# LDA final project

Kaitlin Maciejewski, Morgan de Ferrante, Bingnan Li, Volha Tryputsen

## Contents

Exploratory	T
Smoking vs. Age, Sex	1
(1): Smoking $\sim$ age, sex	1
Is there a relationship between age and smoking status?	1
Does this relationship differ by sex?	2
(2) number of cigarettes $\sim$ age, sex	2
Is there a relationship between the number of cigarettes smoked per day and age?	2
All	2
Smokers only	2
Does this relationship differ by sex?	3
All	3
Smokers only	3
Smoking vs. health outcomes	3
(1) The relationship between current smoking status and systolic blood pressure	3
smoking ~ sysbp	3
smoking $\sim$ sysbp, sex	3
(2) The relationship between current smoking status and diastolic blood pressure	4
smoking $\sim$ diabp	4
smoking $\sim$ diabp, sex	4
(3) The relationship between current smoking status and serum total cholesterol	5
smoking $\sim$ totchol	5
smoking $\sim$ totchol, sex [!!!]	5
Cor plot	6
Missingness	7
a	_
Summary	8
Models	8
(1) Is there a relationship between age and smoking status? Does this relationship differ by sex? .	9
(2) Is there a relationship between the number of cigarettes smoked per day and age? Does this	
relationship differ by sex?	9
(3) Totchol and cursmoke	13
(4) Sysbp and cursmoke	15
(5) Diabp and cursmoke	16

## Exploratory

Smoking vs. Age, Sex

(1): Smoking  $\sim$  age, sex

Is there a relationship between age and smoking status?

ANS: Yes, the proportion of smokers decreases with the age.

```
data$sex <- recode(data$sex, `1` = "male", `2` = "female")
data$cursmoke <- recode(data$cursmoke, `1` = "yes", `0` = "no")

smoke_v_age = data %>%
   select(cursmoke, age) %>%
   ggplot(aes(x = age, y = cursmoke)) +
   geom_jitter(height = 0.1, alpha = 0.1) +
   geom_smooth(lwd = 1.5) +
   theme_bw()
```

#### Does this relationship differ by sex?

ANS: There is a higher proportion of smoker among men compared to women as both age ,but there is no interaction between age and sex.

```
smoke_age_sex = data %>%
select(cursmoke, age, sex) %>%
ggplot(aes(x = age, y = cursmoke, group = sex, color = sex)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

## (2) number of cigarettes ~ age, sex

Is there a relationship between the number of cigarettes smoked per day and age?

ANS: Yes, number of sigarets smoked per day stays constant for 30-50 years old and decreases with age after 50 years old.

#### All

```
n_c_age_all = data %%%
select(cigpday, age) %>%
ggplot(aes(x = age, y = cigpday)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

#### Smokers only

```
n_c_age_smoke = data %>%
select(cigpday, age) %>%
filter(cigpday > 0) %>%
ggplot(aes(x = age, y = cigpday)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

#### Does this relationship differ by sex?

ANS: There is sex effect (men smoke higer number of sigarets per day than women across age), but there is no sex and age interaction.

#### All

```
n_c_age_s_all = data %>%
  select(cigpday, age, sex) %>%
  ggplot(aes(x = age, y = cigpday, group = sex, color = sex)) +
  geom_jitter(height = 0.1, alpha = 0.1) +
  geom_smooth(lwd = 1.5) +
  theme_bw()
```

#### Smokers only

```
n_c_age_s_smoke = data %>%
  select(cigpday, age, sex) %>%
  filter(cigpday > 0) %>%
  ggplot(aes(x = age, y = cigpday, group = sex, color = sex)) +
  geom_jitter(height = 0.1, alpha = 0.1) +
  geom_smooth(lwd = 1.5) +
  theme_bw()
```

## Smoking vs. health outcomes

(1) The relationship between current smoking status and systolic blood pressure.

