D. Makowski Université Paris-Saclay INRAE

Outline

- Definition & main principles
- Several extensions of linear regression
- Trees and forests
- Deep learning

Linear regression

$$Y = X\theta + \varepsilon$$

Y is a N-vector of output values, X is a matrix N by P of inputs (design), θ is a P-vector of parameters, ε is a N-vector of residuals

- Simple linear regression (including one input only),
- Polynomial regression
- Linear model with interactions

$$Y = X\theta + \varepsilon$$

$$\begin{pmatrix} y_1 \\ y_2 \\ \dots \\ y_N \end{pmatrix} = \begin{pmatrix} x_{11} & x_{12} & \dots & x_{1P} \\ x_{21} & x_{22} & \dots & x_{2P} \\ \dots & \dots & \dots & \dots \\ x_{N1} & x_{N2} & \dots & x_{NP} \end{pmatrix} \begin{pmatrix} \theta_1 \\ \theta_2 \\ \dots \\ \theta_P \end{pmatrix} + \begin{pmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \dots \\ \varepsilon_N \end{pmatrix}$$

$$Y = X\theta + \varepsilon$$

$$\begin{pmatrix} y_1 \\ \dots \\ y_N \end{pmatrix} = \begin{pmatrix} 1 & x_1 \\ \dots & \dots \\ 1 & x_N \end{pmatrix} \begin{pmatrix} \theta_1 \\ \theta_2 \end{pmatrix} + \begin{pmatrix} \varepsilon_1 \\ \dots \\ \varepsilon_N \end{pmatrix}$$

Several issues

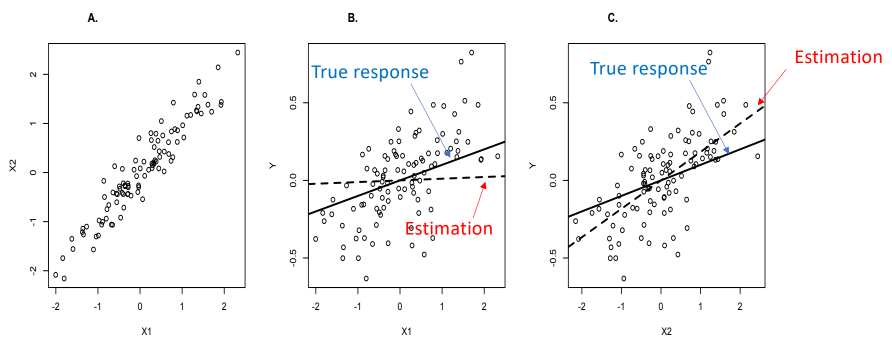
- Inputs X₁, ..., X_P are sometimes (strongly) correlated
- Inputs X_1 , ..., X_p may have non-linear effects (unknown response shape),
- Too many inputs
- Need to estimate extreme responses, not mean response

Several issues

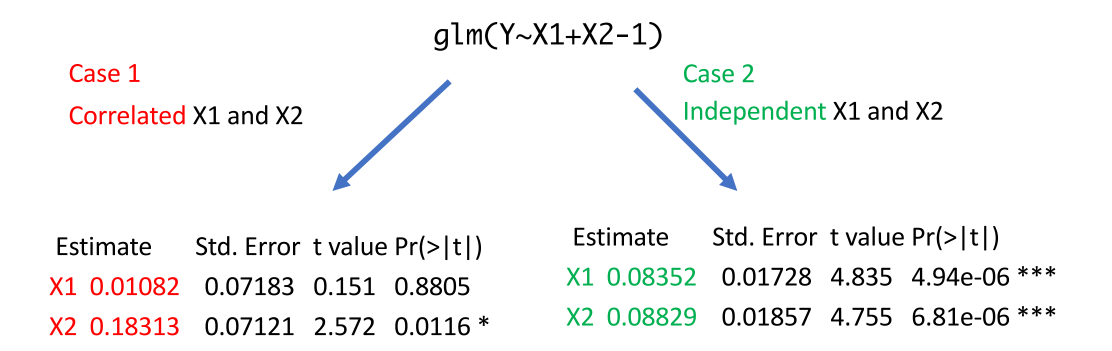
- Inputs X₁, ..., X_P are sometimes (strongly) correlated
- Inputs X_1 , ..., X_p may have non-linear effects (unknown response shape),
- Too many inputs
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Why can correlated inputs be an issue?

100 data generated with
$$Y = 0.1X_1 + 0.1X_2 + \varepsilon$$



Correlation between X1 and X2 = 0.95



Both results are obtained with 100 data

$glm(Y\sim X1+X2-1)$

Case 1

Correlated X1 and X2

Case 2
Independent X1 and X2

Estimate Std. Error t value Pr(>|t|) Estimate Std. Error t value Pr(>|t|) X1 0.01082 0.07183 0.151 0.8805 X1 0.08352 0.01728 4.835 4.94e-06 ***

X2 0.18313 0.07121 2.572 0.0116 * X2 0.08829 0.01857 4.755 6.81e-06 ***

Question 1. Effect of X1 significant in cases 1 and 2?

A. Yes

B. No

$glm(Y\sim X1+X2-1)$

Case 1

Correlated X1 and X2

Case 2
Independent X1 and X2

Estimate Std. Error t value Pr(>|t|)

Y1 0.08

X1 0.01082 0.07183 0.151 0.8805

X2 0.18313 0.07121 2.572 0.0116 *

Estimate Std. Error t value Pr(>|t|)

X1 0.08352 0.01728 4.835 4.94e-06 ***

X2 0.08829 0.01857 4.755 6.81e-06 ***

Question 1. Effect of X1 significant in cases 1 and 2?

A. Yes

B. No

Question 2. Which case shows the most accurate estimated values?

