All subset regression with leaps, bestglm, glmulti, and meifly

```
## Settings for RMarkdown
http://yihui.name/knitr/options#chunk_options
opts_chunk$set(comment = "", warning = FALSE, message = FALSE,
tidy = FALSE,
    echo = T, fig.width = 5, fig.height = 5)
options(width = 100, scipen = 5, digits = 5)
setwd("~/statistics/Rmedstats/")
```

Summary

- For linear regression, use **leaps**, which allows use of adjusted \(R^2 \) and Mallow Cp.
- For logistic regression, use **glmulti**.
- For Cox regression, use **glmulti**.
- This article about glmulti is a good summary of this topic (http://www.jstatsoft.org/v34/i12/paper).

Load and prepare dataset

http://www.umass.edu/statdata/statdata/data/lowbwt.txt

SOURCE: Hosmer and Lemeshow (2000) Applied Logistic Regression: Second Edition. These data are copyrighted by John Wiley & Sons Inc. and must be acknowledged and used accordingly. Data were collected at Baystate Medical Center, Springfield, Massachusetts during 1986.

```
library(gdata)
lbw <-
read.xls("http://www.umass.edu/statdata/statdata/lowbwt.xls")
names(lbw) <- tolower(names(lbw))

## Recoding
lbw <- within(lbw, {
    ## race relabeling
    race.cat <- factor(race, levels = 1:3, labels =
c("white","Black","other"))

    ## ftv (frequency of visit) relabeling
    ftv.cat <- cut(ftv, breaks = c(-Inf, 0, 2, Inf), labels =
c("None","Normal","Many"))
    ftv.cat <- relevel(ftv.cat, ref = "Normal")

    ## ptl
    preterm <- factor(ptl >= 1, levels = c(F,T), labels =
c("0","1+"))
})
```

leaps (regression subset selection)

Regression subset selection including exhaustive search. This is only for linear regression.

Reference: http://www.statmethods.net/stats/regression.html

Perform all subset regression, and choose "nbest" model(s) for each number of predictors up to nvmax.

The result shows how it was performed.

```
Subset selection object
Call: regsubsets.formula(bwt ~ age + lwt + race.cat + smoke +
preterm +
    ht + ui + ftv.cat, data = lbw, nbest = 1, nvmax = NULL,
force.in = NULL,
    force.out = NULL, method = "exhaustive")
10 Variables
             (and intercept)
              Forced in Forced out
age
                  FALSE
                              FALSE
lwt
                              FALSE
                  FALSE
race.catBlack
                  FALSE
                              FALSE
race.catOther
                  FALSE
                              FALSE
smoke
                  FALSE
                              FALSE
preterm1+
                  FALSE
                              FALSE
ht
                              FALSE
                  FALSE
ui
                  FALSE
                              FALSE
ftv.catNone
                              FALSE
                  FALSE
ftv.catMany
                  FALSE
                              FALSE
1 subsets of each size up to 10
Selection Algorithm: exhaustive
```

Best model at each variable number

The best model in the 10-variable case includes all variables, as that is the only way to have 10 variables.

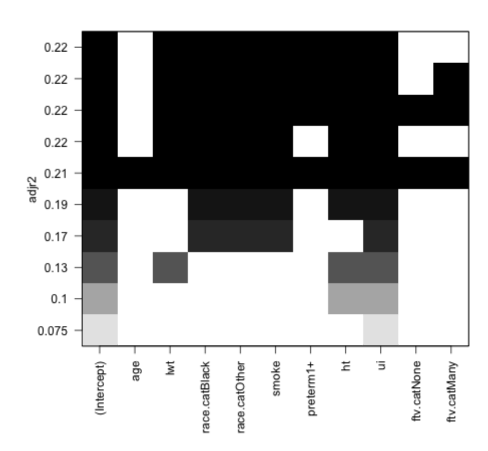
```
summary.out <- summary(regsubsets.out)
as.data.frame(summary.out$outmat)</pre>
```

```
age lwt race.catBlack race.catOther smoke preterm1+
ht ui ftv.catNone ftv.catMany
   (1)
2
   (1)
3
   (1)
*
                               *
                                                     *
4
   (1)
5
                                                     *
   (1)
6
   (1)
                 *
                               *
                                              *
7
                               *
                                                     *
                                                               *
   (1)
   (1)
8
                                              *
                                                     *
                                                               *
9
   (1)
                 *
                                                               *
                *
  *(1)
                                                               *
10
               *
                            *
```

