# Linear regression

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### 1 Introduction

- 1. A statistical method to model relationship between:
  - outcome: numerical variable.
  - predictors/independent variables: numerical, categorical variables.

- 2. A type of Generalized Linear Models (GLMs), which also includes other outcome types, e.g. categorical and count.
- 3. Basically, the linear relationship is structured as follows,

 $numerical\ outcome = numerical\ predictors + categorical\ predictors$ 

### 2 Simple linear regression (SLR)

### 2.1 About SLR

- 1. Model *linear* (straight line) relationship between:
  - outcome: numerical variable.
  - a predictor: numerical variable (only).

*Note*: What if the predictor is a categorical variable? Remember, we already handled that with one-way ANOVA.

2. Formula,

 $numerical\ outcome = intercept + coefficient \times numerical\ predictor$ 

in short,

$$\hat{y} = \beta_0 + \beta_1 x_1$$

where  $\hat{y}$  is the predicted value of the outcome y.

### 2.2 Analysis

#### 2.2.1 Libraries

```
# library
library(foreign)
library(epiDisplay)

## Loading required package: survival

## Loading required package: MASS

## Loading required package: nnet
library(psych)

##

## Attaching package: 'psych'

## The following objects are masked from 'package:epiDisplay':

##

## alpha, cs, lookup
library(lattice)

##

## Attaching package: 'lattice'
```

```
## The following object is masked from 'package:epiDisplay':
##
## dotplot
library(rsq)
library(MASS)
library(car)

## Loading required package: carData
##
## Attaching package: 'car'
## The following object is masked from 'package:psych':
##
## logit
library(broom)
```

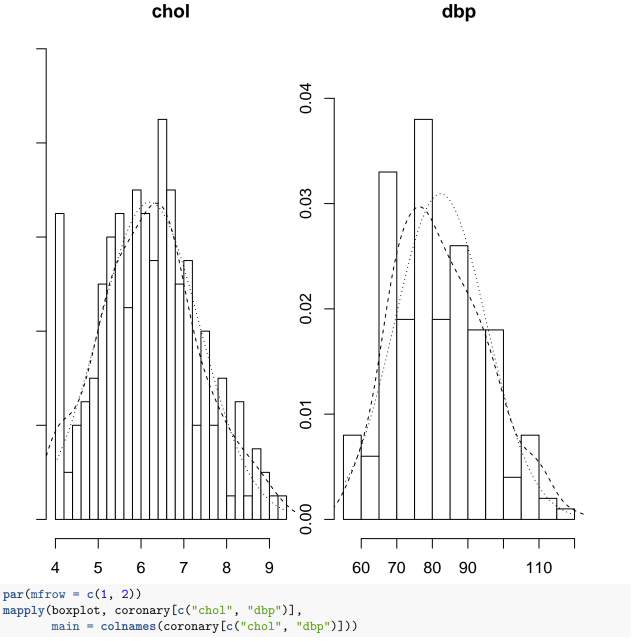
#### 2.2.2 Data set

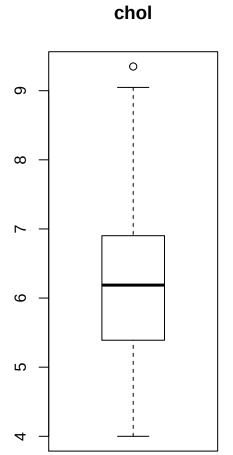
```
# data
coronary = read.dta("coronary.dta")
str(coronary)
                   200 obs. of 9 variables:
## 'data.frame':
## $ id : num 1 14 56 61 62 64 69 108 112 134 ...
## $ cad : Factor w/ 2 levels "no cad", "cad": 1 1 1 1 1 1 2 1 1 1 ...
## $ sbp : num 106 130 136 138 115 124 110 112 138 104 ...
## $ dbp
          : num 68 78 84 100 85 72 80 70 85 70 ...
## $ chol : num 6.57 6.33 5.97 7.04 6.66 ...
## $ age
          : num 60 34 36 45 53 43 44 50 43 48 ...
           : num 38.9 37.8 40.5 37.6 40.3 ...
## $ bmi
## $ race : Factor w/ 3 levels "malay", "chinese",..: 3 1 1 1 3 1 1 2 2 2 ...
## $ gender: Factor w/ 2 levels "woman", "man": 1 1 1 1 2 2 2 1 1 2 ...
## - attr(*, "datalabel")= chr "Written by R.
## - attr(*, "time.stamp")= chr ""
## - attr(*, "formats")= chr "%9.0g" "%9.0g" "%9.0g" "%9.0g" ...
## - attr(*, "types")= int 100 108 100 100 100 100 108 108
## - attr(*, "val.labels")= chr "" "cad" "" "" ...
## - attr(*, "var.labels")= chr "id" "cad" "sbp" "dbp" ...
## - attr(*, "version")= int 7
## - attr(*, "label.table")=List of 3
    ..$ cad : Named int 1 2
##
##
   .. ..- attr(*, "names")= chr "no cad" "cad"
##
    ..$ race : Named int 1 2 3
    ....- attr(*, "names")= chr "malay" "chinese" "indian"
##
    ..$ gender: Named int 12
##
##
    ....- attr(*, "names")= chr "woman" "man"
```

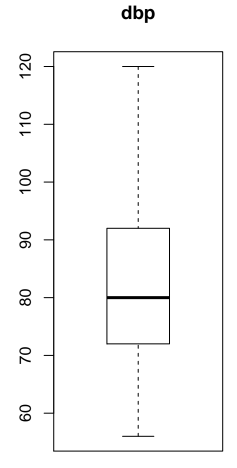
### 2.2.3 Data exploration

#### 2.2.3.1 Descriptive statistics

```
summ(coronary[c("chol", "dbp")])
##
## No. of observations = 200
##
##
     Var. name obs. mean
                           median s.d.
                                          min.
                                                  max.
## 1 chol
               200 6.2
                           6.19
                                   1.18
                                           4
                                                  9.35
## 2 dbp
               200 82.31 80
                                   12.9
                                                  120
                                          56
2.2.3.2 Plots
multi.hist(coronary[c("chol", "dbp")], ncol = 2)
```