A. Case 1

B. Case 2

Idea: use independent combinations of X_1 , X_2 , ..., X_P

- Principal component regression (PCR)
- Partial least square regression (PLSR)

Principal component analysis

- Replace the initial input variables (X₁, ..., X_P) with new independent variables
- The new variables correspond to linear combinations of the old ones
- These linear combinations are chosen so as to have a maximum variance

$$Y = \alpha_0 + \alpha_1 X_1 + \dots + \alpha_P X_P + \varepsilon$$

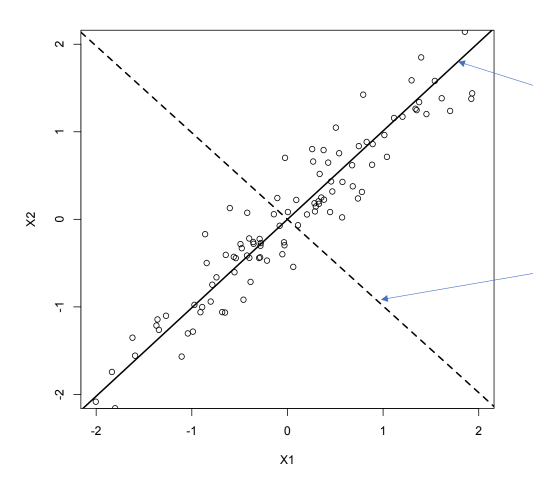
$$Z_1 = \sum_{j=1}^{P} \beta_{1j} X_j$$
 $Z_2 = \sum_{j=1}^{P} \beta_{2j} X_j$ etc.

$$Y = \gamma_0 + \gamma_1 Z_1 + \dots + \gamma_K Z_K + \varepsilon$$



The components are calculated from the eigenvectors of the variance-covariance matrix of

→ Diagonalization of the variance-covariance matrix



Component 1

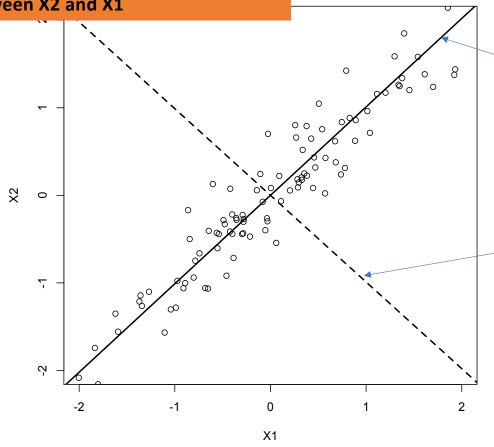
$$Z_1 = 0.704X_1 + 0.711X_2$$

Component 2

$$Z_2 = 0.711X_1 - 0.704X_2$$

Question 1: Component 1 is closer to

- A. The mean of X1 and X2
- B. The difference between X2 and X1



Component 1

$$Z_1 = 0.704X_1 + 0.711X_2$$

Component 2

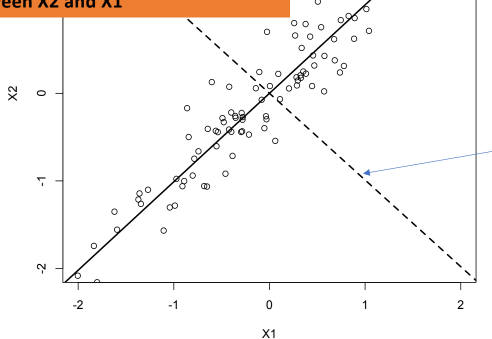
$$Z_2 = 0.711X_1 - 0.704X_2$$

Question 1: Component 1 is closer to

- A. The mean of X1 and X2
- B. The difference between X2 and X1

Question 2: Component 2 is closer to

- A. The mean of X1 and X2
- B. The difference between X2 and X1



Component 1

$$Z_1 = 0.704X_1 + 0.711X_2$$

Component 2

$$Z_2 = 0.711X_1 - 0.704X_2$$

Replace

$$Y = \alpha_1 X_1 + \alpha_2 X_2 + \varepsilon$$

by

$$Y = b_1 CP_1 + \varepsilon$$

= $b_1 (0.704X_1 + 0.711X_2) + \varepsilon$
= $0.097X_1 + 0.098X_2 + \varepsilon$

Example: maize biomass prediction

Objective:

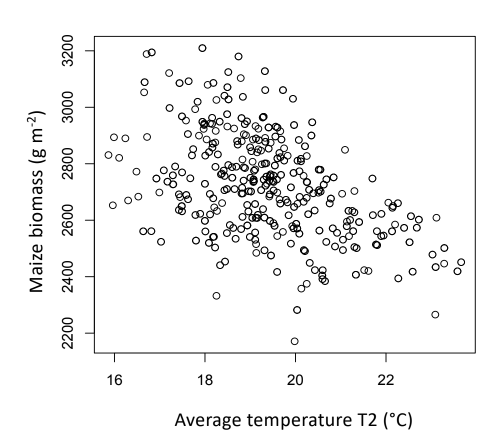
Develop a simple model predicting the final biomass of maize, noted B (g m⁻²), from 6 input variables describing the climatic conditions during the growing season, under optimal water conditions.

Data:

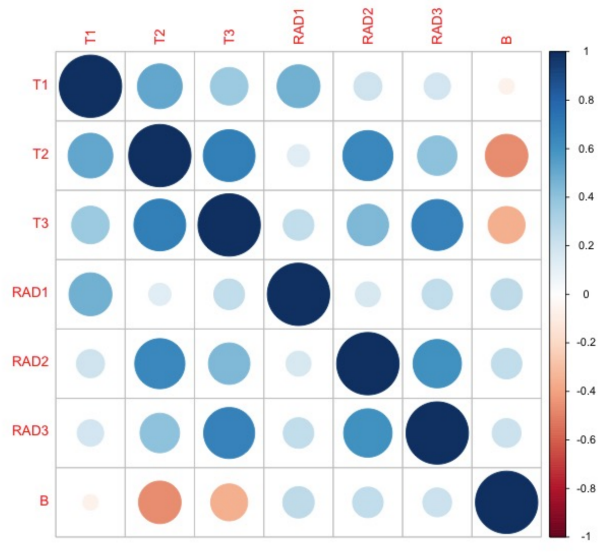
- 680 biomass data obtained for 40 different sites in France and for 17 years (1995 to 2011).
- Mean temperatures during the first part of the growing season (day 1 to day 50), during the second part of the growing season (day 51 to day 100) and during the last part of the growing season (day 101 to day 150). They are noted (T1, T2, T3).
- Average radiations during the same periods (RAD1, RAD2, RAD3).

