Statistique bayésienne avec R - exercice régression données Prostate

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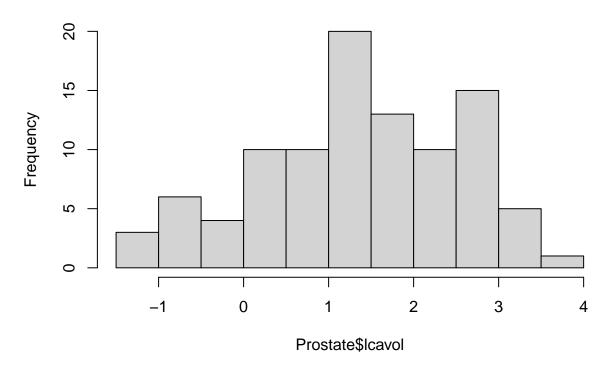
Chargeons les données

library(lasso2)

Un premier examen rapide nous permet de juger que la variable a prédire est approximativement gaussienne, et les variables explicatives sont très corrélées entre elles

hist(Prostate\$lcavol)

Histogram of Prostate\$Icavol



```
cor(Prostate)
            lcavol
                      lweight
                                  age
                                             lbph
## lcavol 1.0000000 0.194128286 0.2249999 0.027349703
                                                  0.53884500 0.675310484
## lweight 0.1941283 1.000000000 0.3075286 0.434934636
                                                  0.10877851
                                                            0.100237795
         0.11765804
                                                            0.127667752
## age
## lbph
         0.0273497 \quad 0.434934636 \quad 0.3501859 \quad 1.000000000 \quad -0.08584324 \quad -0.006999431
## svi
         0.673111185
## lcp
         0.6753105  0.100237795  0.1276678  -0.006999431
                                                  0.67311118 1.000000000
## gleason 0.4324171 -0.001275658 0.2688916 0.077820447
                                                  0.32041222
                                                            0.514830063
         0.4336522 0.050846821 0.2761124 0.078460018 0.45764762
                                                            0.631528245
         ## lpsa
##
              gleason
                         pgg45
          0.432417056 0.43365225 0.7344603
## lcavol
## lweight -0.001275658 0.05084682 0.3541204
          0.268891599 0.27611245 0.1695928
## age
## lbph
          0.077820447 0.07846002 0.1798094
## svi
          0.320412221 0.45764762 0.5662182
## lcp
          0.514830063 0.63152825 0.5488132
## gleason 1.000000000 0.75190451 0.3689868
## pgg45
          0.751904512 1.00000000 0.4223159
          0.368986803 0.42231586 1.0000000
## lpsa
```