```
smoking ~ sysbp
```

ANS: Proportion of smokers decreases with increase of systolic blood presure

```
smoke_sbp = data %>%
select(cursmoke, sysbp) %>%
ggplot(aes(x = sysbp, y = cursmoke)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

ANS: slightly higher sysbp for non-smokers

```
smoke_sysbp_status = data %>%
select(cursmoke, sysbp) %>%
mutate(cursmoke = factor(cursmoke)) %>%
ggplot(aes(y = sysbp, x = cursmoke)) +
geom_boxplot(outlier.colour = "white") +
theme_bw()
```

#### smoking ~ sysbp, sex

ANS: Proportion of smokers decreases with increase of systolic blood presure; the proportion is higher for men (sex effect).

```
smoke_sysbp_sex = data %>%
  select(cursmoke, sysbp, sex) %>%
  ggplot(aes(x = sysbp, y = cursmoke, group = sex, color = sex)) +
  geom_jitter(height = 0.1, alpha = 0.1) +
  geom_smooth(lwd = 1.5) +
  theme_bw()
```

ANS: no differences in sysbp between male and female smokers and non-smokers

## (2) The relationship between current smoking status and diastolic blood pressure.

#### smoking ~ diabp

ANS: Proportion of smokers decreases with increase of diastolic blood presure for BP=100 ad then proportion increases again (latter could be due to not enough data)

```
smoke_dbp = data %>%
  select(cursmoke, diabp) %>%
  ggplot(aes(x = diabp, y = cursmoke)) +
  geom_jitter(height = 0.1, alpha = 0.1) +
  geom_smooth(lwd = 1.5) +
  theme_bw()
```

ANS: no difference

```
smoke_dbp_box = data %>%
select(cursmoke, diabp) %>%
mutate(cursmoke = factor(cursmoke)) %>%
ggplot(aes(y = diabp, x = cursmoke)) +
geom_boxplot(outlier.colour = "white") +
theme_bw()
```

#### smoking ~ diabp, sex

ANS: Proportion of smokers decreases with increase of diastolic blood presure; the proprtions are higher for men (sex effect).

```
smoke_dbp_s = data %%%
select(cursmoke, diabp, sex) %>%
ggplot(aes(x = diabp, y = cursmoke, group = sex, color = sex)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

ANS: no difference

## (3) The relationship between current smoking status and serum total cholesterol.

#### smoking ~ totchol

ANS: Proportion of smokers slightly decreases with increase of total cholesterol values

```
smoke_tc = data %>%
select(cursmoke, totchol) %>%
ggplot(aes(x = totchol, y = cursmoke)) +
geom_jitter(height = 0.1, alpha = 0.1) +
geom_smooth(lwd = 1.5) +
theme_bw()
```

ANS: no difference

```
smoke_tc_bp = data %>%
select(cursmoke, totchol) %>%
mutate(cursmoke = factor(cursmoke)) %>%
ggplot(aes(y = totchol, x = cursmoke)) +
geom_boxplot(outlier.colour = "white") +
theme_bw()
```