https://github.com/davemakowski/TP machinelearning

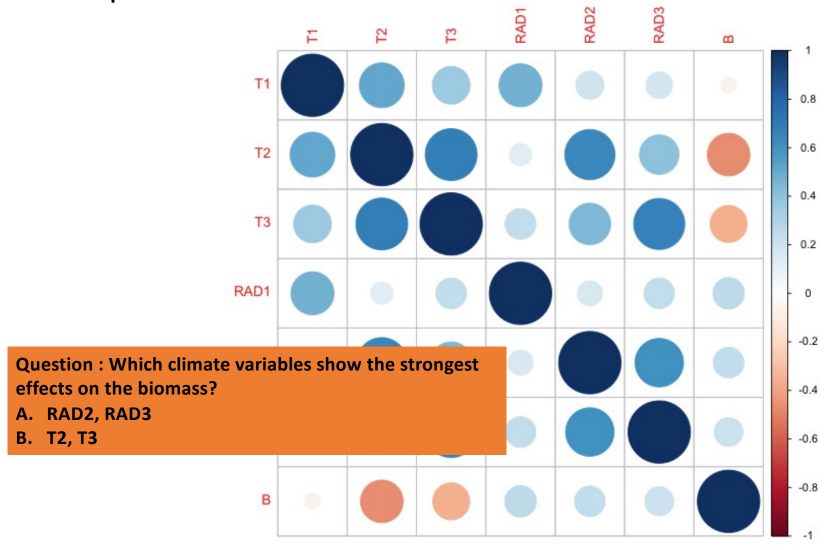
Example: maize biomass



Example: maize biomass



Example: maize biomass



```
library(pls)
Mod_pcr<-pcr(B~T1+T2+T3+RAD1+RAD2+RAD3, data=DataSet,
validation="L00", scale="TRUE")
summary(Mod_pcr)</pre>
```

> summary(Mod_pcr)

Data: X dimension: 680 6

Y dimension: 680 1

Fit method: svdpc

Number of components considered: 6

VALIDATION: RMSEP

Cross-validated using 680 leave-one-out segments.

	(Intercept)	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
CV	182.1	182.3	180.2	138.3	120.4	71.92	64.51
adjCV	182.1	182.3	180.2	138.3	120.4	71.92	64.51

TRAINING: % variance explained

	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
X	49.60384	71.917	84.77	93.65	98.51	100.0
В	0.05284	2.998	43.12	56.92	84.66	87.7

Question: Is the average error of prediction lower than 100 g/m²?

A. No

> summary(Mod_pcr)

B. Yes

Data: X dimension: 680 6 C. It depends on the number of components included

Y dimension: 680 1

Fit method: svdpc

Number of components considered: 6

VALIDATION: RMSEP

Cross-validated using 680 leave-one-out segments.

	(Intercept)	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
CV	182.1	182.3	180.2	138.3	120.4	71.92	64.51
adjCV	182.1	182.3	180.2	138.3	120.4	71.92	64.51

TRAINING: % variance explained

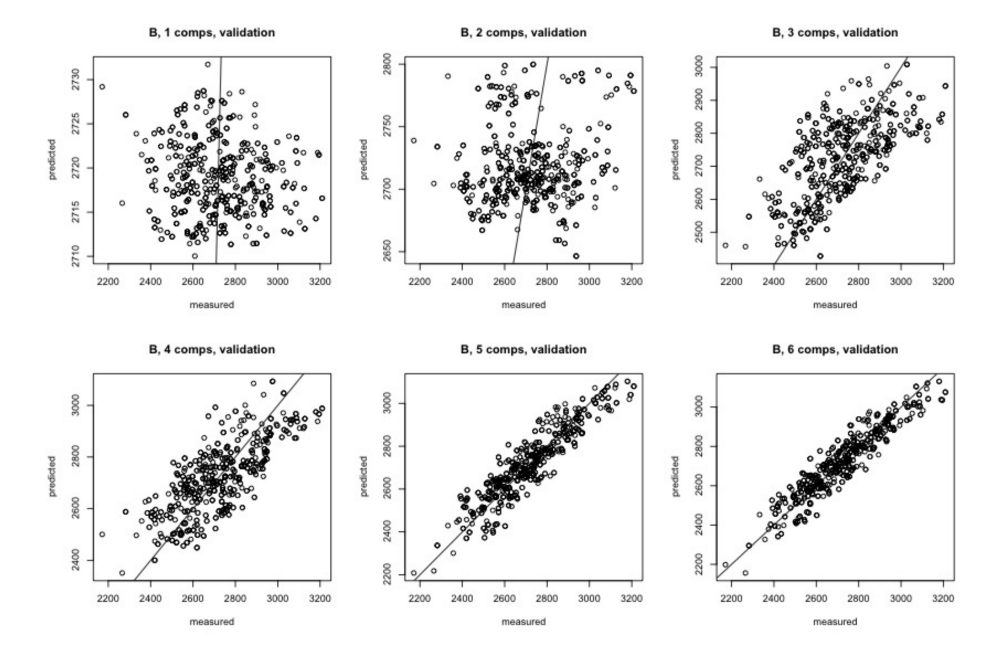
	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
Χ	49.60384	71.917	84.77	93.65	98.51	100.0
В	0.05284	2.998	43.12	56.92	84.66	87.7