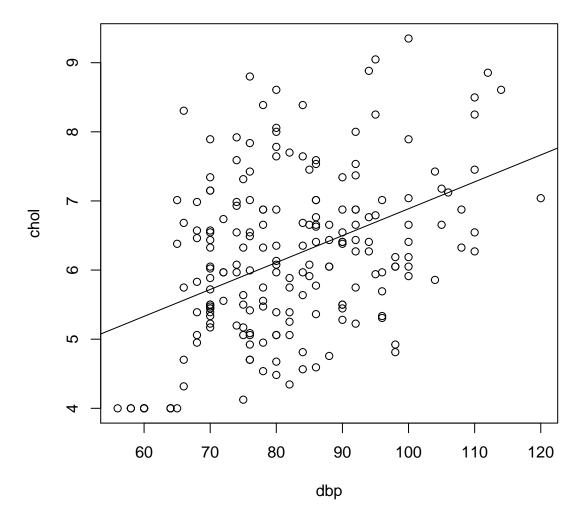
```
## chol dbp
## stats Numeric,5 Numeric,5
## n 200 200
## conf Numeric,2 Numeric,2
## out 9.35 Numeric,0
## group 1 Numeric,0
## names "" ""
par(mfrow = c(1, 1))
```

### 2.2.4 Univariable

```
Fit model,
```

```
# model: chol ~ dbp
slr_chol = glm(chol ~ dbp, data = coronary)
summary(slr_chol)
```

```
##
## Call:
## glm(formula = chol ~ dbp, data = coronary)
## Deviance Residuals:
##
      Min
                1Q
                     Median
                                   3Q
                                           Max
## -1.9967 -0.8304 -0.1292 0.7734
                                        2.8470
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.995134
                          0.492092
                                   6.087 5.88e-09 ***
              0.038919
                         0.005907
                                     6.589 3.92e-10 ***
## dbp
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.154763)
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 228.64 on 198 degrees of freedom
## AIC: 600.34
##
## Number of Fisher Scoring iterations: 2
Confint(slr_chol) # 95% CI
                               2.5 %
##
                                         97.5 %
                Estimate
## (Intercept) 2.99513427 2.03065127 3.95961727
              0.03891876 0.02734161 0.05049591
## dbp
Important results,
  • Coefficient, \beta.
  • 95% CI.
  • P-value.
Obtain R^2, % of variance explained,
rsq(slr_chol, adj = T)
## [1] 0.1756834
Scatter plot,
plot(chol ~ dbp, data = coronary)
abline(slr_chol)
```



this allows assessment of normality, linearity and equal variance assumptions. We expect eliptical/oval shape (normality), equal scatter of dots on both sides of the prediction line (equal variance). Both these indicate linear relationship between chol and dbp.

### 2.2.5 Interpretation

- $\bullet\,$  1mmHg increase in DBP causes 0.04mmol/L increase in cholestrol.
- $\bullet~$  DBP explains 17.6% variance in cholestrol.

### 2.2.6 Model equation

$$chol = 3.0 + 0.04 \times dbp$$

### 3 Multiple linear regression (MLR)

#### 3.1 About MLR

- 1. Model *linear* relationship between:
  - outcome: numerical variable.
  - predictors: numerical, categorical variables.

Note: MLR is a term that refers to linear regression with two or more numerical variables. Whenever we have both numerical and categorical variables, the proper term for the regression model is General Linear Model. However, we will use the term MLR in this workshop.

2. Formula,

 $numerical\ outcome = intercept + coefficients \times numerical\ predictors \\ + coefficients \times categorical\ predictors$ 

in a shorter form,

$$\hat{y} = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_k x_k$$

where we have k predictors.

Whenever the predictor is a categorical variable with more than two levels, we use dummy variable(s). This can be easily specified in R using factor() if the variable is not yet properly specified as such. There is no problem with binary categorical variable.

For a categorical variable with more than two levels, the number of dummy variables (i.e. once turned into several binary variables) equals number of levels minus one. For example, whenever we have four levels, we will obtain three dummy (binary) variables.

### 3.2 Analysis

### 3.2.1 Review data set

```
# data
str(coronary)
```

```
'data.frame':
                    200 obs. of 9 variables:
##
   $ id
                  1 14 56 61 62 64 69 108 112 134 ...
            : num
            : Factor w/ 2 levels "no cad", "cad": 1 1 1 1 1 1 2 1 1 1 ...
##
   $ sbp
                   106 130 136 138 115 124 110 112 138 104 ...
                   68 78 84 100 85 72 80 70 85 70 ...
##
     dbp
            : num
                   6.57 6.33 5.97 7.04 6.66 ...
##
   $
     chol
           : num
##
   $ age
                   60 34 36 45 53 43 44 50 43 48 ...
            : num
            : num 38.9 37.8 40.5 37.6 40.3 ...
##
   $ bmi
           : Factor w/ 3 levels "malay", "chinese", ...: 3 1 1 1 3 1 1 2 2 2 ...
##
##
   $ gender: Factor w/ 2 levels "woman", "man": 1 1 1 1 2 2 2 1 1 2 ...
   - attr(*, "datalabel")= chr "Written by R.
   - attr(*, "time.stamp")= chr ""
##
##
   - attr(*, "formats")= chr "%9.0g" "%9.0g" "%9.0g" "%9.0g" ...
   - attr(*, "types")= int 100 108 100 100 100 100 100 108 108
   - attr(*, "val.labels")= chr "" "cad" "" "" ...
   - attr(*, "var.labels")= chr "id" "cad" "sbp" "dbp" ...
   - attr(*, "version")= int 7
   - attr(*, "label.table")=List of 3
             : Named int 12
     ..$ cad
##
```

```
## ...- attr(*, "names")= chr "no cad" "cad"
## ..$ race : Named int 1 2 3
## ...- attr(*, "names")= chr "malay" "chinese" "indian"
## ..$ gender: Named int 1 2
## ...- attr(*, "names")= chr "woman" "man"
```