By adjusted \(R^2 \), the best model includes lwt, race.cat, preterm, ht, and ui (variables that have black boxes at the higest Y-axis value).

```
## Adjusted R2
plot(regsubsets.out, scale = "adjr2", main = "Adjusted R^2")
```

Adjusted R²

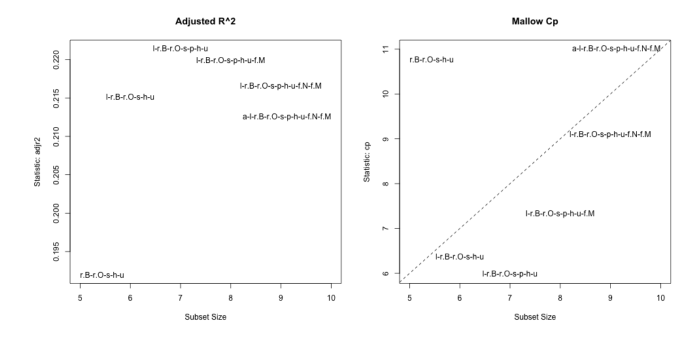


Plot Output from regsubsets Function in leaps package

This is just another way of presenting the same information for adjusted \(R^2 \). The model with 7 variables (counting dummy variables seprately) has the highest adjusted \(R^2 \).

Mallow Cp is used to decide on the number of predictors to include. The stopping rule is to start with the smallest model and gradually increase number of variables, and stop when Mallow Cp is approximately (number of regressors + 1, broken line) for the first time. In this case, the model with 6 regressors is the first one to achieve such a condition.

```
library(car)
layout(matrix(1:2, ncol = 2))
## Adjusted R2
res.legend <-
    subsets(regsubsets.out, statistic="adjr2", legend = FALSE,
min.size = 5, main = "Adjusted R^2")
## Mallow Cp
res.legend <-
    subsets(regsubsets.out, statistic="cp", legend = FALSE,
min.size = 5, main = "Mallow Cp")
abline(a = 1, b = 1, lty = 2)</pre>
```



```
res.legend
```

```
Abbreviation
age
                            a
lwt
race.catBlack
                          r.B
race.catOther
                          r.o
smoke
                            S
preterm1+
                            p
ht
                            h
                            u
ftv.catNone
                          f.N
                          f.M
ftv.catMany
```

See which model has the highest adjusted R2

The model with 7 variables (counting dummy variables separately) has the highest adjusted $\ (R^2)$. Variables marked with TRUE are the ones chosen.

```
which.max(summary.out$adjr2)
```

[1] 7

```
summary.out$which[7,]
```

```
lwt race.catBlack
  (Intercept)
                          age
race.catOther
                       smoke
                                  preterm1+
         TRUE
                       FALSE
                                        TRUE
                                                       TRUE
TRUE
               TRUE
                              TRUE
                           ui
                                ftv.catNone
                                               ftv.catMany
            ht
                                                      FALSE
         TRUE
                        TRUE
                                       FALSE
```

Do regression with the best model

Somehow I had to recreate the best model from the output above.

```
best.model <- lm(bwt ~ lwt + race.cat + smoke + preterm + ht +
ui, data = lbw)
summary(best.model)</pre>
```

```
call:
lm(formula = bwt ~ lwt + race.cat + smoke + preterm + ht + ui,
    data = 1bw
Residuals:
                 Median
    Min
             1Q
                              3Q
                                     Max
                           494.2
-1886.4
        -441.0
                    53.5
                                  1620.9
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                                              <2e-16 ***
(Intercept)
               2871.99
                            243.67
                                     11.79
lwt
                  4.04
                              1.67
                                      2.42
                                              0.0167 *
race.catBlack
               -466.32
                            145.13
                                     -3.21
                                              0.0016 **
               -335.68
                                              0.0032 **
                            112.28
                                     -2.99
race.catOther
               -323.57
                            104.96
                                     -3.08
                                             0.0024 **
smoke
                                     -1.56
                            133.50
preterm1+
               -208.44
                                             0.1202
                                             0.0044 **
ht
               -573.69
                            198.96
                                     -2.88
               -489.96
                            135.93
                                     -3.60
                                             0.0004 ***
ui
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 643 on 181 degrees of freedom
Multiple R-squared: 0.25,
                           Adjusted R-squared: 0.221
F-statistic: 8.64 on 7 and 181 DF, p-value: 0.00000000396
```

bestglm (Best subset GLM)