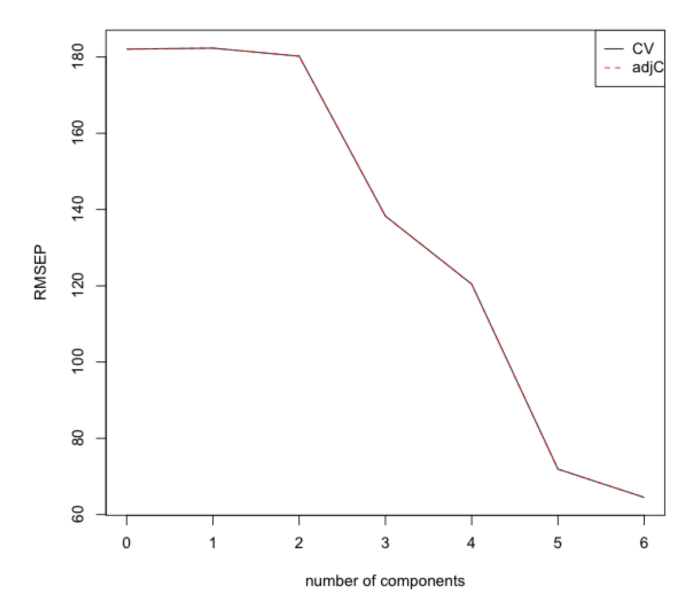
```
par(mfrow=c(2,3))

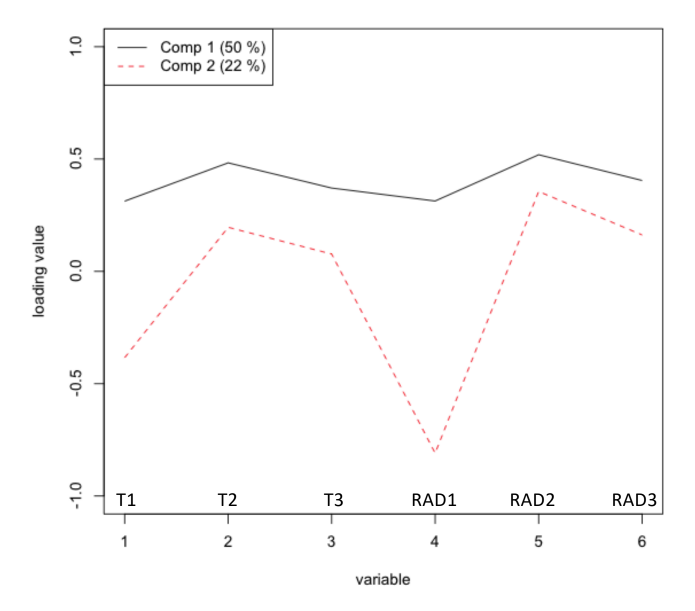
plot(Mod_pcr, line=TRUE, ncomp=1)
plot(Mod_pcr, line=TRUE, ncomp=2)
plot(Mod_pcr, line=TRUE, ncomp=3)
plot(Mod_pcr, line=TRUE, ncomp=4)
plot(Mod_pcr, line=TRUE, ncomp=5)
plot(Mod_pcr, line=TRUE)

par(mfrow=c(1,1))

plot(RMSEP(Mod_pcr), legend ="topright")
plot(Mod_pcr, "loadings", comps=1:2, legendpos="topleft", ylim=c(-1,1))
```







Difference between PCR and PLSR

- In PLSR, the components are determined to be strongly correlated to the values of Y
- Iterative determination of the components in PLSR
- Rarely large differences between PCR and PLSR, but PLSR often require fewer components

```
Mod_pls<-plsr(B~T1+T2+T3+RAD1+RAD2+RAD3, data=DataSet, validation="L00",
scale="TRUE")
summary(Mod_pls)</pre>
```

Question: Is the average error of prediction lower than 100 g/m²?

C. It depends on the number of components included

A. No

B. Yes

> summary(Mod_pls)

Data: X dimension: 680 6

Y dimension: 680 1 Fit method: kernelpls

Number of components considered: 6

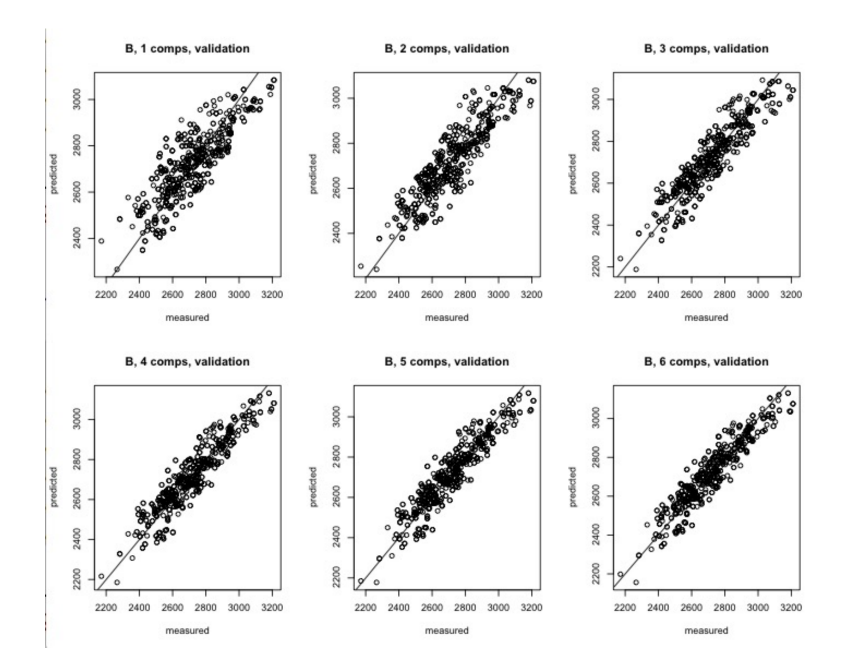
VALIDATION: RMSEP

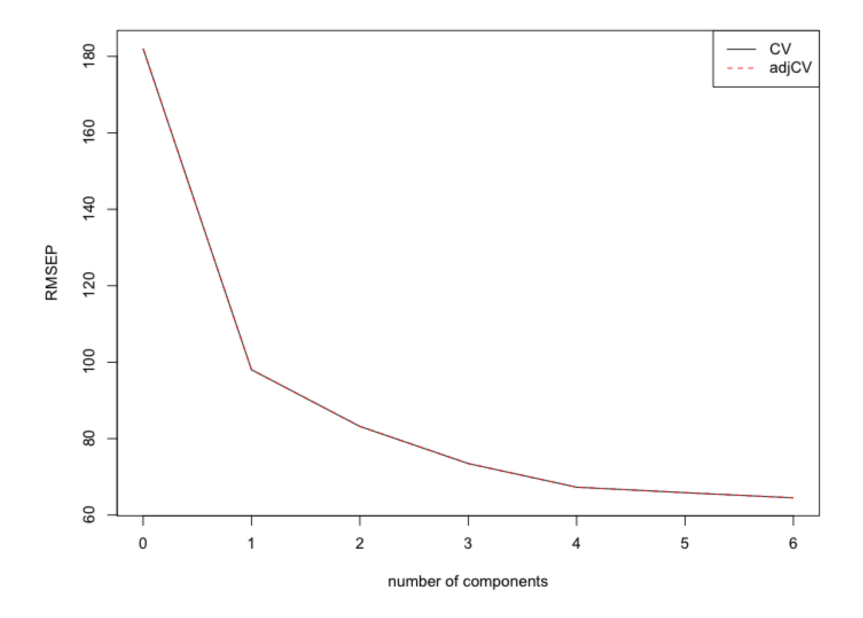
Cross-validated using 680 leave-one-out segments.