We exclude id, cad and age from our data for the purpose of this analysis, keeping only sbp, dbp, bmi, race and gender. We will add age later in the exercise.

```
coronary = subset(coronary, select = -c(id, cad, age))
# remove id, cad, age from our data since we're not going to use them,
# easier to specify multivariable model.
```

### 3.2.2 Data exploration

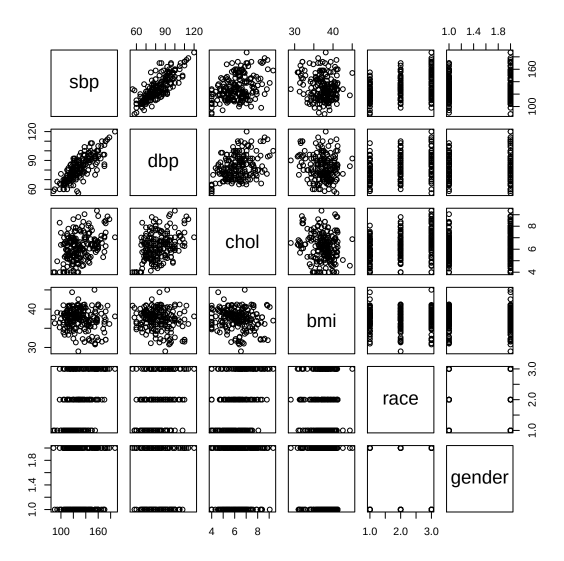
### 3.2.2.1 Descriptive statistics

```
summ(coronary[c("chol", "sbp", "dbp", "bmi")])
##
## No. of observations = 200
##
    Var. name obs. mean
                          median s.d.
                                          min.
                                                 max.
## 1 chol
              200 6.2
                           6.19
                                   1.18
                                          4
                                                 9.35
## 2 sbp
              200 130.18 126
                                   19.81 88
                                                 187
              200 82.31 80
                                   12.9
                                                 120
## 3 dbp
                                          56
## 4 bmi
              200 37.45 37.8
                                   2.68
                                          28.99 45.03
codebook(coronary[c("race", "gender")])
```

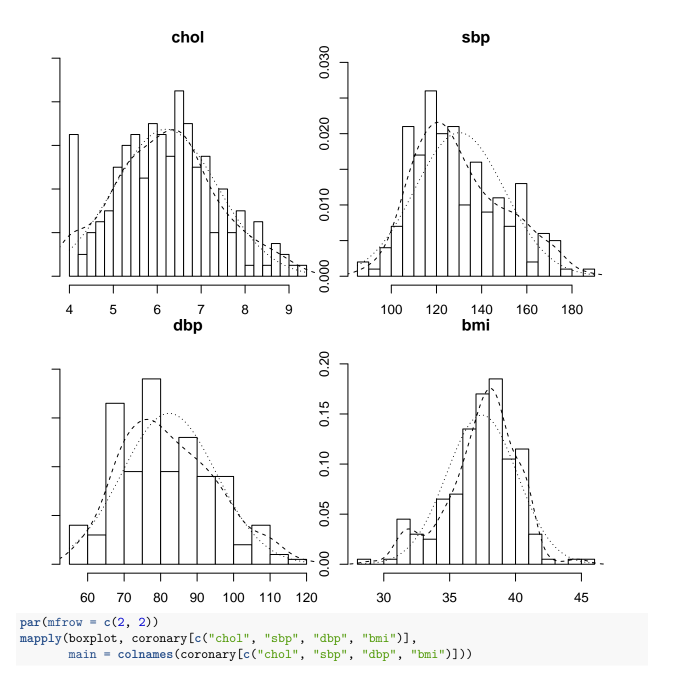
```
##
##
##
## race
          Frequency Percent
##
## malay
                 73
                      36.5
## chinese
                 64
                       32.0
## indian
                 63
                      31.5
## ========
## gender
            :
        Frequency Percent
## woman
                      50
              100
              100
                       50
## man
##
      ______
```

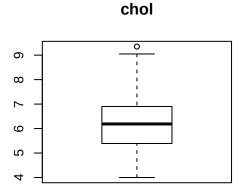
### 3.2.2.2 Plots

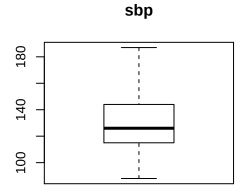
```
plot(coronary)
```

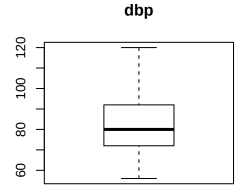


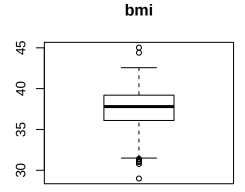
multi.hist(coronary[c("chol", "sbp", "dbp", "bmi")])



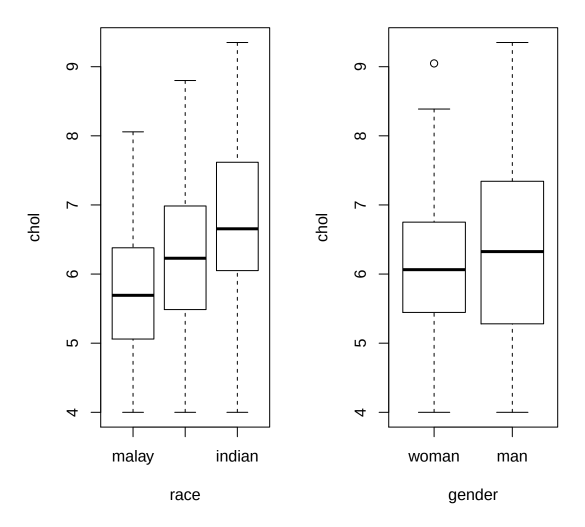








```
##
        chol
                            dbp
                                     bmi
                  sbp
## stats Numeric,5 Numeric,5 Numeric,5
        200
                  200
                            200
## n
## conf Numeric,2 Numeric,2 Numeric,2
## out
        9.35
                  Numeric, 0 Numeric, 0 Numeric, 8
## group 1
                  Numeric, 0 Numeric, 0 Numeric, 8
## names ""
par(mfrow = c(1, 1))
par(mfrow = c(1, 2))
boxplot(chol ~ race, data = coronary)
boxplot(chol ~ gender, data = coronary)
```



```
par(mfrow = c(1, 1))
```

### 3.2.3 Variable selection

### 3.2.3.1 Univariable

Perform SLR for chol, sbp, dbp and bmi on your own as shown above. Now, we are concerned with which variables are worthwhile to include in the multivariable models.

