Comparison Of Two Tests

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```
library(simstudy)
library(dplyr)
library(caret)
library(lmtest)
library(knitr)
```

Define leave one out cross validation prediction error function

Leave one out cross validation prediction error is evaludated by loocv() function, which is built based on

$$LOOCV = \frac{1}{n} \sum_{i=1}^{n} \frac{(Y_i - \hat{Y}_i)^2}{(1 - h_{ii})^2}$$

```
#### leave one out cross validation prediction error function using trick
loocv=function(fit){
   h=lm.influence(fit)$h
   mean((residuals(fit)/(1-h))^2)
}
```

Define likehood ratio test function

(1) First step: Evaluate the maximum likelihood estimation of σ^2 , that is

$$\hat{\sigma_{\alpha}^2} = \frac{1}{n} RSS_{\alpha} = \frac{1}{n} (Y - X_{\alpha} \hat{\beta_{\alpha}})^T (Y - X_{\alpha} \hat{\beta_{\alpha}})$$

for any certain model M_{α} . Let's call this funtion 'sd_mle()'.

(2) Second step: Assuming normality, the maximum log-likelihood value for model α , $l(\hat{\beta}_{\alpha}, \hat{\sigma}_{\alpha}^2)$ say, for the full-rank regression model is the log probability density of Y with maximum likelihood estimation as parameters, namely,

$$\begin{split} l(\hat{\beta_{\alpha}}, \hat{\sigma_{\alpha}^2}) &= log \bigg\{ (2\pi \hat{\sigma_{\alpha}^2})^{-\frac{n}{2}} exp \bigg\{ -\frac{1}{2\hat{\sigma_{\alpha}^2}} ||Y - X_{\alpha} \hat{\beta_{\alpha}}||^2 \bigg\} \bigg\} \\ &= -\frac{n}{2} log (2\pi \hat{\sigma_{\alpha}^2}) - \frac{1}{2\hat{\sigma_{\alpha}^2}} ||Y - X_{\alpha} \hat{\beta_{\alpha}}||^2 \\ &= \sum_{i=1}^n -\frac{1}{2} log (2\pi \hat{\sigma_{\alpha}^2}) - \frac{1}{2\hat{\sigma_{\alpha}^2}} (Y_i - X_{i,\alpha} \hat{\beta_{\alpha}})^2 \\ &= \sum_{i=1}^n dnorm(Y_i; X_{i,\alpha} \hat{\beta_{\alpha}}, \hat{\sigma_{\alpha}^2}) \end{split}$$

, where l(.,.) is the log-likelihood function. Let's call this funtion 'ml()'.

(3) Third step: Evaluate log likelihood ratio test statistic according to definition, namely,

$$LLR = -2(l(\hat{\beta_{\alpha}}, \hat{\sigma_{\alpha}^2}) - l(\hat{\beta_{\lambda}}, \hat{\sigma_{\lambda}^2}))$$

, where we assume $M_{\alpha} \subset M_{\lambda}$.Let's call this funtion 'lrt()'.

```
sd_mle <- function(object) {
    sqrt(mean(residuals(object)^2))
}

ml <- function(object) {
    sum(dnorm(model.response(model.frame(object)),
        mean = fitted(object), sd = sd_mle(object), log = TRUE))
}

lrt <- function(model1,model2) {
    -2 * (ml(model1) - ml(model2))
}</pre>
```

Define A Full Rank Dataset

The true linear model is y=3+2x1+5x2+3x3+1.5x4+e, which is the full model.

```
#### Define A Full Rank Dataset

def <- defData(varname = "x1", dist="uniform",formula = "10;20") ## x1 is from unifrom distribution

def <- defData(def,varname = "x2", dist="uniform",formula = "0;3")

def <- defData(def,varname = "x3", dist="uniform",formula = "0;5")

def <- defData(def,varname = "x4", dist="uniform",formula = "5;10")

def <- defData(def, varname = "y", formula = "3+2*x1+5*x2+3*x3+1.5*x4", variance = 1)

## The true linear model is y=3+2*x1+5*x2+3*x3+1.5*x4+e</pre>
```

Generate a dataset with size 1000

Here I wanted to do a single time simulation to see exactly what values the test statistics are under the alternative hypothesis, i.e., under full model. In this simulation, I generated a dataset of size 1000 and fit model 1~4, respectively.

To test the power of a certain test, basically we set the null hypothesis as model 1~3 respectively, because they are all incorrect (insufficient) model compared with the model under alternative hypothesis, which we set as the ture model. The goal is to see exactly to what extent can we reject the null hypothesis in each test.