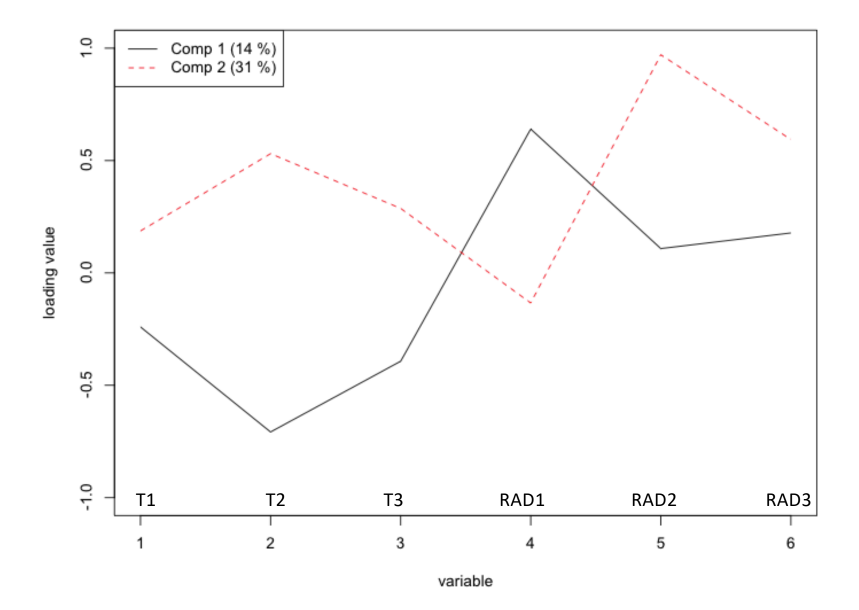
	(Intercept)	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
CV	182.1	98.03	83.19	73.48	67.29	65.85	64.51
adjCV	182.1	98.02	83.20	73.48	67.29	65.85	64.51

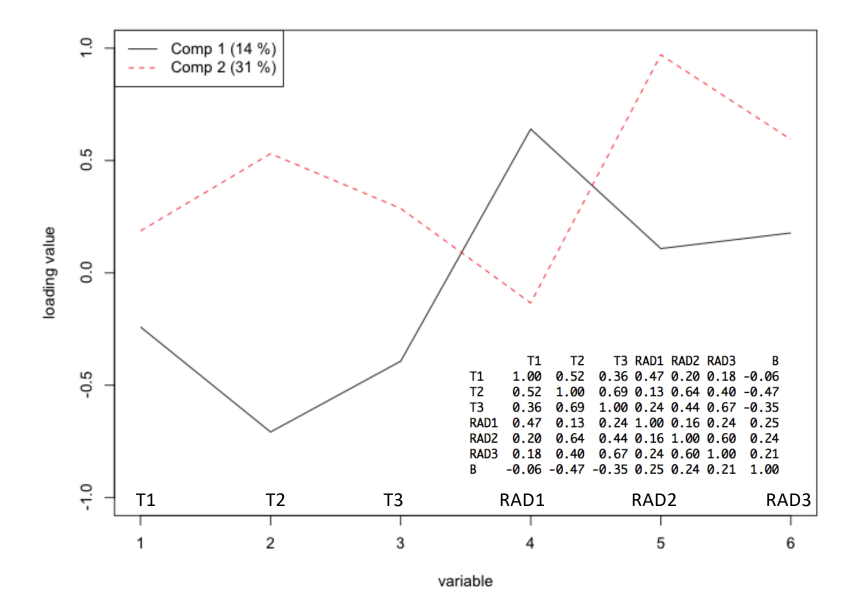
TRAINING: % variance explained

	1 comps	2 comps	3 comps	4 comps	5 comps	6 comps
Χ	13.75	45.09	74.99	89.61	97.43	100.0
В	71.76	79.45	83.97	86.57	87.17	87.7









Several issues

- Inputs X₁, ..., X_P are sometimes (strongly) correlated
- Inputs X₁, ..., X_p may have non-linear effects (unknown response shape),
- Too many inputs
- Need to estimate extreme responses, not mean response

Generalized Additive Model (GAM)

$$E(Y) = \mu + s(X_1) + \dots + s(X_P)$$

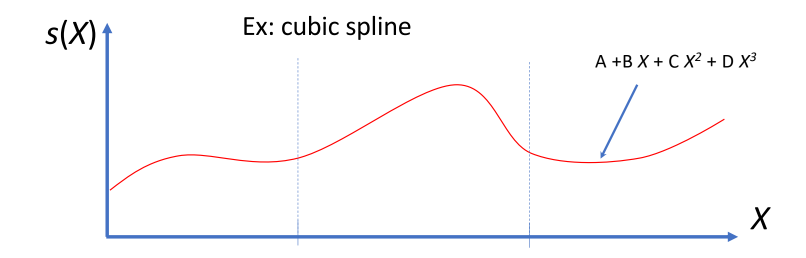
The expected value of Y expressed as a smooth function of the inputs s(X).

- Different types of functions s(X) can be used,
- Advantage of GAM: results easily interpreted.

Generalized Additive Model (GAM)

$$E(Y) = \mu + s(X_1) + \dots + s(X_P)$$

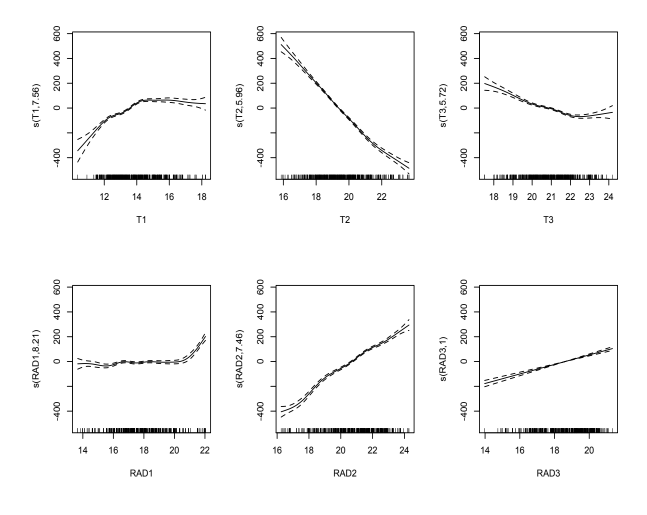
s(X) = spline = piece-wise polynomial function



```
library(mgcv)
```