```
smoking ~ totchol, sex [!!!]
```

ANS: Proportion of smokers has nonlinear relationship with total cholesterol for women; proprtions increases with increase in total cholesterol for men (sex by totchol interaction effect).

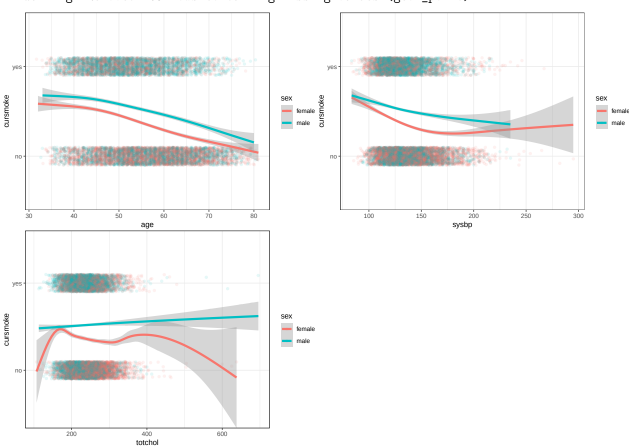
```
smoke_tc_sex = data %>%
  select(cursmoke, totchol, sex) %>%
  ggplot(aes(x = totchol, y = cursmoke, group = sex, color = sex)) +
  geom_jitter(height = 0.1, alpha = 0.1) +
  geom_smooth(lwd = 1.5) +
  theme_bw()
```

ANS: no difference

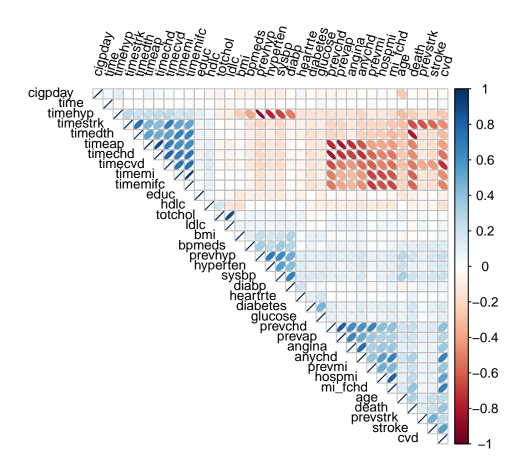
```
smoke_age_sex
smoke_sysbp_sex
smoke_tc_sex
```

## Warning: Removed 409 rows containing non-finite values (stat\_smooth).

## Warning: Removed 409 rows containing missing values (geom\_point).



## Cor plot



## Missingness

```
prop <- round(colSums(is.na(data))/dim(data)[1], 3)
knitr::kable(sort(prop, decreasing = TRUE)[1:9], col.names = "Proportion of NAs")</pre>
```

	Proportion of NAs
hdlc	0.740
ldlc	0.740
glucose	0.124
bpmeds	0.051
totchol	0.035
educ	0.025
cigpday	0.007
bmi	0.004
heartrte	0.001

```
prob.data <- data %>%
    group_by(period) %>%
    summarise(sysbp_prob = sum(sysbp, na.rm = TRUE)/n())
prob.data
```

```
## # A tibble: 3 x 2
## period sysbp_prob
```

```
<int>
                  <dbl>
## 1
                   133.
          1
## 2
          2
                   137.
## 3
          3
                   140.
table(data$period)
##
##
      1
           2
## 4434 3930 3263
```

## **Summary**

```
summary(age)
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
           48.00
                    54.00
                           54.79
                                   62.00
                                           81.00
table(data$cursmoke)
##
    no yes
## 6598 5029
table(data$sex)
##
## female
           male
    6605
           5022
summary(cigpday)
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                                   NA's
                                            Max.
##
     0.00
             0.00
                     0.00
                             8.25
                                   20.00
                                           90.00
summary(totchol)
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                                   NA's
                                            Max.
##
    107.0 210.0
                    238.0
                           241.2
                                   268.0
                                           696.0
                                                     409
summary(diabp)
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
           75.00
                   82.00
                            83.04
                                   90.00 150.00
summary(sysbp)
##
     Min. 1st Qu. Median
                           Mean 3rd Qu.
                                            Max.
     83.5 120.0 132.0 136.3 149.0
                                           295.0
```

## Models

```
library(lme4)
library(dplyr)
```

(1) Is there a relationship between age and smoking status? Does this relationship differ by sex?