Best subset glm using AIC, BIC, EBIC, BICq or Cross-Validation. For the normal case, the 'leaps' is used. Otherwise, a slower exhaustive search. The 'xtable' package is needed for vignette 'SimExperimentBICq.Rnw' accompanying this package.

References:

- http://cran.r-project.org/web/packages/bestglm/
- http://cran.r-project.org/web/packages/bestglm/vignettes/bestglm.pdf

Load bestglm

```
library(bestglm)
```

Reformat data

The outcome variable must be named y, no extraneous variables should be present in the dataset.

```
lbw.for.bestqlm <- within(lbw, {</pre>
    id
         <- NULL
                         # Delete
    low
         <- NULL
    race <- NULL
    ptl
         <- NULL
    ftv <- NULL
                      # bwt into y
# Delete bwt
         <- bwt
    bwt <- NULL
})
## Reorder variables
lbw.for.bestglm <-</pre>
    lbw.for.bestglm[,
c("age","lwt","race.cat","smoke","preterm","ht","ui","ftv.cat","y")]
```

Perform all-subset linear (gaussian) regression based on Akaike Information Criteria (AIC)

```
Morgan-Tatar search since factors present with more than 2 levels.
```

Show top 5 models
res.bestglm\$BestModels

```
ui ftv.cat Criterion
    age lwt race.cat smoke preterm
                                       ht
                                                            2450.2
1 FALSE TRUE
                 TRUE
                                TRUE TRUE TRUE
                                                  FALSE
                        TRUE
2 FALSE TRUE
                                                            2450.7
                 TRUE
                        TRUE
                               FALSE TRUE TRUE
                                                  FALSE
                                                            2452.1
  TRUE TRUE
                                TRUE TRUE TRUE
                 TRUE
                        TRUE
                                                  FALSE
4
                               FALSE TRUE TRUE
                                                  FALSE
                                                            2452.5
  TRUE TRUE
                 TRUE
                        TRUE
5 FALSE TRUE
                                                            2453.3
                 TRUE
                        TRUE
                                TRUE TRUE TRUE
                                                   TRUE
```

Show result for the best model

The model identical to the one chosen by the adjusted \(R^2 \) was selected.

```
summary(res.bestglm$BestModel)
```

```
call:
lm(formula = y \sim ., data = data.frame(xy[, c(bestset[-1],
FALSE),
    drop = FALSE, y = y)
Residuals:
                 Median
    Min
             10
                              30
                                     Max
-1886.4
         -441.0
                    53.5
                           494.2
                                  1620.9
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                                              <2e-16 ***
(Intercept)
               2871.99
                            243.67
                                      11.79
Ìwt
                  4.04
                              1.67
                                       2.42
                                              0.0167 *
race.catBlack
                                      -3.21
               -466.32
                            145.13
                                              0.0016 **
                                     -2.99
                                              0.0032 **
race.catOther
               -335.68
                            112.28
smoke
                -323.57
                            104.96
                                      -3.08
                                              0.0024 **
                            133.50
preterm1+
                -208.44
                                      -1.56
                                              0.1202
                -573.69
                            198.96
                                     -2.88
                                              0.0044 **
ht
                                              0.0004 ***
               -489.96
                            135.93
                                     -3.60
นา์
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 643 on 181 degrees of freedom
Multiple R-squared: 0.25,
                           Adjusted R-squared: 0.221
F-statistic: 8.64 on 7 and 181 DF, p-value: 0.00000000396
```