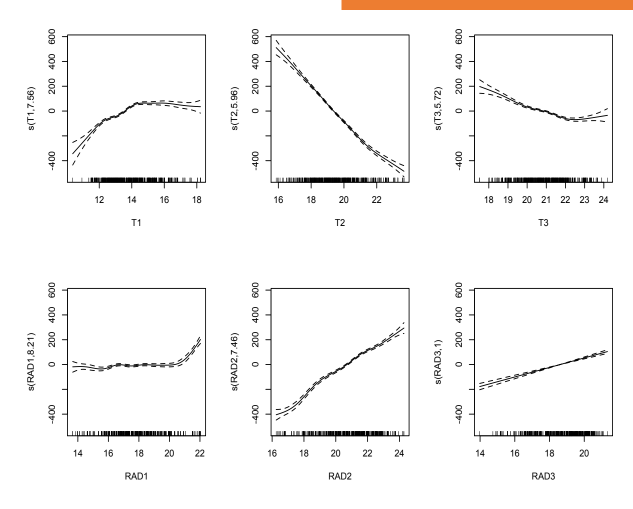
```
Mod_gam<-
gam(B~s(T1)+s(T2)+s(T3)+s(RAD1)+s(RAD2)+s(RAD3),
data=DataSet)
summary(Mod_gam)
plot(Mod_gam)</pre>
```

```
Family: gaussian
Link function: identity
Formula:
B \sim s(T1) + s(T2) + s(T3) + s(RAD1) + s(RAD2) + s(RAD3)
Parametric coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 2718.716 1.944 1399 <2e-16 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' '1
Approximate significance of smooth terms:
         edf Ref.df
                        F p-value
s(T1) 7.563 8.499 53.94 <2e-16 ***
s(T2) 5.962 7.148 282.88 <2e-16 ***
s(T3) 5.720 6.914 29.84 <2e-16 ***
s(RAD1) 8.206 8.826 21.84 <2e-16 ***
s(RAD2) 7.460 8.410 167.23 <2e-16 ***
s(RAD3) 1.000 1.000 197.49 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
R-sq.(adj) = 0.922 Deviance explained = 92.7%
```



Question: Is the effect of T1 on the biomass linear?

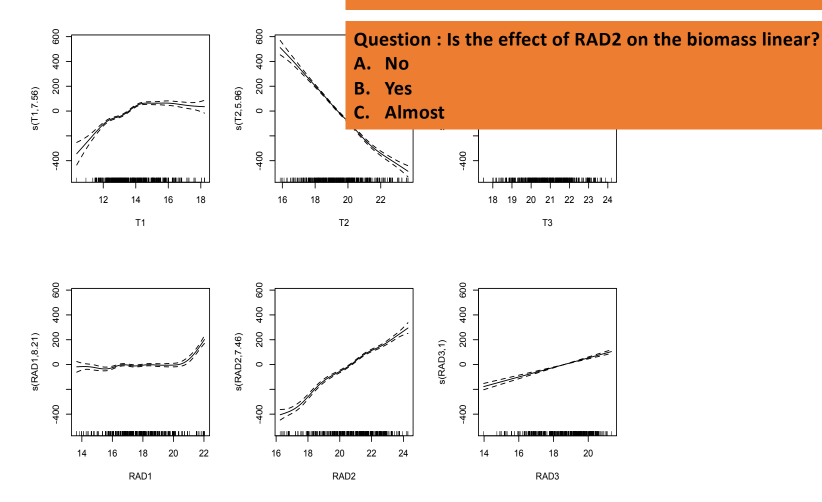
- B. Yes
- C. Almost



Example: maize biom A. No

Question: Is the effect of T1 on the biomass linear?

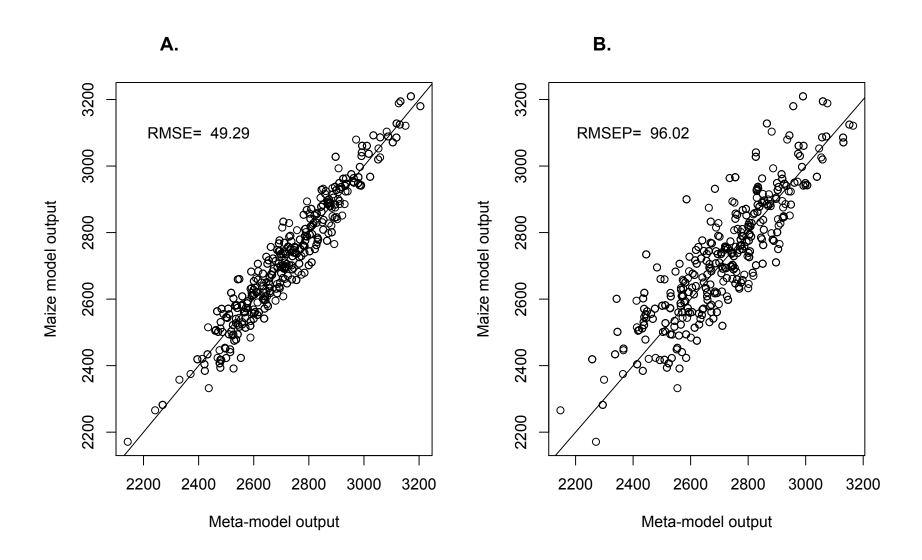
- B. Yes
- C. Almost



```
RMSE_gam<-sqrt(mean((DataSet$B-predict(Mod_gam))^2))
RMSE_gam
#Cross-validation
B_pred_gam<-rep(NA,length(DataSet$B))</pre>
List_year<-unique(DataSet$Year)
for (i in 1:length(List_year))
Training_i<-DataSet[DataSet$Year!=List_year[i],]
Test_i<-DataSet[DataSet$Year==List_year[i],]
Mod_i \leftarrow gam(B \sim s(T1) + s(T2) + s(T3) + s(RAD1) + s(RAD2) + s(RAD3), data=Training_i)
B_gam_i<-predict(Mod_i, newdata=Test_i)</pre>
B_pred_gam[DataSet$Year==List_year[i]]<-B_gam_i</pre>
RMSEP_gam<-sqrt(mean((DataSet$B-B_pred_gam)^2))
RMSEP_gam
```

```
RMSE_gam<-sqrt(mean((DataSet$B-predict(Mod_gam))^2))
RMSE_gam
#Cross-validation
B_pred_gam<-rep(NA,length(DataSet$B))
                                              Question: The training dataset includes
                                              A. All data
List_year<-unique(DataSet$Year)
                                              B. All data but the ith
                                              C. Only the ith data
for (i in 1:length(List_year))
Training_i<-DataSet[DataSet$Year!=List_year[i],]
Test_i<-DataSet[DataSet$Year==List_year[i],]
Mod_i \leftarrow gam(B \sim s(T1) + s(T2) + s(T3) + s(RAD1) + s(RAD2) + s(RAD3), data=Training_i)
B_gam_i<-predict(Mod_i, newdata=Test_i)</pre>
B_pred_gam[DataSet$Year==List_year[i]]<-B_gam_i</pre>
RMSEP_gam<-sqrt(mean((DataSet$B-B_pred_gam)^2))
RMSEP_gam
```