```
my.data <- read.csv("../final_data/frmgham2.csv")</pre>
library(gee)
model.q1 <- gee(CURSMOKE ~ AGE + as.factor(SEX) + as.factor(educ)</pre>
                + BMI + DIABETES + HEARTRTE + PREVCHD + PREVSTRK
                + PREVHYP + TIMEDTH,
                id = RANDID,
                data = my.data,
                family=binomial,
                 corstr = "unstructured")
##
        (Intercept)
                                AGE as.factor(SEX)2 as.factor(educ)2
##
      5.2810549757
                      -0.0570318456
                                       -0.7246241605
                                                         0.0503157656
## as.factor(educ)3 as.factor(educ)4
                                                 BMI
                                                             DIABETES
      -0.2263406725 -0.2104931713 -0.0940121564 -0.1114780579
##
##
          HEARTRTE
                            PREVCHD
                                            PREVSTRK
                                                              PREVHYP
      0.0173911758 -0.0569530253
                                       -0.2633153119
##
                                                        -0.2252617826
##
            TIMEDTH
##
      -0.0001021991
knitr::kable(summary(model.q1)$coefficients[,c(1,4,5)], digits = 3)
```

F 1.40		
5.149	0.278	18.543
-0.051	0.002	-21.106
-0.716	0.060	-11.892
0.083	0.071	1.168
-0.182	0.086	-2.116
-0.212	0.097	-2.185
-0.084	0.007	-12.152
-0.060	0.102	-0.595
0.009	0.001	6.031
-0.259	0.090	-2.872
-0.189	0.176	-1.073
-0.049	0.041	-1.187
0.000	0.000	-6.341
	-0.716 0.083 -0.182 -0.212 -0.084 -0.060 0.009 -0.259 -0.189 -0.049	-0.051 0.002 -0.716 0.060 0.083 0.071 -0.182 0.086 -0.212 0.097 -0.084 0.007 -0.060 0.102 0.009 0.001 -0.259 0.090 -0.189 0.176 -0.049 0.041

```
model.q1[["working.correlation"]]

## [,1] [,2] [,3]

## [1,] 1.0000000 0.7563191 0.5136670

## [2,] 0.7563191 1.0000000 0.5713635

## [3,] 0.5136670 0.5713635 1.0000000
```

(2) Is there a relationship between the number of cigarettes smoked per day and age? Does this relationship differ by sex?

If we think cig per day as count data, it follows poisson distribution. Then we can fit GEE model as well:

```
##
       (Intercept)
                              AGE as.factor(SEX)2 as.factor(educ)2
      4.914149e+00
                     -3.395990e-02
                                   -7.680017e-01
                                                    5.765596e-02
## as.factor(educ)3 as.factor(educ)4
                                                         DIABETES
                                              BMI
##
     -1.227240e-01 -8.833837e-02
                                  -4.493994e-02 -1.608903e-01
                                         PREVSTRK
##
          HEARTRTE
                          PREVCHD
                                                          PREVHYP
##
      1.281640e-02 -9.883049e-02 -1.115583e-01 -1.079832e-01
           TIMEDTH
##
##
     -5.152275e-05
```

knitr::kable(summary(model.q2\_1)\$coefficients[,c(1,4,5)], digits = 3)

	Estimate	Robust S.E.	Robust z
(Intercept)	4.523	0.170	26.626
AGE	-0.024	0.001	-15.826
as.factor(SEX)2	-0.795	0.039	-20.223
as.factor(educ)2	0.096	0.044	2.172
as.factor(educ)3	-0.072	0.057	-1.245
as.factor(educ)4	-0.067	0.063	-1.059
BMI	-0.039	0.005	-8.032
DIABETES	-0.110	0.072	-1.536
HEARTRTE	0.007	0.001	8.453
PREVCHD	-0.241	0.066	-3.635
PREVSTRK	-0.130	0.110	-1.181
PREVHYP	-0.017	0.028	-0.587
TIMEDTH	0.000	0.000	-5.505

## model.q2\_1[["working.correlation"]]

```
## [,1] [,2] [,3]
## [1,] 1.0000000 0.7425318 0.5006078
## [2,] 0.7425318 1.0000000 0.6615306
## [3,] 0.5006078 0.6615306 1.0000000
```

Using mixed effect model using cig per day instead of smoking status:

```
data = my.data)
summary(model.saturated)
## Linear mixed model fit by REML ['lmerMod']
## Formula:
## CIGPDAY ~ as.factor(SEX) + AGE + BPMEDS + as.factor(educ) + TOTCHOL +
##
      BMI + GLUCOSE + DIABETES + HEARTRTE + PREVAP + PREVCHD +
      PREVMI + PREVSTRK + STROKE + PREVHYP + (1 | RANDID)
##
##
     Data: my.data
##
## REML criterion at convergence: 67692.5
##
## Scaled residuals:
##
      Min
               1Q Median
                               ЗQ
                                      Max
## -4.4444 -0.3150 -0.1078 0.2067 6.2548
##
## Random effects:
## Groups
            Name
                        Variance Std.Dev.
## RANDID
                                 9.787
             (Intercept) 95.78
## Residual
                        35.77
                                 5.981
## Number of obs: 9310, groups: RANDID, 4213
## Fixed effects:
                    Estimate Std. Error t value
## (Intercept)
                   23.237579 1.393589 16.675
## as.factor(SEX)2 -7.062484 0.339110 -20.827
## AGE
                   -0.178024 0.013735 -12.962
## BPMEDS
                    0.147068 0.340662
                                         0.432
## as.factor(educ)2 0.602137
                               0.400578
                                          1.503
## as.factor(educ)3 -0.984441 0.482907 -2.039
## as.factor(educ)4 -1.015103 0.548951 -1.849
## TOTCHOL
                    0.011407 0.002557
                                         4.461
## BMI
                   -0.313257 0.035227 -8.893
## GLUCOSE
                   -0.010167 0.004203 -2.419
## DIABETES
                               0.570383 -0.446
                   -0.254321
## HEARTRTE
                    0.070622 0.008232
                                         8.579
## PREVAP
                   -3.058426 0.985947 -3.102
## PREVCHD
                    0.948740 1.050262
                                         0.903
## PREVMI
                   -2.594434 0.894896 -2.899
## PREVSTRK
                   -1.029740
                               0.917968 - 1.122
## STROKE
                    0.943808
                               0.586941
                                         1.608
## PREVHYP
                   -0.205972
                               0.240530 -0.856
#using variables that selected
model.mixed2 <- lmer(CIGPDAY~ AGE + as.factor(SEX) + SYSBP</pre>
                     + DIABP + TOTCHOL + as.factor(educ)
                     + (1|RANDID),
                     data = my.data,
                     na.action = "na.omit")
summary(model.mixed2)
## Linear mixed model fit by REML ['lmerMod']
## Formula:
## CIGPDAY ~ AGE + as.factor(SEX) + SYSBP + DIABP + TOTCHOL + as.factor(educ) +
```

```
(1 | RANDID)
##
##
     Data: my.data
##
## REML criterion at convergence: 78607.2
## Scaled residuals:
             10 Median
      Min
                             30
                                    Max
## -4.6343 -0.3025 -0.1015 0.1642 6.6280
##
## Random effects:
## Groups
           Name
                       Variance Std.Dev.
## RANDID
            (Intercept) 97.30
                               9.864
## Residual
                       36.52
                               6.043
## Number of obs: 10868, groups: RANDID, 4306
##
## Fixed effects:
##
                   Estimate Std. Error t value
## (Intercept)
                  -0.199583 0.012148 -16.429
## AGE
## as.factor(SEX)2 -6.448164 0.332812 -19.375
## SYSBP
                  -0.001364 0.006670 -0.205
## DIABP
                  -0.018833 0.011514 -1.636
## TOTCHOL
                   0.011959 0.002373
                                       5.039
## as.factor(educ)2 1.055542 0.392720
                                       2.688
## as.factor(educ)3 -0.625323 0.474072 -1.319
## as.factor(educ)4 -0.827192  0.539360 -1.534
##
## Correlation of Fixed Effects:
##
             (Intr) AGE
                          a.(SEX SYSBP DIABP TOTCHO as.()2 as.()3
## AGE
             -0.490
## as.fc(SEX)2 -0.144 0.019
## SYSBP
              0.023 -0.460 -0.064
## DIABP
             -0.452 0.314 0.076 -0.679
## TOTCHOL
             -0.368 -0.095 -0.094 0.047 -0.154
## as.fctr(d)2 -0.225  0.110 -0.055  0.008  0.001  0.002
## as.fctr(d)3 -0.161 0.051 -0.092 0.028 -0.004 -0.007 0.351
library(dplyr)
##################
# Initial Models #
##################
# totchol_fit <- gee(totchol ~ cursmoke + age + factor(sex) + factor(educ) + bmi +
             diabetes + heartrie + prevchd + prevhyp + prevstrk + death,
#
            id = randid,
#
            family = "gaussian",
#
            na.action = "na.omit")
# round(2 * pnorm(abs(coef(summary(totchol_fit))[,5]), lower.tail = FALSE), 3)
#
#
```