Logistic regression

Do the same, but as a logistic regression model. The resulting model was identical to the best linear model in this case.

```
## Prepare data
lbw.for.best.logistic <- within(lbw, {</pre>
        <- NULĹ
                        # Delete
    id
    bwt
        <- NULL
    race <- NULL
    ptl <- NULL
    ftv <- NULL
         <- low
                       # bwt into y
    low <- NULL
                     # Delete bwt
})
## Reorder variables
lbw.for.best.logistic <-</pre>
    lbw.for.best.logistic[,
c("age","lwt","race.cat","smoke","preterm","ht","ui","ftv.cat","y")]
## Perform
res.best.logistic <-
    bestglm(Xy = lbw.for.best.logistic,
                                        # binomial family for
            family = binomial,
logistic
                                        # Information criteria
            IC = "AIC",
for
            method = "exhaustive")
```

Morgan-Tatar search since family is non-gaussian. Note: factors present with more than 2 levels.

```
## Show top 5 models
res.best.logistic$BestModels
```

```
age lwt race.cat smoke preterm
                                     ht
                                           ui ftv.cat
Criterion
1 FALSE TRUE
                TRUE TRUE
                              TRUE TRUE TRUE
                                                FALSE
211.85
2 FALSE TRUE
                TRUE
                      TRUE
                              TRUE TRUE FALSE
                                                FALSE
212.48
3 TRUE TRUE
                TRUE
                      TRUE
                              TRUE TRUE TRUE
                                                FALSE
212.83
4 TRUE TRUE
                              TRUE TRUE FALSE
                TRUE TRUE
                                                FALSE
213.15
5 FALSE TRUE
                              TRUE TRUE TRUE
                TRUE TRUE
                                                 TRUE
214.37
```

```
## Show result for the best model: Same model was chosen
summary(res.best.logistic$BestModel)
```

```
call:
glm(formula = y \sim ., family = family, data = Xi, weights =
weights)
Deviance Residuals:
            1Q Median
                             3Q
                                    Max
-1.731
       -0.784
               -0.514
                          0.954
                                  2.198
Coefficients:
              Estimate Std. Error z value Pr(>|z|)
                                              0.8969
(Intercept)
              -0.12533
                           0.96756
                                     -0.13
              -0.01592
                           0.00695
                                     -2.29
lwt
                                              0.0221 *
race.catBlack
               1.30086
                           0.52848
                                      2.46
                                              0.0138 *
               0.85441
                           0.44091
                                      1.94
                                              0.0526
race.catOther
                           0.40447
smoke
               0.86658
                                      2.14
                                       2.51
preterm1+
               1.12886
                           0.45039
                                              0.0122
                           0.70737
                                              0.0083 **
ht
               1.86690
                                      2.64
               0.75065
                           0.45882
                                      1.64
                                              0.1018
ui
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 234.67
                            on 188
                                    degrees of freedom
Residual deviance: 197.85
                            on 181
                                    degrees of freedom
AIC: 213.9
Number of Fisher Scoring iterations: 4
```

glmulti (Model selection and multimodel inference made easy)

Automated model selection and model-averaging. Provides a wrapper for glm and other functions, automatically generating all possible models (under constraints set by the user) with the specified response and explanatory variables, and finding the best models in terms of some Information Criterion (AIC, AICc or BIC). Can handle very large numbers of candidate models. Features a Genetic Algorithm to find the best models when an exhaustive screening of the candidates is not feasible.

References

- http://www.jstatsoft.org/v34/i12/paper
- http://cran.r-project.org/web/packages/glmulti/index.html