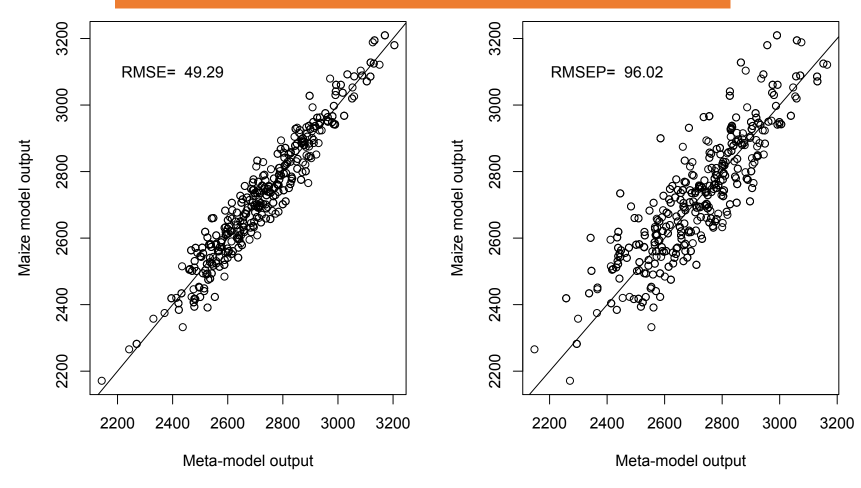
```
RMSE_gam<-sqrt(mean((DataSet$B-predict(Mod_gam))^2))
RMSE_gam
#Cross-validation
B_pred_gam<-rep(NA,length(DataSet$B))
List_year<-unique(DataSet$Year)
                                                                   Question: The test dataset includes
                                                                   A. All data
                                                                   B. All data but the ith
                                                                   C. Only the ith data
for (i in 1:length(List_year))
Training_i<-DataSet[DataSet$Year!=List_year[i],]
Test_i<-DataSet[DataSet$Year==List_year[i],]</pre>
Mod_i \leftarrow gam(B \sim s(T1) + s(T2) + s(T3) + s(RAD1) + s(RAD2) + s(RAD3), data=Training_i)
B_gam_i<-predict(Mod_i, newdata=Test_i)</pre>
B_pred_gam[DataSet$Year==List_year[i]]<-B_gam_i</pre>
RMSEP_gam<-sqrt(mean((DataSet$B-B_pred_gam)^2))
RMSEP_gam
```



Question : Are the prediction errors lower than 50 g/m² in average?

A. Yes

B. No



Several issues

- Inputs X₁, ..., X_P are sometimes (strongly) correlated
- Inputs X_1 , ..., X_p may have non-linear effects (unknown response shape),
- Too many inputs
- Need to estimate extreme responses, not mean response

Other powerful regression techniques: Penalized regression methods

LASSO, ridge and elastic net

$$Y = X\theta + \varepsilon$$
$$\hat{Y} = X\hat{\theta}$$

 $\hat{\theta}$ est estimé en minimisant $\sum_{i} (Y_i - \hat{Y}_i)^2 + \lambda G$ Penalty term

$$G = \sum_{j} |\theta_i|$$
 LASSO

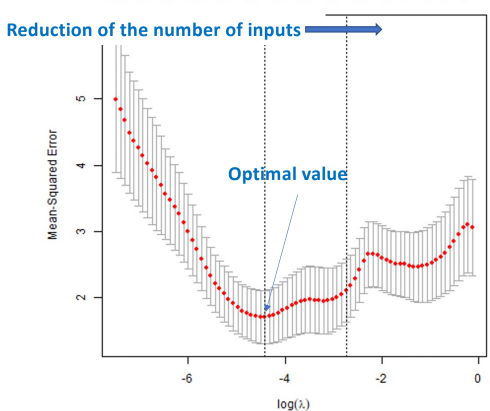
$$G = \sum_{j} \theta_i^2$$
 Ridge

$$G = \sum_{j} \alpha |\theta_{i}| + \sum_{j} (1 - \alpha)\theta_{i}^{2}$$
 Elastic net

Additional parameter (hyper-parameter)

Penalty level (hyper-parameter) optimized by cross validation

32 33 32 33 31 30 29 26 19 18 12 9 7 5 4



Example « spores »

Y=quantity of fungus spores X=34 inputs representing climatic conditions

From Data science pour l'agriculture et l'environnement - Méthodes et applications avec R et Python. Editions Ellipses. https://www.editions-ellipses.fr/

R package glmnet

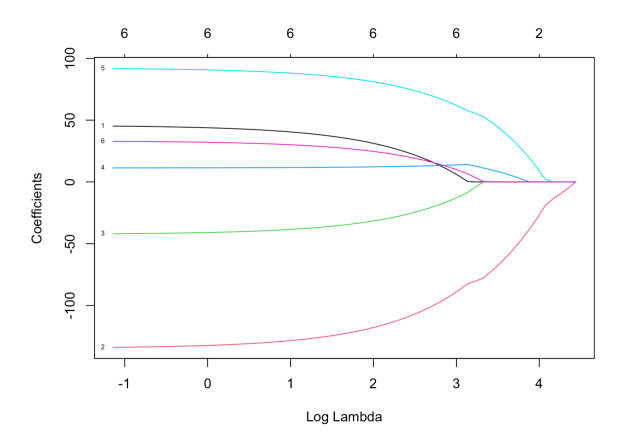
```
cv <- cv.glmnet(x, y, alpha = 1)
model <- glmnet(x, y, alpha = 1, lambda = cv$lambda.min)</pre>
```

library(glmnet)
X=as.matrix(DataSet[,3:8])
Y=DataSet\$B

#Fit LASSO

model <- glmnet(X, Y, alpha = 1)