```
# sysbp_fit <- gee(sysbp ~ cursmoke + age + factor(sex) + factor(educ) + bmi +
#
                       diabetes + heartrie + prevchd + prevstrk + death,
#
                     id = randid,
#
                     family = "qaussian",
#
                     na.action = "na.omit")
# round(2 * pnorm(abs(coef(summary(sysbp_fit))[,5]), lower.tail = FALSE), 3)
#
# diabp fit <- qee(diabp ~ cursmoke + age + factor(sex) + factor(educ) + bmi +
                       diabetes + heartrie + prevchd + prevstrk +death,
#
#
                     id = randid,
#
                     family = "gaussian",
#
                     na.action = "na.omit")
#
# round(2 * pnorm(abs(coef(summary(diabp_fit))[,5]), lower.tail = FALSE), 3)
```

## (3) Totchol and cursmoke

```
# Models After Removing Non Significant Terms #
totchol_fit <- gee(totchol ~ cursmoke + age + factor(sex) + bmi +</pre>
                 diabetes + heartrte + prevhyp ,
                id = randid,
                family = "gaussian",
                corstr = "unstructured",
                na.action = "na.omit")
##
   (Intercept)
                cursmoke
                                age factor(sex)2
                                                      bmi
##
   162.1032803
                2.3672102
                           0.6268036
                                     12.3710689
                                                 0.7814908
##
      diabetes
                heartrte
                             prevhyp
                0.1891382
                           5.2800067
##
    -5.8194286
knitr::kable(summary(totchol_fit)$coefficients[,c(1,4,5)], digits = 3)
```

	Estimate	Robust S.E.	Robust z
(Intercept)	149.257	5.279	28.274
cursmoke	3.430	1.009	3.399
age	0.648	0.053	12.309
factor(sex)2	13.474	1.219	11.050
bmi	1.404	0.144	9.748
diabetes	-6.628	2.638	-2.513
heartrte	0.117	0.032	3.626
prevhyp	3.451	0.903	3.820

knitr::kable(round(2 \* pnorm(abs(coef(summary(totchol\_fit))[,5]), lower.tail = FALSE), 3))

	Х
(Intercept)	0.000
cursmoke	0.001
age	0.000