Regression linéaire avec sélection de variables

```
modele1=lm(lcavol~.,data=Prostate)
summary(modele1)
##
## Call:
## lm(formula = lcavol ~ ., data = Prostate)
##
## Residuals:
       Min
                 1Q
                      Median
## -1.88603 -0.47346 -0.03987 0.55719 1.86870
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -2.260101
                          1.259683 -1.794
                                             0.0762 .
## lweight
              -0.073166
                          0.174450
                                   -0.419
                                             0.6759
## age
               0.022736
                          0.010964
                                    2.074
                                             0.0410
## lbph
              -0.087449
                          0.058084
                                   -1.506
                                             0.1358
## svi
              -0.153591
                          0.253932 -0.605
                                             0.5468
## lcp
              0.367300
                          0.081689
                                    4.496 2.10e-05 ***
                                     1.236
                                             0.2196
## gleason
              0.190759
                          0.154283
## pgg45
              -0.007158
                          0.004326 - 1.654
                                             0.1016
## lpsa
               0.572797
                          0.085790
                                    6.677 2.11e-09 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.6998 on 88 degrees of freedom
## Multiple R-squared: 0.6769, Adjusted R-squared: 0.6475
## F-statistic: 23.04 on 8 and 88 DF, p-value: < 2.2e-16
```

```
library(car)
## Loading required package: carData
vif(modele1)
## lweight
                        1bph
                                 svi
                                          lcp gleason
                age
                                                         pgg45
                                                                   lpsa
## 1.471496 1.306232 1.392115 2.166568 2.557640 2.433439 2.918987 1.922553
Nous pouvons réaliser une sélection de variables à l'aide d'une procédure step
modele2=step(modele1)
## Start: AIC=-60.7
## lcavol ~ lweight + age + lbph + svi + lcp + gleason + pgg45 +
##
      lpsa
##
##
            Df Sum of Sq
                           RSS
                                   AIC
                 0.0861 43.179 -62.507
## - lweight 1
## - svi
             1
                  0.1792 43.272 -62.299
## - gleason 1 0.7486 43.842 -61.031
## <none>
                        43.093 -60.701
## - lbph 1
                1.1100 44.203 -60.234
## - pgg45 1
                1.3403 44.433 -59.730
          1
## - age
               2.1058 45.199 -58.073
## - lcp
           1 9.9002 52.993 -42.641
## - lpsa
                 21.8300 64.923 -22.946
           1
##
## Step: AIC=-62.51
## lcavol ~ age + lbph + svi + lcp + gleason + pgg45 + lpsa
##
##
            Df Sum of Sq
                           RSS
## - svi
           1 0.1752 43.354 -64.115
## - gleason 1
                  0.8357 44.015 -62.648
## <none>
                         43.179 -62.507
## - pgg45 1
                1.3195 44.499 -61.588
## - lbph 1 1.4818 44.661 -61.234
## - age
           1
                2.0198 45.199 -60.073
          1
## - lcp
                9.8752 53.054 -44.529
          1 23.1542 66.333 -22.862
## - lpsa
##
## Step: AIC=-64.11
## lcavol ~ age + lbph + lcp + gleason + pgg45 + lpsa
##
            Df Sum of Sq
##
                           RSS
                                   AIC
## <none>
                         43.354 -64.115
## - gleason 1
                  0.9571 44.311 -63.997
                 1.3338 44.688 -63.175
## - lbph
            1
## - pgg45
             1
                1.4298 44.784 -62.967
                  1.9355 45.290 -61.878
## - age
             1
## - lcp
             1
                 10.9352 54.289 -44.297
## - lpsa
             1
                 24.9001 68.254 -22.093
summary(modele2)
```

