 0.459
## autotheft -0.60 0.559 -0.484
```

- x<sub>1</sub> is a weighted sum of all variables. The weights are approximately equal (when data is centered and scaled).
   Hence x<sub>1</sub> is a measure of the average crime rate.
- $x_2$  gets negative contributions violent crime and positive contributions from economical crime. A state with much voilence tends to have a negative value of  $x_2$ ; a state with economical crime tends to have a positive value of  $x_3$ .
- From thereof the picture is more blurred.

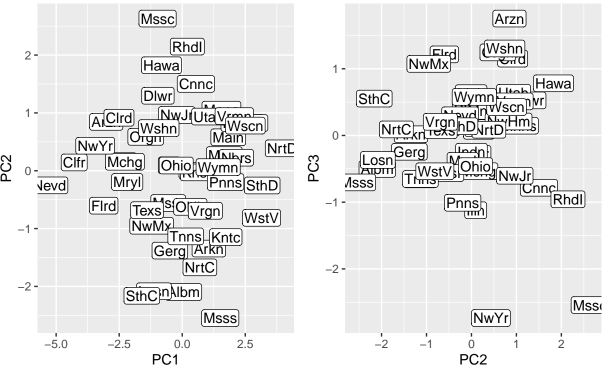
## 4.4 Principal components / scores

The principal components / scores:

```
X <- pca$x
head(X, 5)
##
               PC1
                    PC2
                            PC3
                                   PC4
                                        PC5
                                              PC6
## Alabama
              0.05 -2.10 -0.502
                                0.251 0.498 -0.43
                                                   0.118
## Alaska
             -2.42 0.17 0.070 1.160 1.470 1.50
                                                   0.465
             -3.01 0.84 1.752 -0.116 0.280 -1.07
              1.05 -1.35 0.018 0.022 0.023 0.39 -0.311
## Arkansas
## California -4.28  0.14 -0.276  0.025  0.058  0.38 -0.464
```

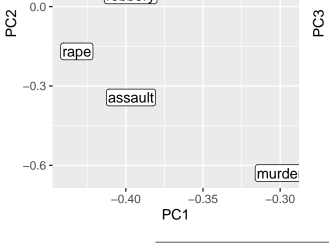
## 4.5 Scores plot

```
state2 <- abbreviate(state, 4)</pre>
state2 |> head(10)
##
        Alabama
                       Alaska
                                   Arizona
                                                Arkansas
                                                            California
                                                                            Colorado
##
         "Albm"
                       "Alsk"
                                    "Arzn"
                                                  "Arkn"
                                                                 "Clfr"
                                                                              "Clrd"
## Connecticut
                    Delaware
                                   Florida
                                                 Georgia
         "Cnnc"
                       "Dlwr"
                                     "Flrd"
                                                  "Gerg"
do_plot2 <- function(dat, x, y, lab){</pre>
    dat <- dat |> as.data.frame()
    dat |> ggplot(aes(x=!!sym(x), y=!!sym(y), label=lab)) +
         geom_point() +
         geom_label(label.padding = unit(0.1, "lines"))
}
X <- pca$x
p1 <- X |> do_plot2("PC1", "PC2", state2)
p2 <- X |> do_plot2("PC2", "PC3", state2)
cowplot::plot_grid(p1, p2, nrow=1)
```

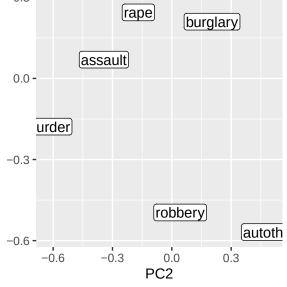


## 4.6 Loading plots





robbery



# 5 Example: NIRmilk

In these data NIR (near infrared) measurements are made at 152 wavelengths on 17 milk samples: Milk runs trough a glass tube. Near infrared light is sent through the tube. The transmittance (fraction of electromagnetic power) at different wavelengths is recorded.

The samples are also analyzed for contents of fat, lactose, protein and drymatter.

A natural question is: Can fat, lactose, protein and drymatter content be predicted from the NIR measurements (which are easy and cheap to obtain).

PCA is an excellent tool in this connections