```
        x

        factor(sex)2
        0.000

        bmi
        0.000

        diabetes
        0.012

        heartrte
        0.000

        prevhyp
        0.000
```

```
QIC(totchol_fit)
##
        QIC
## 84588.89
totchol_fit2 <- gee(totchol ~ cursmoke + age + factor(sex) + bmi +</pre>
                     diabetes + heartrte + prevhyp ,
                   id = randid,
                   family = "gaussian",
                   corstr = "exchangeable",
                   na.action = "na.omit")
##
    (Intercept)
                    cursmoke
                                       age factor(sex)2
                                                                  bmi
##
    162.1032803
                   2.3672102
                                 0.6268036
                                             12.3710689
                                                            0.7814908
##
       diabetes
                    heartrte
                                   prevhyp
     -5.8194286
                                 5.2800067
##
                   0.1891382
knitr::kable(summary(totchol_fit2)$coefficients[,c(1,4,5)], digits = 3)
```

	Estimate	Robust S.E.	Robust z
(Intercept)	157.548	5.162	30.518
cursmoke	3.824	1.035	3.696
age	0.402	0.051	7.824
factor(sex)2	13.291	1.222	10.874
bmi	1.656	0.143	11.578
diabetes	-7.648	2.322	-3.293
heartrte	0.118	0.033	3.607
prevhyp	2.423	0.894	2.710

knitr::kable(round(2 \* pnorm(abs(coef(summary(totchol\_fit2))[,5]), lower.tail = FALSE), 3))

```
\mathbf{X}
(Intercept)
               0.000
cursmoke
               0.000
age
               0.000
factor(sex)2
               0.000
bmi
               0.000
diabetes
               0.001
heartrte
               0.000
               0.007
prevhyp
```

```
QIC(totchol_fit2)
```

## QIC

## (4) Sysbp and cursmoke

```
sysbp_fit <- gee(sysbp ~ cursmoke + age + factor(sex) + bmi +</pre>
                  diabetes + heartrte + prevchd + prevstrk + death,
                id = randid,
                family = "gaussian",
                corstr = "unstructured",
                na.action = "na.omit")
                                     age factor(sex)2
##
  (Intercept)
                  cursmoke
                                                              bmi
    40.1191453
                               0.7236371 2.9078810
                                                         1.3070103
##
                -1.1246140
##
      diabetes
                                                             death
                  heartrte
                                 prevchd
                                            prevstrk
##
      4.8530466
                  0.2464253
                               2.4405511
                                            8.6116353
                                                        7.0492511
knitr::kable(summary(sysbp_fit)$coefficients[,c(1,4,5)], digits = 3)
```

	Estimate	Robust S.E.	Robust z
(Intercept)	42.702	2.163	19.740
cursmoke	-0.702	0.433	-1.619
age	0.741	0.022	33.618
factor(sex)2	3.272	0.542	6.042
bmi	1.373	0.062	22.301
diabetes	3.664	1.127	3.250
heartrte	0.175	0.016	11.231
prevchd	1.670	0.876	1.906
prevstrk	4.894	2.004	2.443
death	7.650	0.639	11.967

knitr::kable(round(2 \* pnorm(abs(coef(summary(sysbp\_fit))[,5]), lower.tail = FALSE), 3))

	Х
(Intercept)	0.000
cursmoke	0.105
age	0.000
factor(sex)2	0.000
bmi	0.000
diabetes	0.001
heartrte	0.000
prevchd	0.057
prevstrk	0.015
death	0.000

```
id = randid,
                family = "gaussian",
                corstr = "exchangeable",
                na.action = "na.omit")
                                    age factor(sex)2
##
   (Intercept)
                   cursmoke
                                                              bmi
##
     40.1191453 -1.1246140
                               0.7236371
                                           2.9078810
                                                        1.3070103
##
      diabetes
                  heartrte
                                prevchd
                                           prevstrk
                                                            death
                                                        7.0492511
##
     4.8530466
                  0.2464253
                               2.4405511
                                           8.6116353
knitr::kable(summary(sysbp_fit2)$coefficients[,c(1,4,5)], digits = 3)
```

	Estimate	Robust S.E.	Robust z
(Intercept)	43.000	2.176	19.758
cursmoke	-0.772	0.431	-1.789
age	0.741	0.021	34.477
factor(sex)2	3.321	0.545	6.093
bmi	1.370	0.063	21.783
diabetes	3.173	1.114	2.848
heartrte	0.172	0.016	10.967
prevchd	1.553	0.862	1.800
prevstrk	4.201	2.094	2.006
death	7.794	0.644	12.109

knitr::kable(round(2 \* pnorm(abs(coef(summary(sysbp\_fit2))[,5]), lower.tail = FALSE), 3))

	Х
(Intercept)	0.000
cursmoke	0.074
age	0.000
factor(sex)2	0.000
bmi	0.000
diabetes	0.004
heartrte	0.000
prevchd	0.072
prevstrk	0.045
death	0.000

### QIC(sysbp\_fit2)