##

```
## Call:
## lm(formula = lcavol ~ age + lbph + lcp + gleason + pgg45 + lpsa,
       data = Prostate)
##
## Residuals:
##
       Min
                  1Q
                     Median
                                    3Q
                                            Max
## -1.90430 -0.51715 -0.02241 0.57388 1.87347
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -2.535920
                           1.111608 -2.281
                                              0.0249 *
                           0.010598
                                     2.004
                                              0.0480 *
                0.021243
               -0.088505
## lbph
                           0.053188 -1.664
                                              0.0996 .
## lcp
                0.344851
                           0.072379
                                     4.765 7.22e-06 ***
                           0.150234
                                      1.410
## gleason
                0.211762
                                              0.1621
## pgg45
               -0.007353
                           0.004268
                                     -1.723
                                              0.0884 .
                0.544422
                           0.075723
                                     7.190 1.84e-10 ***
## lpsa
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.6941 on 90 degrees of freedom
## Multiple R-squared: 0.6749, Adjusted R-squared: 0.6532
## F-statistic: 31.14 on 6 and 90 DF, \, p-value: < 2.2e-16
La variable gleason n'étant pas significative dans le modèle obtenu par la sélection de variables (step), je la
supprime.
modele3=lm(lcavol~age+lbph+lcp+pgg45+lpsa,data=Prostate)
summary(modele3)
##
## Call:
## lm(formula = lcavol ~ age + lbph + lcp + pgg45 + lpsa, data = Prostate)
##
## Residuals:
##
                  1Q
                       Median
## -1.97761 -0.49431 -0.04114 0.55594
                                        2.00901
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -1.299521
                           0.686526 - 1.893
                                              0.0616 .
## age
                0.022662
                           0.010607
                                      2.137
                                              0.0353 *
## lbph
                           0.053473 -1.669
               -0.089254
                                              0.0985
               0.350498
                                     4.824 5.63e-06 ***
## lcp
                           0.072659
               -0.003649
                           0.003381 - 1.079
                                              0.2834
## pgg45
## lpsa
                0.549946
                           0.076031
                                     7.233 1.43e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.6978 on 91 degrees of freedom
## Multiple R-squared: 0.6677, Adjusted R-squared: 0.6495
## F-statistic: 36.57 on 5 and 91 DF, p-value: < 2.2e-16
modele3=lm(lcavol~age+lbph+lcp+lpsa,data=Prostate)
summary(modele3)
```

```
##
## Call:
## lm(formula = lcavol ~ age + lbph + lcp + lpsa, data = Prostate)
## Residuals:
##
                      Median
       Min
                 1Q
                                   3Q
## -1.92180 -0.49945 -0.09019 0.58214 2.01844
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -1.21022
                          0.68213 -1.774
               0.02003
                          0.01033
                                    1.939
                                            0.0556 .
## lbph
              -0.08935
                          0.05352
                                   -1.669
                                            0.0984 .
                                    5.006 2.67e-06 ***
## lcp
               0.30907
                          0.06175
## lpsa
               0.54273
                          0.07580
                                    7.160 1.93e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.6984 on 92 degrees of freedom
## Multiple R-squared: 0.6635, Adjusted R-squared: 0.6488
## F-statistic: 45.35 on 4 and 92 DF, p-value: < 2.2e-16
```

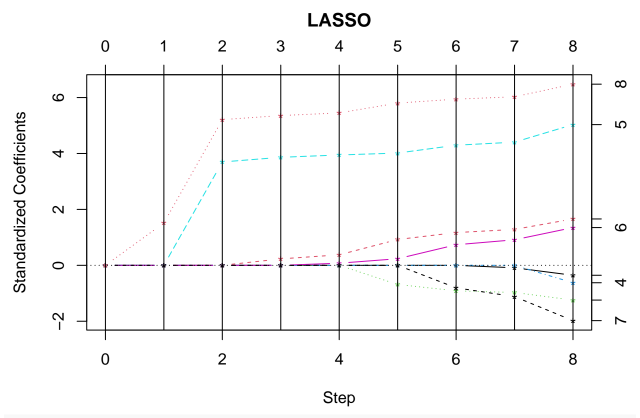
Regression PCR / PLS

```
library(pls)
## Attaching package: 'pls'
## The following object is masked from 'package:stats':
##
##
       loadings
modele5=pcr(lcavol~.,data=Prostate,validation='L00',scale=TRUE)
summary(modele5)
## Data:
           X dimension: 97 8
## Y dimension: 97 1
## Fit method: svdpc
## Number of components considered: 8
##
## VALIDATION: RMSEP
## Cross-validated using 97 leave-one-out segments.
##
          (Intercept) 1 comps 2 comps 3 comps 4 comps 5 comps
                                                                    6 comps
## CV
                       0.8469
                                 0.8495
                1.185
                                          0.8126
                                                   0.8189
                                                            0.8362
                                                                     0.7921
## adiCV
                1.185
                        0.8468
                                 0.8493
                                          0.8124
                                                   0.8187
                                                            0.8361
                                                                     0.7918
          7 comps 8 comps
## CV
           0.7286
                    0.7315
## adjCV
          0.7283
                    0.7312
##
## TRAINING: % variance explained
           1 comps 2 comps 3 comps 4 comps 5 comps 6 comps 7 comps 8 comps
             40.89
## X
                      61.69
                               74.79
                                        82.79
                                                 88.86
                                                          93.82
                                                                   97.21
                                                                           100.00
## lcavol
            50.51
                      50.90
                               55.99
                                        56.20
                                                 56.20
                                                          60.89
                                                                    66.86
                                                                             67.69
```

```
modele6=plsr(lcavol~.,data=Prostate,validation='LOO',scale=TRUE)
summary(modele6)
           X dimension: 97 8
## Data:
## Y dimension: 97 1
## Fit method: kernelpls
## Number of components considered: 8
##
## VALIDATION: RMSEP
## Cross-validated using 97 leave-one-out segments.
         (Intercept) 1 comps 2 comps 3 comps 4 comps 5 comps 6 comps
## CV
               1.185 0.8105 0.7614 0.7369
                                                 0.7326
                                                          0.7302
                                                                   0.7312
## adjCV
               1.185
                       0.8104
                              0.7610 0.7365
                                                 0.7323
                                                          0.7299
                                                                   0.7309
##
         7 comps 8 comps
## CV
          0.7316
                   0.7315
## adjCV
          0.7313
                   0.7312
##
## TRAINING: % variance explained
##
          1 comps 2 comps 3 comps 4 comps 5 comps 6 comps 7 comps
## X
            40.48
                     52.21
                              62.07
                                       78.46
                                                83.72
                                                        90.76
                                                                 93.94
                                                                         100.00
## lcavol
            55.85
                     64.04
                              67.10
                                       67.56
                                                67.68
                                                         67.69
                                                                 67.69
                                                                          67.69
```