We want to choose only variables with P-values < 0.25 to be included in MLR. Obtaining the P-values for each variable is easy by LR test,

```
slr_chol0 = glm(chol ~ 1, data = coronary)
summary(slr_chol0)

##
## Call:
## glm(formula = chol ~ 1, data = coronary)
```

```
##
## Deviance Residuals:
##
       \mathtt{Min}
                  1Q
                        Median
## -2.19854 -0.80854 -0.01104 0.69021
                                           3.15146
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 6.19854
                                    74.06 <2e-16 ***
                          0.08369
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.400874)
##
      Null deviance: 278.77 on 199 degrees of freedom
##
## Residual deviance: 278.77 on 199 degrees of freedom
## AIC: 637.99
## Number of Fisher Scoring iterations: 2
names(coronary)
## [1] "sbp"
               "dbp"
                        "chol"
                                 "bmi"
                                          "race"
                                                   "gender"
add1(slr_chol0, scope = ~ sbp + dbp + bmi + race + gender, test = "LRT")
## Single term additions
##
## Model:
## chol ~ 1
##
         Df Deviance
                        AIC scaled dev. Pr(>Chi)
              278.77 637.99
## <none>
                                 33.855 5.938e-09 ***
              235.36 606.14
## sbp
          1
                               39.648 3.042e-10 ***
## dbp
          1
              228.64 600.34
                                 4.792 0.02859 *
## bmi
          1
             272.17 635.20
## race
          2
              241.68 613.43
                                28.561 6.280e-07 ***
## gender 1
              277.45 639.04
                                 0.952
                                          0.32933
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

All variables are significant and < .25 except gender. So proceed with the rest of the variables, excluding gender.

#### 3.2.3.2 Multivariable

Perform MLR with all selected variables,

```
# all
mlr_chol = glm(chol ~ sbp + dbp + bmi + race, data = coronary)
#mlr_chol = glm(chol ~ ., data = coronary) # shortcut
summary(mlr_chol)

##
## Call:
## glm(formula = chol ~ sbp + dbp + bmi + race, data = coronary)
##
## Deviance Residuals:
## Min 1Q Median 3Q Max
```

```
## -2.17751 -0.73860 -0.02674
                                 0.63163
                                            2.90926
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
               4.842338
                          1.265149
                                     3.827 0.000175
               0.000975
                          0.006990
                                     0.139 0.889210
## sbp
## dbp
               0.028350
                          0.010327
                                     2.745 0.006615 **
## bmi
               -0.038537
                          0.028170
                                    -1.368 0.172879
## racechinese 0.354039
                          0.183169
                                     1.933 0.054710 .
## raceindian
              0.716327
                          0.200346
                                     3.575 0.000441 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.089387)
##
##
       Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 211.34 on 194 degrees of freedom
## AIC: 592.61
## Number of Fisher Scoring iterations: 2
rsq(mlr_chol, adj = T)
```

### ## [1] 0.2223518

Focus on,

- Coefficients,  $\beta$ s.
- 95% CI.
- P-values.

For model fit,

- $R^2 \%$  of variance explained by the model.
- Akaike Information Criterion, AIC for comparison with other models. This is not useful alone, but for comparison with other models. The model with the lowest AIC is the best model.

### 3.2.3.3 Stepwise

As you can see, not all variables are significant. How to select? We proceed with stepwise automatic selection,

```
# stepwise
# both
mlr_chol_stepboth = step(mlr_chol, direction = "both")
## Start: AIC=592.61
## chol ~ sbp + dbp + bmi + race
##
##
          Df Deviance
## - sbp
               211.36 590.63
           1
## - bmi
           1
               213.38 592.53
## <none>
               211.34 592.61
## - dbp
           1
               219.55 598.23
## - race 2
               225.30 601.40
## Step: AIC=590.63
## chol ~ dbp + bmi + race
```

```
##
##
        Df Deviance
                       ATC
## - bmi 1 213.40 590.55
              211.36 590.63
## <none>
         1
## + sbp
              211.34 592.61
## - race 2
              227.04 600.94
## - dbp
          1
              235.88 610.58
##
## Step: AIC=590.55
## chol ~ dbp + race
##
         Df Deviance
                       AIC
              213.40 590.55
## <none>
             211.36 590.63
## + bmi 1
## + sbp 1 213.38 592.53
## - race 2
              228.64 600.34
## - dbp 1 241.68 613.43
summary(mlr_chol_stepboth) # racechinese marginally sig.
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
## Deviance Residuals:
##
     Min 1Q Median
                                 3Q
## -2.1378 -0.7068 -0.0289 0.5997
                                      2.7778
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 3.298028  0.486213  6.783 1.36e-10 ***
              0.031108 0.006104
                                  5.096 8.14e-07 ***
## racechinese 0.359964 0.182149
                                  1.976 0.049534 *
## raceindian 0.713690 0.190883
                                  3.739 0.000243 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.088777)
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.40 on 196 degrees of freedom
## AIC: 590.55
##
## Number of Fisher Scoring iterations: 2
mlr_chol_stepforward = step(slr_chol0, scope = ~ sbp + dbp + bmi + race + gender,
                       direction = "forward")
## Start: AIC=637.99
## chol ~ 1
##
##
           Df Deviance
                         AIC
## + dbp
            1
                228.64 600.34
## + sbp
            1
                235.36 606.14
          2 241.68 613.43
## + race
```