```
## QIC
## 68837.55
```

## (5) Diabp and cursmoke

```
corstr = "unstructured",
                na.action = "na.omit")
                     cursmoke factor(sex)2 factor(educ)2 factor(educ)3
##
     (Intercept)
##
      47.3480563
                   -1.0968017
                                 -0.8594048
                                                0.7880155
                                                              0.6921165
## factor(educ)4
                          bmi
                                   diabetes
                                                 heartrte
                                                              prevstrk
      0.5395935
                    0.9078065
                                 -1.6162433
                                                0.1550842
                                                              4.3067909
##
##
          death
##
      2.8662538
knitr::kable(summary(diabp_fit)$coefficients[,c(1,4,5)], digits = 3)
```

	Estimate	Robust S.E.	Robust z
(Intercept)	48.560	1.142	42.511
cursmoke	-0.607	0.243	-2.501
factor(sex)2	-0.591	0.295	-1.999
factor(educ)2	0.864	0.344	2.513
factor(educ)3	0.769	0.400	1.924
factor(educ)4	0.470	0.450	1.043
bmi	0.971	0.035	28.015
diabetes	-1.731	0.598	-2.895
heartrte	0.112	0.009	12.846
prevstrk	1.842	1.207	1.526
death	3.219	0.336	9.582

knitr::kable(round(2 \* pnorm(abs(coef(summary(diabp\_fit))[,5]), lower.tail = FALSE), 3))

	Х
(Intercept)	0.000
cursmoke	0.012
factor(sex)2	0.046
factor(educ)2	0.012
factor(educ)3	0.054
factor(educ)4	0.297
bmi	0.000
diabetes	0.004
heartrte	0.000
prevstrk	0.127
death	0.000

na.action = "na.omit")

```
##
     (Intercept)
                      cursmoke factor(sex)2 factor(educ)2 factor(educ)3
##
      47.3480563
                    -1.0968017
                                  -0.8594048
                                                 0.7880155
                                                               0.6921165
## factor(educ)4
                           bmi
                                    diabetes
                                                  heartrte
                                                                prevstrk
##
       0.5395935
                     0.9078065
                                  -1.6162433
                                                 0.1550842
                                                               4.3067909
##
           death
       2.8662538
##
```

knitr::kable(summary(diabp\_fit2)\$coefficients[,c(1,4,5)], digits = 3)

	Estimate	Robust S.E.	Robust z
(Intercept)	48.526	1.156	41.980
cursmoke	-0.525	0.242	-2.166
factor(sex)2	-0.518	0.298	-1.738
factor(educ)2	0.857	0.346	2.476
factor(educ)3	0.776	0.403	1.927
factor(educ)4	0.531	0.453	1.171
bmi	0.991	0.035	28.079
diabetes	-2.072	0.582	-3.564
heartrte	0.106	0.009	12.102
prevstrk	1.049	1.215	0.863
death	3.317	0.338	9.800

knitr::kable(round(2 \* pnorm(abs(coef(summary(diabp\_fit2))[,5]), lower.tail = FALSE), 3))

	х
(Intercept)	0.000
cursmoke	0.030
factor(sex)2	0.082
factor(educ)2	0.013
factor(educ)3	0.054
factor(educ)4	0.241
bmi	0.000
diabetes	0.000
heartrte	0.000
prevstrk	0.388
death	0.000

### QIC(diabp\_fit2)

## QIC ## 53508