Load package

```
library(glmulti)
```

All-subset linear regression using Im() based on AIC

```
glmulti.lm.out <-
    glmulti(bwt ~ age + lwt + race.cat + smoke + preterm + ht +
ui + ftv.cat, data = lbw,
    level = 1,  # No interaction

considered

method = "h",  # Exhaustive approach
    crit = "aic",  # AIC as criteria
    confsetsize = 5,  # Keep 5 best models
    plotty = F, report = F, # No plot or interim

reports

fitfunction = "lm")  # lm function

## Show 5 best models (Use @ instead of $ for an $4 object)
glmulti.lm.out@formulas</pre>
```

```
[[1]]
bwt ~ 1 + race.cat + preterm + lwt + smoke + ht + ui
<environment: 0x11a320bb8>

[[2]]
bwt ~ 1 + race.cat + lwt + smoke + ht + ui
<environment: 0x11a320bb8>

[[3]]
bwt ~ 1 + race.cat + preterm + age + lwt + smoke + ht + ui
<environment: 0x11a320bb8>

[[4]]
bwt ~ 1 + race.cat + age + lwt + smoke + ht + ui
<environment: 0x11a320bb8>

[[5]]
bwt ~ 1 + race.cat + preterm + ftv.cat + lwt + smoke + ht + ui
<environment: 0x11a320bb8>
```

```
## Show result for the best model
summary(glmulti.lm.out@objects[[1]])
```

```
call:
fitfunc(formula = as.formula(x), data = data)
Residuals:
    Min
             1Q
                 Median
                              3Q
                                     Max
-1886.4
                   53.5
                           494.2
                                  1620.9
         -441.0
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
                                             <2e-16 ***
               2871.99
                            243.67
(Intercept)
                                     11.79
                                             0.0016 **
race.catBlack
               -466.32
                            145.13
                                     -3.21
                            112.28
               -335.68
                                     -2.99
                                             0.0032 **
race.catOther
                                     -1.56
               -208.44
                            133.50
                                             0.1202
preterm1+
lwt
                  4.04
                              1.67
                                      2.42
                                             0.0167 *
                                             0.0024 **
                            104.96
smoke
               -323.57
                                     -3.08
                                             0.0044 **
ht
               -573.69
                            198.96
                                     -2.88
               -489.96
                                     -3.60
                                             0.0004 ***
ui
                            135.93
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 643 on 181 degrees of freedom
Multiple R-squared: 0.25,
                           Adjusted R-squared: 0.221
F-statistic: 8.64 on 7 and 181 DF, p-value: 0.00000000396
```

All-subset logistic regression using glm() based on AIC

```
qlmulti.logistic.out <-</pre>
    glmulti(low ~ age + lwt + race.cat + smoke + preterm + ht +
ui + ftv.cat, data = 1bw,
                                        # No interaction
             level = 1.
considered
             method = "h"
                                        # Exhaustive approach
             crit = "aic".
                                        # AIC as criteria
             confsetsize = 5,
                                        # Keep 5 best models
             plotty = F, report = F, # No plot or interim
reports
             fitfunction = "glm",  # glm function
family = binomial)  # binomial family for
logistic regression
## Show 5 best models (Use @ instead of $ for an S4 object)
glmulti.logistic.out@formulas
```

```
[[1]]
low ~ 1 + race.cat + preterm + lwt + smoke + ht + ui
<environment: 0x11a2588e8>
[[2]]
low ~ 1 + race.cat + preterm + lwt + smoke + ht
<environment: 0x11a2588e8>
[[3]]
low ~ 1 + race.cat + preterm + age + lwt + smoke + ht + ui
<environment: 0x11a2588e8>
[[4]]
low ~ 1 + race.cat + preterm + age + lwt + smoke + ht
<environment: 0x11a2588e8>
[[5]] low \sim 1 + race.cat + preterm + ftv.cat + lwt + smoke + ht + ui
<environment: 0x11a2588e8>
## Show result for the best model
summary(glmulti.logistic.out@objects[[1]])
call:
fitfunc(formula = as.formula(x), family = ...1, data = data)
Deviance Residuals:
            1Q Median
                             30
                                    Max
-1.731 - 0.784
                         0.954
                                  2.198
               -0.514
Coefficients:
              Estimate Std. Error z value Pr(>|z|)
(Intercept)
              -0.12533
                          0.96756
                                     -0.13
                                             0.8969
race.catBlack 1.30086
                           0.52848
                                      2.46
                                             0.0138 *
                                      1.94
               0.85441
                          0.44091
                                             0.0526
race.catOther
                                      2.51
                                             0.0122 *
preterm1+
               1.12886
                          0.45039
                                     -2.29
                                             0.0221 *
                          0.00695
lwt
              -0.01592
                                             0.0322 *
smoke
               0.86658
                          0.40447
                                     2.14
               1.86690
                          0.70737
                                      2.64
                                             0.0083 **
ht
ui
               0.75065
                          0.45882
                                      1.64
                                             0.1018
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 234.67

Number of Fisher Scoring iterations: 4

Residual deviance: 197.85

AIC: 213.9

on 188

degrees of freedom

on 181 degrees of freedom

Load pbc survival data in survival package

```
library(survival)
pbc <- within(pbc, {
    status.dichotomous <- status > 1
    survival.vector <- Surv(time, status.dichotomous)
})</pre>
```

All-subset Cox regression using coxph() based on AIC