```
nir_milk <- doBy::nir_milk</pre>
ynir <- nir_milk$y |> as_tibble()
xnir <- nir_milk$x |> as_tibble()
ynir |> head(3)
## # A tibble: 3 x 5
##
    sample fat protein lactose
                                     dm
##
            <dbl>
     <chr>>
                    <dbl>
                            <dbl> <dbl>
## 1 s01
             4.17
                     3.64
                             4.53 13.1
## 2 s02
             4.23
                     3.55
                             5.56 14.1
             3.90
## 3 s03
                     4.30
                             5.49
                                  14.5
xnir[,1:6] |> head(3)
## # A tibble: 3 x 6
    sample X964 X968
                           X972
                                   X976 X979
##
##
    <chr>
             <dbl> <dbl> <dbl>
                                  <dbl> <dbl>
## 1 s01
            0.0536 0.0556 0.0608 0.0782 0.109
## 2 s02
            0.0693 0.0713 0.0775 0.0980 0.135
            0.0677 0.0703 0.0773 0.0989 0.136
## 3 s03
```

```
ynir0 <- ynir |> select(-sample)
xnir0 <- xnir |> select(-sample)
samp <- xnir$sample</pre>
wave <- xnir0 |> colnames() |> gsub("X", "", x=_) |> as.numeric()
samp |> head(3)
## [1] "s01" "s02" "s03"
wave |> head(3)
## [1] 964 968 972
xnir_long <- xnir |>
    pivot_longer(-sample, names_to="wave", values_to="value") |>
    mutate(wavelength = as.numeric(gsub("X", "", wave)))
head(xnir_long, 3)
## # A tibble: 3 x 4
##
     sample wave
                   value wavelength
##
     <chr> <chr>
                  <dbl>
                               <dbl>
            X964 0.0536
                                 964
## 1 s01
                  0.0556
                                 968
## 2 s01
            X968
## 3 s01
            X972
                  0.0608
                                 972
plot_xnir <- xnir_long |>
    ggplot(aes(x=wavelength, y=value, group=sample, colour=sample))
plot_grid(plot_xnir + geom_line(),
          plot_xnir + geom_smooth(span=.25, se=F), nrow=1)
                                                           0.4 -
   0.4
                                                           0.3 -
   0.3 -
                                                        value 0.2 ·
  0.2
   0.1
                                                           0.1 -
   0.0 -
                                                           0.0 -
                        1200
                                      1400
                                                                                1200
         1000
                                                                  1000
                                                                                              1400
                        wavelength
                                                                                wavelength
```

A good question is if we should scale the variables to have the same variance or not.

- 1. All measurements are the same quantity but at different wavelengths which suggests that scaling might not be necessary.
- 2. On the other hand, the plot indicates that the variances are different for different wavelengths.

```
xnir_center <- doBy::scale_df(xnir, center = TRUE, scale = FALSE)</pre>
```

```
xnir_center_long <-
    xnir_center |>
    pivot_longer(-sample, names_to = "wave", values_to = "value") |>
    mutate(wavelength = as.numeric(gsub("X", "", wave)))
```

```
plot_xnir_center <- xnir_center_long |>
    ggplot(aes(wavelength, value, group=sample, colour=sample))
plot_grid(plot_xnir_center + geom_line(),
          plot_xnir_center + geom_smooth(span=.2, se=F),
          plot_xnir_center + geom_smooth(span=.7, se=F), nrow=1)
                                                                              0.06 -
                                         0.050 -
    0.05 -
                                         0.025
                                                                               0.03 -
    0.00
                                         0.000
                                     value
value
                                                                              0.00
                                         -0.025
   -0.05 -
                                                                              -0.03
                                        -0.050 -
                                        -0.075 -
          1000
                 1200
                         1400
                                                1000
                                                       1200
                                                             1400
                                                                                    1000
                                                                                            1200
                                                                                                   1400
               wavelength
                                                     wavelength
                                                                                          wavelength
```

We continue with unscaled data (not terribly important in this case)

```
pca0 <- prcomp(xnir0, rank=6)</pre>
summary(pca0)
## Importance of first k=6 (out of 17) components:
##
                            PC1 PC2
                                         PC3
                                                 PC4
                                                          PC5
## Standard deviation
                          0.158\ 0.122\ 0.0958\ 0.02138\ 0.00488\ 0.00371
## Proportion of Variance 0.506 0.298 0.1853 0.00923 0.00048 0.00028
## Cumulative Proportion 0.506 0.804 0.9898 0.99898 0.99946 0.99974
pca_scaled <- prcomp(xnir0, scale=T, rank=6)</pre>
summary(pca_scaled)
## Importance of first k=6 (out of 17) components:
##
                            PC1
                                 PC2 PC3
                                                        PC5
                                                                PC6
## Standard deviation
                          8.679 6.934 5.246 1.3434 0.38062 0.24169
## Proportion of Variance 0.492 0.314 0.180 0.0118 0.00095 0.00038
## Cumulative Proportion 0.492 0.807 0.986 0.9983 0.99923 0.99962
```

Hence, 80% of the total variation in a 150–dimensional data set is explained by the first two principal components and practically all variation is explained by the first three principal components. This is quite a substantial reduction in dimension.

## 5.1 Plotting rotations

Recall: rotations / loadings are the weights given to each measurement when forming the principal components. We plot loadings against wavelengths:

```
rot <- pca0$rotation[, 1:3] |> as.data.frame()
rot$wave <- wave</pre>
rot_long <- rot |> pivot_longer(-wave, names_to = "PC")
rot_long |> head(3)
## # A tibble: 3 x 3
##
      wave PC
##
     <dbl> <chr>
                    <dbl>
       964 PC1
## 1
                   0.0108
##
       964 PC2
                   0.0268
## 3
       964 PC3
                  -0.124
plot_rotation <- rot_long |> ggplot(aes(x=wave, y=value, color=PC)) +
    theme(legend.position="bottom")
plot_grid(plot_rotation + geom_line()
          plot_rotation + geom_smooth(span=.5),
          plot_rotation + geom_smooth(span=2), nrow=1)
                                                                                 0.2 -
    0.2
                                           0.2
                                                                                 0.1
                                           0.1
    0.1
                                      value
                                                                            value
                                          0.0 -
                                                                                0.0 -
    0.0
                                          -0.1
                                                                                -0.1
                                          -0.2
         1000
                 1200
                         1400
                                               1000
                                                       1200
                                                                1400
                                                                                     1000
                                                                                              1200
                                                                                                      1400
                   wave
                                                         wave
                                                                                               wave
```

# 6 Principal component regression (PCR)

Next we turn to the regression problem: We want to predict, say, the fat content from the NIR measurements.

We have 17 samples and 153 predictor variables.

In the usual regression setting the model matrix ynir is  $17 \times 153$  (an extra column of 1s corresponding to the intercept is added).

This is an example of a p larger than n problem; there are more predictors than observations.

A straight forward alternative is principal component regression or PCR

- $1. \;$  First derive principal components of the explanatory variables and
- 2. Then use these principal components as explanatory variables.

Combine the first three principal components with the ynirs and regress the ynirs on (some of) the principal components.

```
doBy::truncate0(cor(ynir0, pca0$x[,1:6]), tol=0.4)
## 4 x 6 sparse Matrix of class "dgCMatrix"
           PC1 PC2 PC3 PC4 PC5 PC6
##
## fat
          0.47 . 0.84
## protein . 0.61 0.74
## lactose 0.93 .
## dm
          0.61 .
                    0.79
nir_ext <- cbind(pca0$x[, 1:6], ynir0)</pre>
nir_ext |> head(3)
##
      PC1 PC2
                    PC3
                           PC4
                                   PC5
                                            PC6 fat protein lactose dm
## 1 -0.11 -0.18 -0.0079 0.0074 0.00099 0.00064 4.2
                                                       3.6
                                                               4.5 13
## 2 0.18 -0.11 -0.0881 0.0069 0.00181 0.00025 4.2
                                                               5.6 14
                                                       3.5
## 3 0.13 0.12 -0.0454 0.0156 0.01020 -0.00358 3.9
                                                        4.3
                                                               5.5 14
cor(ynir0, pca0$x[,1:4])
            PC1
                    PC2
                         PC3
##
                                  PC4
## fat
          0.474 -0.2610 0.84 -0.0095
## protein 0.045 0.6146 0.74 0.2364
## lactose 0.928 0.1677 -0.32 -0.0027
## dm
          0.614 0.0032 0.79 0.0370
```

Now, we can try to make a (multiple) regression explaining the ynirs not directly in terms of the wavelengths but in terms of the principal components (which in turn are derived from the wavelengths):