```
## + bmi
               272.17 635.20
                278.77 637.99
## <none>
               277.45 639.04
## + gender 1
##
## Step: AIC=600.34
## chol ~ dbp
##
##
           Df Deviance
                         AIC
## + race
           2 213.40 590.55
               228.64 600.34
## <none>
## + gender 1
               226.64 600.58
               226.96 600.87
## + sbp
            1
## + bmi
               227.04 600.94
            1
##
## Step: AIC=590.55
## chol ~ dbp + race
##
##
           Df Deviance
                         AIC
## <none>
               213.40 590.55
               211.36 590.63
## + bmi
            1
## + gender 1
               212.47 591.67
## + sbp
            1
               213.38 592.53
summary(mlr_chol_stepforward) # same with both
##
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
##
## Deviance Residuals:
##
      Min 1Q Median
                                 ЗQ
                                        Max
## -2.1378 -0.7068 -0.0289 0.5997
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.298028  0.486213  6.783 1.36e-10 ***
              ## dbp
## racechinese 0.359964  0.182149  1.976  0.049534 *
## raceindian 0.713690 0.190883
                                 3.739 0.000243 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.088777)
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.40 on 196 degrees of freedom
## AIC: 590.55
## Number of Fisher Scoring iterations: 2
# backward
mlr_chol_stepback = step(mlr_chol, direction = "backward")
## Start: AIC=592.61
## chol ~ sbp + dbp + bmi + race
```

```
##
##
         Df Deviance
                        ATC
              211.36 590.63
## - sbp
         1
              213.38 592.53
## - bmi
          1
## <none>
              211.34 592.61
## - dbp
              219.55 598.23
          1
## - race 2
              225.30 601.40
##
## Step: AIC=590.63
## chol ~ dbp + bmi + race
##
         Df Deviance
                        AIC
          1 213.40 590.55
## - bmi
              211.36 590.63
## <none>
## - race 2
              227.04 600.94
## - dbp
          1
              235.88 610.58
##
## Step: AIC=590.55
## chol ~ dbp + race
         Df Deviance
##
                        AIC
## <none>
              213.40 590.55
## - race 2
              228.64 600.34
## - dbp
          1
              241.68 613.43
summary(mlr_chol_stepback) # same with both
##
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
##
## Deviance Residuals:
##
      \mathtt{Min}
              1Q
                    Median
                                  3Q
                                          Max
## -2.1378 -0.7068 -0.0289
                              0.5997
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 3.298028 0.486213
                                   6.783 1.36e-10 ***
              0.031108
                        0.006104
                                    5.096 8.14e-07 ***
## racechinese 0.359964 0.182149
                                    1.976 0.049534 *
## raceindian 0.713690
                                    3.739 0.000243 ***
                        0.190883
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 1.088777)
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.40 on 196 degrees of freedom
## AIC: 590.55
##
## Number of Fisher Scoring iterations: 2
Looking at all these results, we choose:
    chol ~ dbp + race
```

which has the lowest AIC.

```
mlr_chol1 = glm(chol ~ dbp + race, data = coronary)
summary(mlr_chol1)
##
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
##
## Deviance Residuals:
##
      Min
                 10
                     Median
                                   3Q
                                           Max
                               0.5997
## -2.1378 -0.7068 -0.0289
                                        2.7778
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.298028
                          0.486213
                                     6.783 1.36e-10 ***
              0.031108
                          0.006104
                                     5.096 8.14e-07 ***
## racechinese 0.359964
                          0.182149
                                     1.976 0.049534 *
                          0.190883
                                     3.739 0.000243 ***
## raceindian 0.713690
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 1.088777)
##
       Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.40 on 196 degrees of freedom
## AIC: 590.55
##
## Number of Fisher Scoring iterations: 2
```

#### 3.2.3.4 Confounder

If we include a variable and it causes notable change (>20%) in the coefficients of other variables, it is a confounder. When the confounder is significant and the main effect variable is also significant, we keep the confounder in the model.

Formula for % change,

```
100 * (model_small - model_large) / model_large
Hosmer, Lemeshow, & Sturdivant (2013)
```

Start by including common demographic adjustment, gender,

```
# + gender
mlr_chol2 = glm(chol ~ dbp + race + gender, data = coronary)
summary(mlr_chol2) # higher AIC, gender insig.
##
## Call:
## glm(formula = chol ~ dbp + race + gender, data = coronary)
##
## Deviance Residuals:
##
        Min
                                        3Q
                   10
                         Median
                                                 Max
## -2.06350 -0.71634 -0.04471
                                             2.70974
                                   0.64533
##
## Coefficients:
```