Regression LASSO

```
library('lars')
## Loaded lars 1.3
model_lasso=lars(as.matrix(Prostate[,-1]),Prostate$lcavol,type="lasso",trace=F,normalize=TRUE)
plot(model_lasso,xvar='step', plottype='coeff')
```

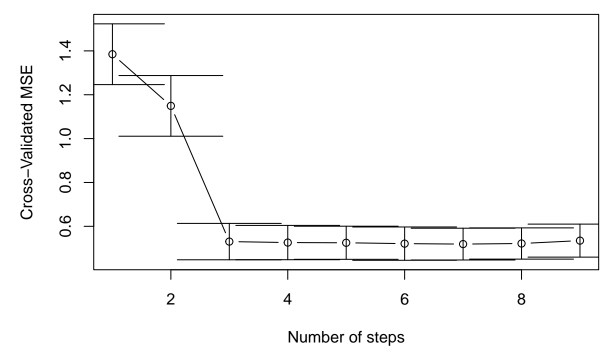


```
print(model_lasso)
```

```
##
## Call:
\#\# lars(x = as.matrix(Prostate[, -1]), y = Prostate$lcavol, type = "lasso",
       trace = F, normalize = TRUE)
## R-squared: 0.677
## Sequence of LASSO moves:
        lpsa lcp age gleason lbph pgg45 lweight svi
## Var
           8
               5
                   2
                           6
                                3
                                      7
                                               1
               2
                                5
                                       6
                                               7
## Step
           1
                           4
```

Cherchons le lambda optimal

```
cv=cv.lars(as.matrix(Prostate[,-1]),Prostate$lcavol,K=10,trace=F,plot.it=T,se=T,type=c("lasso"),mode='s
```



meilleur CVMSE semble être à la troisième étape.

On peut examiner les valeurs des variables sélectionnées à la troisième étape

```
print(model_lasso$lambda[3])
```

Le

```
## [1] 1.242805
```

```
print(model_lasso$beta[3,])
```

```
## lweight age lbph svi lcp gleason pgg45 lpsa
## 0.0000000 0.0000000 0.0000000 0.2698036 0.0000000 0.0000000 0.4606733
```