```
m1 <- lm(fat ~ PC1 + PC3 , data = nir_ext)</pre>
```

The model explains practically all variation in data:

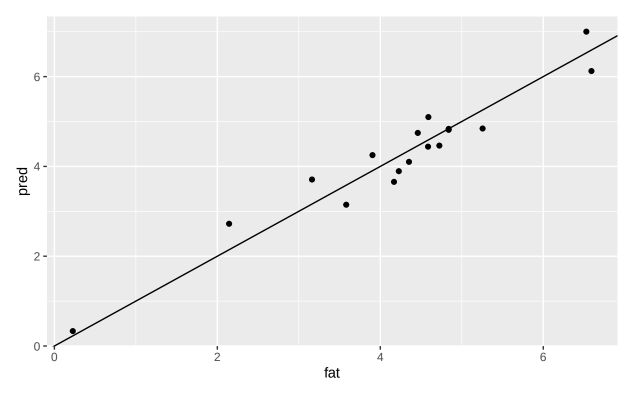
```
m1 |> broom::glance() |> pander::pander()
```

Table 1: Table continues below

r.squared	adj.r.squared	sigma	statistic	p.value	df	logLik
0.931	0.9212	0.4168	94.51	7.414e-09	2	-7.595

AIC	BIC	deviance	df.residual	nobs
23.19	26.52	2.432	14	17

```
nir_ext2 <- nir_ext |> modelr::add_predictions(m1)
nir_ext2 |> ggplot(aes(x=fat, y=pred)) +
    geom_point() + geom_abline(slope=1, intercept=0)
```



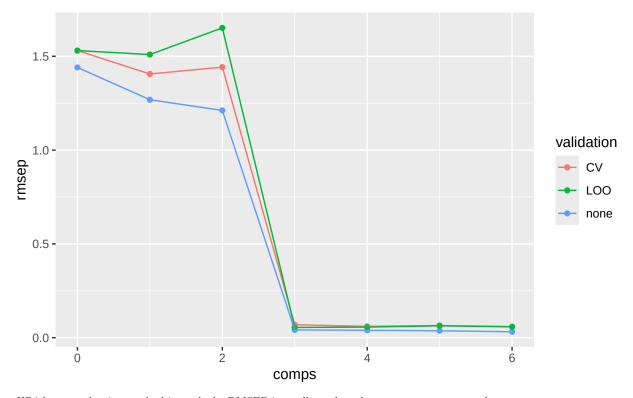
Notice: Since the first three PCs explain practically all variation in data we need not include any additional PCs in the regression model. However, it could be the case that not all of the PCs had a significant effect as a predictor, but that is not the case here.

## 7 Cross validation

The predictive ability of the regression model above is assessed on the basis of the data and here we should do cross validation. The pls package has a function for this purpose.

```
library(pls)
ynir$X <- as.matrix(xnir0)</pre>
ynir[1:3,1:4]
## # A tibble: 3 \times 4
##
    sample fat protein lactose
##
    <chr> <dbl>
                    <dbl>
                             <dbl>
## 1 s01
             4.17
                     3.64
                              4.53
## 2 s02
             4.23
                     3.55
                              5.56
## 3 s03
             3.90
                      4.30
                              5.49
ynir$X[1:3, 1:5]
         X964 X968 X972 X976 X979
## [1,] 0.054 0.056 0.061 0.078 0.11
## [2,] 0.069 0.071 0.078 0.098 0.13
## [3,] 0.068 0.070 0.077 0.099 0.14
```

Best predictive results with three principal components:



Whichever evaluation method is used, the RMSEP is smallest when three components are used.

Notice that scores we compute "manually" are the same as those computed by pcr:

```
## Comp 1 Comp 2 Comp 3 Comp 4 Comp 5 Comp 6
## 1 -0.11 -0.18 -0.0079 0.0074 0.00099 0.00064
## 2 0.18 -0.11 -0.0881 0.0069 0.00181 0.00025
## 3 0.13 0.12 -0.0454 0.0156 0.01020 -0.00358

pca0$x[, 1:6] |> head(3)

## PC1 PC2 PC3 PC4 PC5 PC6
## [1,] -0.11 -0.18 -0.0079 0.0074 0.00099 0.00064
## [2,] 0.18 -0.11 -0.0881 0.0069 0.00181 0.00025
## [3,] 0.13 0.12 -0.0454 0.0156 0.01020 -0.00358
```

## 8 Take-home message

- PCR is not a feature selection method as such: It does not extract significant wavelengths and throw the rest away.
- But for these data, it probably makes very little sense to talk about a few significant wavelengths. Instead it is perhaps relevant to look at collections of wavelengths.
- Can see PCA as a necessary preprocessing step (dimension reduction) before regression.
- Very common to do such dimension reductions before other analyses.