```
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.203032 0.497111
                                     6.443 8.94e-10 ***
                                     5.149 6.37e-07 ***
               0.031533
                          0.006124
                         0.182369
## racechinese 0.353052
                                     1.936
                                             0.0543 .
## raceindian 0.692724
                          0.192293
                                     3.602
                                             0.0004 ***
                                     0.925
                                             0.3560
## genderman
              0.137663
                          0.148790
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for gaussian family taken to be 1.089578)
##
       Null deviance: 278.77 on 199 degrees of freedom
##
## Residual deviance: 212.47 on 195 degrees of freedom
## AIC: 591.67
##
## Number of Fisher Scoring iterations: 2
coef(mlr_chol2); coef(mlr_chol1)
## (Intercept)
                                                     genderman
                       dbp racechinese raceindian
##
     3.2030318
                 0.0315331
                             0.3530516
                                         0.6927239
                                                     0.1376627
## (Intercept)
                       dbp racechinese raceindian
## 3.29802826 0.03110811 0.35996365 0.71369024
100 * (coef(mlr_chol1) - coef(mlr_chol2)[1:4])/coef(mlr_chol2)[1:4] # change < 20%
## (Intercept)
                       dbp racechinese
                                      raceindian
##
      2.965828
                 -1.347773
                              1.957792
                                          3.026647
# no notable change in coeffs, gender is not a confounder
Now, we can try adding sbp & bmi to mlr_chol1 and see what happens to the coefficients. We will use
update() function here.
mlr_chol3 = update(mlr_chol1, . ~ . + sbp)
summary(mlr_chol3) # higher AIC, sbp insig.
##
## Call:
## glm(formula = chol ~ dbp + race + sbp, data = coronary)
##
## Deviance Residuals:
##
       Min
                  1Q
                         Median
                                       3Q
                                                Max
## -2.12850 -0.71572 -0.03242
                                  0.59676
                                            2.77189
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
                          0.529556
## (Intercept) 3.269724
                                     6.174 3.78e-09 ***
                          0.010281
                                     2.916 0.003963 **
## dbp
              0.029978
## racechinese 0.357407
                         0.183561
                                     1.947 0.052963 .
                                     3.516 0.000545 ***
## raceindian 0.705445
                          0.200635
## sbp
              0.000958
                          0.007005
                                     0.137 0.891365
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.094256)
```

```
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.38 on 195 degrees of freedom
## AIC: 592.53
## Number of Fisher Scoring iterations: 2
coef(mlr_chol3); coef(mlr_chol1)
## (Intercept)
                       dbp racechinese
                                        raceindian
                                                             sbp
## 3.2697237312 0.0299783153 0.3574065705 0.7054452332 0.0009580065
## (Intercept)
                      dbp racechinese raceindian
## 3.29802826 0.03110811 0.35996365 0.71369024
100 * (coef(mlr_chol1) - coef(mlr_chol3)[1:4])/coef(mlr_chol3)[1:4] # change < 20%
## (Intercept)
                      dbp racechinese raceindian
   0.8656550 3.7687027 0.7154536
                                     1.1687670
# no notable change in coeffs, sbp is not a confounder
mlr chol4 = update(mlr chol1, . ~ . + bmi)
summary(mlr_chol4) # slighly higher AIC, bmi insig.
##
## Call:
## glm(formula = chol ~ dbp + race + bmi, data = coronary)
## Deviance Residuals:
       Min
                  1Q
                       Median
                                     3Q
                                             Max
## -2.18698 -0.73076 -0.01935
                               0.63476
                                          2.91524
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.870859 1.245373 3.911 0.000127 ***
              ## racechinese 0.356642 0.181757 1.962 0.051164 .
## raceindian 0.724716
                        0.190625
                                   3.802 0.000192 ***
             -0.038530
                         0.028099 -1.371 0.171871
## bmi
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 1.083909)
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 211.36 on 195 degrees of freedom
## AIC: 590.63
##
## Number of Fisher Scoring iterations: 2
coef(mlr_chol4); coef(mlr_chol1)
## (Intercept)
                     dbp racechinese raceindian
## 4.87085865 0.02950027 0.35664168 0.72471631 -0.03853042
## (Intercept)
                     dbp racechinese raceindian
## 3.29802826 0.03110811 0.35996365 0.71369024
```

```
100 * (coef(mlr_chol1) - coef(mlr_chol4)[1:4])/coef(mlr_chol4)[1:4] # change < 20%
## (Intercept)
                       dbp racechinese raceindian
## -32.290619
                             0.931459
                                       -1.521432
                  5.450250
# no notable change in coeffs of other vars (ignore intercept!)
# bmi is not a confounder
Our chosen model:
    mlr_chol1: chol ~ dbp + race
summary(mlr_chol1)
##
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
## Deviance Residuals:
      Min
                1Q
                     Median
                                  3Q
                                          Max
## -2.1378 -0.7068 -0.0289 0.5997
                                       2.7778
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.298028  0.486213  6.783 1.36e-10 ***
                                   5.096 8.14e-07 ***
              0.031108 0.006104
## racechinese 0.359964
                        0.182149
                                    1.976 0.049534 *
## raceindian 0.713690
                        0.190883
                                   3.739 0.000243 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 1.088777)
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 213.40 on 196 degrees of freedom
## AIC: 590.55
##
## Number of Fisher Scoring iterations: 2
Confint(mlr_chol1) # 95% CI of the coefficients
##
                Estimate
                               2.5 %
                                         97.5 %
## (Intercept) 3.29802826 2.345067995 4.25098852
              0.03110811 0.019143668 0.04307255
## racechinese 0.35996365 0.002958566 0.71696873
## raceindian 0.71369024 0.339566932 1.08781356
Compare this model with the no-variable model and all-variable model by LR test and AIC comparison,
# LR test
anova(slr_chol0, mlr_chol1, test = "LRT") # sig. better than no var at all!
## Analysis of Deviance Table
## Model 1: chol ~ 1
## Model 2: chol ~ dbp + race
   Resid. Df Resid. Dev Df Deviance Pr(>Chi)
## 1
         199
                  278.77
```

```
## 2
           196
                   213.40 3
                               65.373 5.755e-13 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# model with no var at all is called Null Model
anova(mlr_chol, mlr_chol1, test = "LRT") # no sig. dif with all vars model,
## Analysis of Deviance Table
##
## Model 1: chol ~ sbp + dbp + bmi + race
## Model 2: chol ~ dbp + race
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
           194
## 1
                   211.34
## 2
           196
                   213.40 -2 -2.0593
                                        0.3886
# model with 2 vars (dbp arepsilon race) is just as good as full model (with all the vars)
# model with all vars is called Saturated Model
# AIC
AIC(slr_chol0, mlr_chol1, mlr_chol)
##
             df
                     AIC
## slr_chol0 2 637.9921
## mlr_chol1 5 590.5459
## mlr_chol
              7 592.6065
# our final model has the lowest AIC
```

#### 3.2.3.5 Multicollinearity, MC

Multicollinearity is the problem of repetitive/redundant variables – high correlations between predictors. MC is checked by Variance Inflation Factor (VIF). VIF > 10 indicates MC problem.

#### 3.2.3.6 Interaction, \*

Interaction is the predictor variable combination that requires interpretation of regression coefficients separately based on the levels of the predictor (e.g. separate analysis for each race group, Malay vs Chinese vs Indian). This makes interpreting our analysis complicated. So, most of the time, we pray not to have interaction in our regression model.