```
md < -c()
n = 1000
```

```
dt <- genData(n, def)
dt <- dt%>%select(y,x1,x2,x3,x4) ##generate dataset n=1000
fit1 <- lm(y~ x1, data = dt)
fit2 <- lm(y~ x1+x2, data = dt)
fit3 <- lm(y~ x1+x2+x3, data = dt)
fit4 <- lm(y~ x1+x2+x3+x4, data = dt)
md<-c(loocv(fit1),loocv(fit2),loocv(fit3),loocv(fit4)) ##loocv prediction error for each model</pre>
```

Calculate the test statistic in one time simulation

Evaluate each test statistic based on

New Test Statistic =
$$(\frac{n}{\sigma^2}\hat{\Gamma}_{\alpha,n} - 2d_{\alpha}) - (\frac{n}{\sigma^2}\hat{\Gamma}_{\lambda,n} - 2d_{\lambda})$$
, and
$$LLR = -2(l(\hat{\beta}_{\alpha}, \hat{\sigma}_{\alpha}^2) - l(\hat{\beta}_{\lambda}, \hat{\sigma}_{\lambda}^2))$$

and compare the corresponding value of two test statistics.

```
ts1 = (n*md[1]-2*1) - (n*md[4]-2*4)

ts2 = (n*md[2]-2*2) - (n*md[4]-2*4)

ts3 = (n*md[3]-2*3) - (n*md[4]-2*4)

lr1 = lrt(fit1,fit4)

lr2 = lrt(fit2,fit4)

lr3 = lrt(fit3,fit4)

ts1;lr1;ts2;lr2;ts3;lr3

## [1] 40661.32

## [1] 3759.252

## [1] 24493.39

## [1] 3265.798

## [1] 4579.586

## [1] 1740.793
```

Now do the same simulation once again, only with sample size of 5000 this time. ## Generate a dataset with size 5000

```
md < -c()
n = 5000
```

```
dt <- genData(n, def)
dt <- dt%>%select(y,x1,x2,x3,x4) ##generate dataset n=1000
fit1 <- lm(y~ x1, data = dt)
fit2 <- lm(y~ x1+x2, data = dt)
fit3 <- lm(y~ x1+x2+x3, data = dt)
fit4 <- lm(y~ x1+x2+x3+x4, data = dt)
md<-c(loocv(fit1),loocv(fit2),loocv(fit3),loocv(fit4)) ##loocv prediction error for each model</pre>
```

Calculate the test statistic in one time simulation

```
ts1 = (n*md[1]-2*1) - (n*md[4]-2*4)

ts2 = (n*md[2]-2*2) - (n*md[4]-2*4)

ts3 = (n*md[3]-2*3) - (n*md[4]-2*4)

lr1 = lrt(fit1,fit4)

lr2 = lrt(fit2,fit4)

lr3 = lrt(fit3,fit4)

ts1;lr1;ts2;lr2;ts3;lr3

## [1] 212415.4

## [1] 18906.95

## [1] 117538

## [1] 16037.23

## [1] 23609.33

## [1] 8756.283
```

Interpretation: from the test statistics value above, we can see that in this one-time simulation, our new test statistics are much greater than their corresponding traditional maximum likelihood ratio test statistics, which implies

that our new test enjoys a much higher power. Besides, the difference becomes larger as the difference between dimensions and sample size n become larger.

1000 times simulations

Now let's make 1000 times simulations, in each time to see whether our new test statistic is greater than its counterpart log likelihood ratio test statistic. Since they share the same criterion, a larger value implies a greater power of the test.

```
df1 \leftarrow c()
df2 <- c()
df3 \leftarrow c()
for(i in 1:1000){
  dt <- genData(100, def)%>%select(y,x1,x2,x3,x4)
  ##generate dataset n=100
  ##The true linear model is y=3+2*x1+5*x2+3*x3+1.5*x4+e
  fit1 <- lm(y \sim x1, data = dt)
 fit2 <- lm(y \sim x1+x2, data = dt)
  fit3 <- lm(y \sim x1+x2+x3, data = dt)
  fit4 <- lm(y ~ x1+x2+x3+x4, data = dt)
  md<-c(loocv(fit1),loocv(fit2),loocv(fit3),loocv(fit4))</pre>
  ts1 = (n*md[1]-2*1) - (n*md[4]-2*4) ##new test statistics
  ts2 = (n*md[2]-2*2) - (n*md[4]-2*4)
  ts3 = (n*md[3]-2*3) - (n*md[4]-2*4)
  lr1 = lrt(fit1,fit4)
  lr2 = lrt(fit2,fit4)
 lr3 = lrt(fit3,fit4)
 df1 <- c(df1,ifelse (ts1 > lr1, 1, 0 ) )
df2 <- c(df2,ifelse (ts2 > lr2, 1, 0 ) )
df3 <- c(df3,ifelse (ts3 > lr3, 1, 0 ) )
## In each simulation record if new test statistics is larger than traditional maximum likelihood ratio
}
```

Show the result

```
table(df1)

## df1
## 1
## 1000

table(df2)
## df2
```

```
## 1
## 1000
```

table(df3)

df3 ## 1 ## 1000

Interpretation: from the result, we can see that in this 1000 times mulations, our new test statistics are greater than traditional maximum likelihood ratio test statistics each single time, which implies that the new test has the same level as maximum likelihood ratio test while enjoys a higher power and its performance is very stable. So it can beat likelihood ratio test on power.