Ré-estimons le modèle avec les variables sélectionnées par le LASSO

```
modele4=lm(lcavol~lpsa+lcp,data=Prostate)
summary(modele4)
```

```
##
## lm(formula = lcavol ~ lpsa + lcp, data = Prostate)
##
## Residuals:
##
       Min
                  1Q
                       Median
                                    30
                                             Max
##
   -1.65744 -0.54398 -0.05502 0.57163
                                        2.07959
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                0.09135
                           0.20527
                                      0.445
                                               0.657
                           0.07501
## lpsa
                0.53162
                                      7.087 2.49e-10 ***
## 1cp
                0.32837
                           0.06193
                                      5.303 7.54e-07 ***
## ---
                     '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 0.7092 on 94 degrees of freedom
## Multiple R-squared: 0.6455, Adjusted R-squared: 0.6379
```

```
## F-statistic: 85.57 on 2 and 94 DF, p-value: < 2.2e-16
On peut finalement comparer les modèles suivant le critère de notre choix (AdjR2,AIC,BIC).
library(broom)
rbind(glance(modele1),glance(modele2),glance(modele3),glance(modele4))
## # A tibble: 4 x 12
    r.squared adj.r.squa~1 sigma stati~2 p.value
                                                                   AIC
                                                                         BIC devia~3
##
                                                       df logLik
##
         <dbl>
                      <dbl> <dbl>
                                    <dbl>
                                              <dbl> <dbl> <dbl> <dbl> <dbl> <
         0.677
                      0.647 0.700
                                                        8 -98.3 217.
## 1
                                     23.0 1.34e-18
                                                                        242.
                                                                                 43.1
## 2
                      0.653 0.694
                                     31.1 5.52e-20
                                                       6 -98.6 213.
         0.675
                                                                        234.
                                                                                43.4
## 3
         0.663
                      0.649 0.698
                                     45.3 5.51e-21
                                                        4 -100.
                                                                  213.
                                                                        228.
                                                                                 44.9
         0.645
                                                        2 -103.
## 4
                      0.638 0.709
                                     85.6 6.81e-22
                                                                  214.
                                                                        224.
                                                                                 47.3
## # ... with 2 more variables: df.residual <int>, nobs <int>, and abbreviated
      variable names 1: adj.r.squared, 2: statistic, 3: deviance
Le meilleur modèle suivant les critères AIC et R2ajusté est le modèle 4, avec les 2 variables lpsa et lcp
On pourrait aussi implétenter une validation croisée
library(lmvar)
modele1=lm(lcavol~.,data=Prostate,x=TRUE,y=TRUE)
modele2=step(modele1,trace = FALSE)
modele3=lm(lcavol~age+lbph+lcp+lpsa,data=Prostate,x=TRUE,y=TRUE)
modele4=lm(lcavol~lpsa+lcp,data=Prostate,x=TRUE,y=TRUE)
cv.lm(modele1)
## Mean absolute error
                                 0.6009087
## Sample standard deviation :
                                 0.1277847
##
## Mean squared error
                              : 0.5246905
## Sample standard deviation
                             : 0.1908469
##
## Root mean squared error
                              : 0.7104631
## Sample standard deviation : 0.1488202
cv.lm(modele2)
## Mean absolute error
                              : 0.5904426
## Sample standard deviation :
                                 0.1667396
##
## Mean squared error
                              : 0.5251093
## Sample standard deviation :
                                 0.2646416
##
## Root mean squared error
                                 0.7050636
## Sample standard deviation : 0.1763666
cv.lm(modele3)
## Mean absolute error
                              : 0.6018834
## Sample standard deviation : 0.1134667
##
## Mean squared error
                              : 0.5293942
## Sample standard deviation : 0.1924361
##
## Root mean squared error
                              : 0.7171432
## Sample standard deviation : 0.1295279
```

cv.lm(modele4)

Mean absolute error : 0.5834911 ## Sample standard deviation : 0.1224547

##

Mean squared error : 0.5031042
Sample standard deviation : 0.1777004

##

Root mean squared error : 0.6982231
Sample standard deviation : 0.1316084

Le meilleur modèle suivant le critère de validation croisée est le modèle 4, avec les 3 variables séléctionnées par LASSO