```
summary(glm(chol ~ dbp*race, data = coronary)) # dbp*race not sig.
##
## Call:
## glm(formula = chol ~ dbp * race, data = coronary)
##
## Deviance Residuals:
##
        Min
                   1Q
                         Median
                                        3Q
                                                 Max
## -2.10485 -0.77524
                       -0.02423
                                   0.58059
                                             2.74380
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept)
                   2.11114
                              0.92803
                                      2.275 0.024008 *
## dbp
                   0.04650
                              0.01193 3.897 0.000134 ***
## racechinese
                   1.95576
                              1.28477
                                      1.522 0.129572
## raceindian
                                      1.920 0.056266 .
                   2.41530
                              1.25766
## dbp:racechinese -0.02033
                              0.01596 -1.273 0.204376
## dbp:raceindian -0.02126
                              0.01529 -1.391 0.165905
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for gaussian family taken to be 1.087348)
##
##
      Null deviance: 278.77 on 199 degrees of freedom
## Residual deviance: 210.95 on 194 degrees of freedom
## AIC: 592.23
##
## Number of Fisher Scoring iterations: 2
# in R, it is easy to fit interaction by *
# dbp*race will automatically include all vars involved i.e. equal to
# glm(chol ~ dbp + race + dbp:race, data = coronary)
# use : to just include just the interaction
```

There is no interaction here because the included interaction term was insignificant.

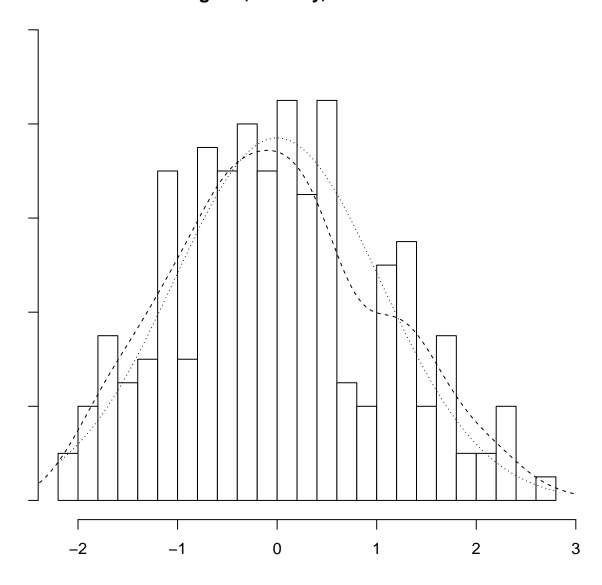
### 3.2.4 Model fit assessment: Residuals

#### 3.2.4.1 Histogram

Raw residuals: Normality assumption.

```
rraw_chol = resid(mlr_chol1) # unstandardized
multi.hist(rraw_chol)
```

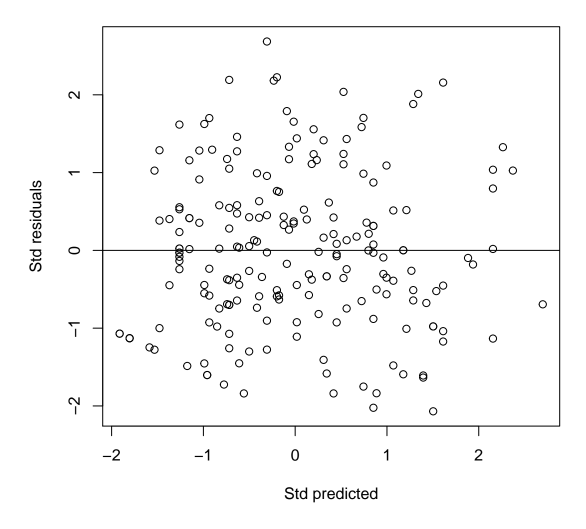
## Histogram, Density, and Normal Fit



### 3.2.4.2 Scatter plots

 $Standardized\ residuals\ vs\ Standardized\ predicted\ values:\ Overall-normality,\ linearity\ and\ equal\ variance\ assumptions.$ 

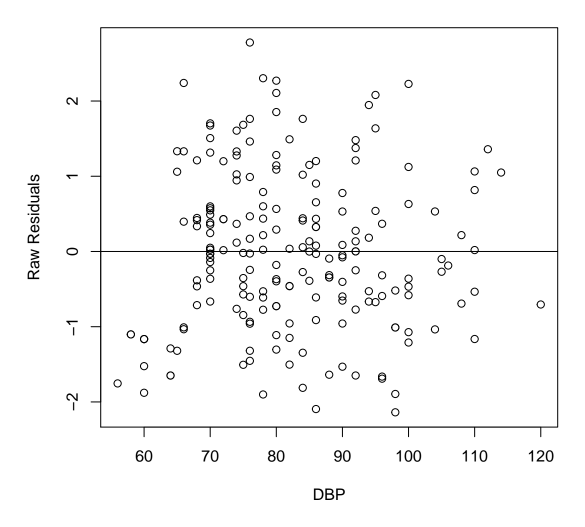
```
rstd_chol = rstandard(mlr_chol1) # standardized residuals
pstd_chol = scale(predict(mlr_chol1)) # standardized predicted values
plot(rstd_chol ~ pstd_chol, xlab = "Std predicted", ylab = "Std residuals")
abline(0, 0) # normal, linear, equal variance
```



The dots should form elliptical/oval shape (normality) and scattered roughly equal above and below the zero line (equal variance). Both these indicate linearity.

Raw residuals vs Numerical predictor by each predictors: Linearity assumption.