#Plot coefficients for different penalty levels plot(model, xvar="lambda", label=TRUE)

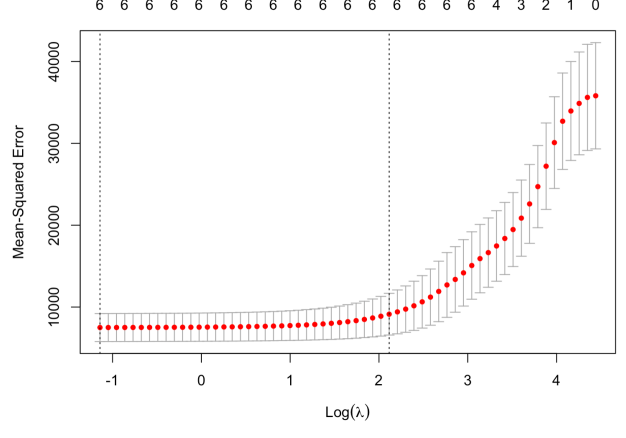


#Regression coefficients for two levels of penalty

```
coef(model, s=exp(4))
coef(model, s=exp(3.5))
```

```
> coef(model, s=exp(4))
7 x 1 sparse Matrix of class "dgCMatrix"
(Intercept) 3050.708642
T1
T2
            -26.696864
T3
RAD1
RAD2
              9.025979
RAD3
> coef(model, s=exp(3.5))
7 x 1 sparse Matrix of class "dgCMatrix"
(Intercept) 2962.356869
T1
T2
            -67.604088
T3
             8.409119
RAD1
RAD2
             44.438812
RAD3
```

#Selection of a lambda value by year-by-year cross-validation



Several issues

- Inputs X₁, ..., X_P are sometimes (strongly) correlated
- Inputs X_1 , ..., X_p may have non-linear effects (unknown response shape),
- Too many inputs
- Need to estimate extreme responses, not mean response

Quantile regression

 Useful to estimate the response of Y to X for upper or lower quantiles of Y

Relevant when using a limited number of inputs, i.e., one or two

Useful for risk analysis

Can be easily implemented with the R package quantreg

```
library(quantreg)  \label{eq:mod} mod <- rq(B^T2+I(T2^2), \ data=DataSet, \ tau=c(0.05,0.1, \ 0.5, \ 0.9, \ 0.95)) \\ print(coef(mod))
```

```
tau= 0.05 tau= 0.10 tau= 0.50 tau= 0.90 tau= 0.95 (Intercept) 2660.507311 1658.472676 247.708516 2892.652836 1967.910435 T2 14.875536 115.001848 295.144193 95.822782 198.461026 I(T2^2) -1.254486 -3.646621 -8.611448 -4.796667 -7.505065
```

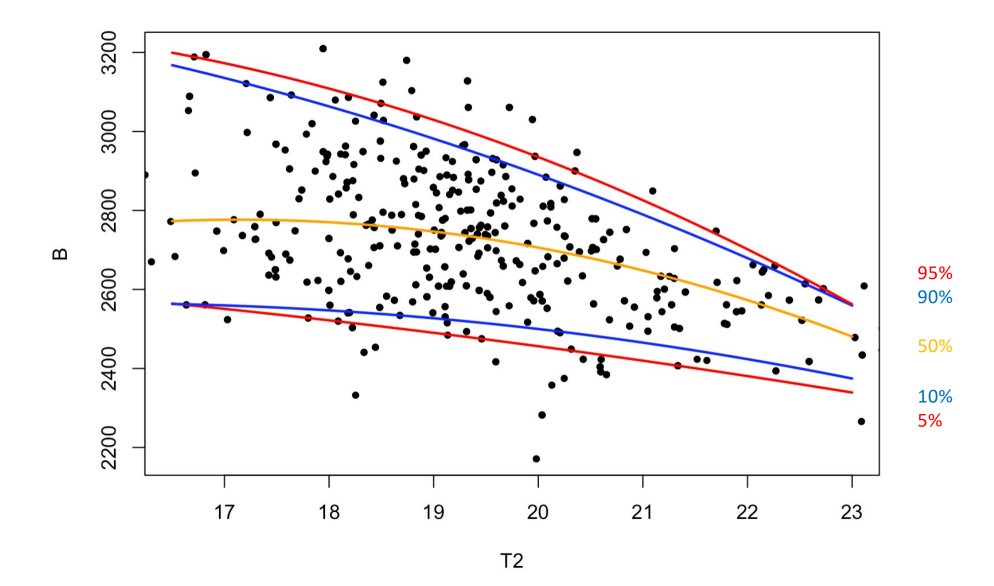
```
Xvec=seq(16.5,23,by=0.1)
Para=coef(mod)[,1]
lines(Xvec,Para[1]+Para[2]*Xvec+Para[3]*Xvec^2, col="red", lwd=2)

Para=coef(mod)[,2]
lines(Xvec,Para[1]+Para[2]*Xvec+Para[3]*Xvec^2, col="blue", lwd=2)

Para=coef(mod)[,3]
lines(Xvec,Para[1]+Para[2]*Xvec+Para[3]*Xvec^2, col="orange", lwd=2)

Para=coef(mod)[,4]
lines(Xvec,Para[1]+Para[2]*Xvec+Para[3]*Xvec^2, col="blue", lwd=2)

Para=coef(mod)[,5]
lines(Xvec,Para[1]+Para[2]*Xvec+Para[3]*Xvec^2, col="red", lwd=2)
```



Conclusion

- How to deal with strongly correlated inputs?
- → PCR, PLSR
- How to analyze the non linear relationships between Y and $X_1, ..., X_P$?
- \rightarrow GAM
- How to obtain accurate predictions with a simplified model?
- → Penalized regressions
- How to analyse responses at upper or lower quantiles?
- → Quantile regression