```
glmulti.coxph.out <-
    glmulti(survival.vector ~ trt + age + sex + ascites +
hepato + spiders, data = pbc,
    level = 1,  # No interaction

considered

method = "h",  # Exhaustive approach
    crit = "aic",  # AIC as criteria
    confsetsize = 5,  # Keep 5 best models
    plotty = F, report = F, # No plot or interim

reports

fitfunction = "coxph") # coxph function

## Show 5 best models (Use @ instead of $ for an $4 object)
glmulti.coxph.out@formulas</pre>
```

```
[[1]]
survival.vector ~ 1 + age + ascites + hepato + spiders
<environment: 0x104baef88>
[[2]]
survival.vector ~ 1 + trt + age + ascites + hepato + spiders
<environment: 0x104baef88>
[[3]]
survival.vector \sim 1 + \text{sex} + \text{age} + \text{ascites} + \text{hepato} + \text{spiders}
<environment: 0x104baef88>
[[4]]
survival.vector ~ 1 + sex + trt + age + ascites + hepato +
spiders
<environment: 0x104baef88>
[[51]
survival.vector ~ 1 + sex + ascites + hepato + spiders
<environment: 0x104baef88>
```

```
## Show result for the best model
summary(glmulti.coxph.out@objects[[1]])
```

```
call:
fitfunc(formula = as.formula(x), data = data)
  n= 312, number of events= 125
   (106 observations deleted due to missingness)
           coef exp(coef) se(coef)
                                             Pr(>|z|)
                                      Ζ
                           0.00864 3.03
                                              0.00242 **
        0.02620
                  1.02655
age
                           0.26109 5.73 0.0000000099 ***
ascites 1.49687
                  4.46770
hepato 0.84318
                  2.32375
                           0.21198 3.98 0.0000695829 ***
                                              0.00065 ***
                  1.94646 0.19530 3.41
spiders 0.66601
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
        exp(coef) exp(-coef) lower .95 upper .95
age
             1.03
                       0.974
                                  1.01
ascites
             4.47
                       0.224
                                   2.68
                                             7.45
                                             3.52
             2.32
                       0.430
hepato
                                  1.53
             1.95
                       0.514
                                  1.33
                                             2.85
spiders
Concordance= 0.771 (se = 0.029)
Rsquare= 0.27 (max possible= 0.983)
Likelihood ratio test= 98.3
                                         p=0
Wald test
                     = 121
                            on 4 df,
                                        p=0
Score (logrank) test = 156
                           on 4 df.
                                        0=q
```

meifly (Interactive model exploration using GGobi)

Exploratory model analysis. Fit and graphical explore ensembles of linear models.

This function just conduct all-subset regression, thus it can handle coxph without problems, but users will have to do model comparison using the result object. Interaction terms cannot be handled, thus inclusion of interaction terms needs creation of product term beforehand.

References:

http://cran.r-project.org/web/packages/meifly/index.html

Load meifly package

```
library(meifly)
```

Fit all subsets (main effects only)

For x, give a data.frame without the outcome variable.

Show the result

As expected, it is the same as the model chosen here (http://rpubs.com/kaz_yos/exhaustive).

```
## Extract AIC from each model
fitall.out.aic <- t(sapply(fitall.out, extractAIC))

## Create an order list of increasing AIC
final.out.order <- order(fitall.out.aic[,2])

## Show the result for the best model
fitall.out[final.out.order][1]</pre>
```

```
$`58`
call:
coxph(formula = y \sim age + ascites + hepato + spiders, data =
data.
    model = FALSE
          coef exp(coef) se(coef)
                          0.00864 3.03 0.0024000000
                     1.03
        0.0262
age
                     4.47
                           0.26109 5.73 0.0000000099
ascites 1.4969
                          0.21198 3.98 0.0000700000
hepato 0.8432
                     2.32
spiders 0.6660
                     1.95 0.19530 3.41 0.0006500000
Likelihood ratio test=98.3 on 4 df, p=0 n= 312, number of
events= 125
   (106 observations deleted due to missingness)
attr(,"class")
[1] "ensemble"
```

Exhaustive model selection without using packages

See: http://rpubs.com/kaz_yos/exhaustive