```
plot(rraw_chol ~ coronary$dbp, xlab = "DBP", ylab = "Raw Residuals")
abline(0, 0)
```



### 3.2.5 Interpretation

Now we have decided on our final model, rename the model,

```
# rename the selected model
mlr_chol_final = mlr_chol1
and interpret the model,
summary(mlr_chol_final)
##
## Call:
## glm(formula = chol ~ dbp + race, data = coronary)
##
## Deviance Residuals:
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -2.1378 -0.7068 -0.0289
                                0.5997
                                          2.7778
```

```
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
  (Intercept) 3.298028
                          0.486213
                                     6.783 1.36e-10 ***
##
## dbp
               0.031108
                          0.006104
                                     5.096 8.14e-07 ***
                                     1.976 0.049534 *
## racechinese 0.359964
                          0.182149
## raceindian 0.713690
                                     3.739 0.000243 ***
                          0.190883
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for gaussian family taken to be 1.088777)
##
##
       Null deviance: 278.77
                             on 199
                                      degrees of freedom
## Residual deviance: 213.40 on 196
                                      degrees of freedom
  AIC: 590.55
##
## Number of Fisher Scoring iterations: 2
Confint(mlr_chol_final) # 95% CI of the coefficients
##
                                2.5 %
                                          97.5 %
                 Estimate
## (Intercept) 3.29802826 2.345067995 4.25098852
               0.03110811 0.019143668 0.04307255
## racechinese 0.35996365 0.002958566 0.71696873
## raceindian 0.71369024 0.339566932 1.08781356
rsq(mlr_chol_final, adj = T)
```

#### ## [1] 0.2227869

- 1mmHg increase in DBP causes 0.03mmol/L increase in cholestrol, controlling for the effect of race.
- Being Chinese causes 0.36mmol/L increase in cholestrol in comparison to Malay, controlling for the effect of DBP
- Being Indian causes 0.71mmol/L increase in cholestrol in comparison to Malay, controlling for the effect of DBP.
- DBP and race explains 22.3% variance in cholestrol.

Turn the results into data frames results using broom,

```
tib_mlr = tidy(mlr_chol_final, conf.int = T); tib_mlr
```

```
## # A tibble: 4 x 7
##
     term
                  estimate std.error statistic p.value conf.low conf.high
##
     <chr>>
                     <dbl>
                                <dbl>
                                          <dbl>
                                                    <dbl>
                                                             <dbl>
                                                                        <dbl>
                    3.30
                             0.486
                                           6.78 1.36e-10
                                                           2.35
                                                                       4.25
## 1 (Intercept)
## 2 dbp
                    0.0311
                             0.00610
                                           5.10 8.14e- 7
                                                           0.0191
                                                                       0.0431
## 3 racechinese
                    0.360
                                           1.98 4.95e- 2
                             0.182
                                                           0.00296
                                                                       0.717
## 4 raceindian
                    0.714
                             0.191
                                           3.74 2.43e- 4
                                                           0.340
                                                                       1.09
```

Display the results using kable in a nice table,

knitr::kable(tib\_mlr)

term	estimate	std.error	statistic	p.value	conf.low	conf.high
(Intercept)	3.2980283	0.4862132	6.783091	0.0000000	2.3450680	4.2509885
dbp	0.0311081	0.0061044	5.095998	0.0000008	0.0191437	0.0430726
racechinese	0.3599636	0.1821488	1.976207	0.0495342	0.0029586	0.7169687

term	estimate	std.error	statistic	p.value	conf.low	conf.high
raceindian	0.7136902	0.1908827	3.738893	0.0002425	0.3395669	1.0878136

We can export the results into a .csv file for use later (e.g. to prepare a table for journal articles etc.),

```
write.csv(tib_mlr, "mlr_final.csv")
```

### 3.2.6 Model equation

Cholestrol level in mmol/L can be predicted by its predictors as given by,

```
chol = 3.30 + 0.03 \times dbp + 0.36 \times race \ (chinese) + 0.71 \times race \ (indian)
```

#### 3.2.7 Prediction

It is easy to predict in R using our fitted model above. First we view the predicted values for our sample,

```
coronary$pred_chol = predict(mlr_chol_final)
head(coronary)

## sbp dbp chol bmi race gender pred_chol
## 1 106 68 6.5725 38.9 indian woman 6.127070
## 2 130 78 6.3250 37.8 malay woman 5.724461
## 3 136 84 5 9675 40 5 malay woman 5 911109
```

## 3 136 84 5.9675 40.5 malay woman 5.911109 ## 4 138 100 7.0400 37.6 malay woman 6.408839 ## 5 115 85 6.6550 40.3 indian man 6.655908 ## 6 124 72 5.9675 37.6 malay man 5.537812

Now let us try predicting for any values for dbp and race,

```
str(coronary[c("dbp", "race")])
```

```
## 'data.frame': 200 obs. of 2 variables:
## $ dbp : num 68 78 84 100 85 72 80 70 85 70 ...
## $ race: Factor w/ 3 levels "malay", "chinese", ..: 3 1 1 1 3 1 1 2 2 2 ...
# simple, dbp = 90, race = indian
predict(mlr_chol_final, list(dbp = 90, race = "indian"))
```

## 6.811448

More data points

```
new_data = data.frame(dbp = c(90, 90, 90), race = c("malay", "chinese", "indian"))
new_data
```

```
## dbp race
## 1 90 malay
## 2 90 chinese
## 3 90 indian
```

```
predict(mlr_chol_final, new_data)
```

```
## 1 2 3
## 6.097758 6.457722 6.811448
```

```
new_data$pred_chol = predict(mlr_chol_final, new_data)
new_data
```

```
## dbp race pred_chol
## 1 90 malay 6.097758
## 2 90 chinese 6.457722
## 3 90 indian 6.811448
```

### 4 Exercises

- 1. Present the results in a table (follow Arifin et al. (2016))
- 2. Obtain the coefficient for 5mmHg increase in DBP.
- 3. Add age to the multivariable model. What happens?

### References

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