EMM CANTOS

Yi Li

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```
library(regmedint)
```

Warning: package 'regmedint' was built under R version 4.1.2

0. Data Source

Original paper: Vallurupalli M, MacFadyen JG, Glynn RJ, et al. Effects of Interleukin- 1β Inhibition on Incident Anemia: Exploratory Analyses From a Randomized Trial. Ann Intern Med. 2020;172(8):523. doi:10.7326/M19-2945

1. Variables

Treatment: canakinumab use (binary)

Mediator: change in log CRP (continuous)

Outcome: anemia (survival time)

Baseline confounders: age, sex, baseline log CRP, heart failure, hypertension, diabetes

```
## age female base_logCRP chf htn dm
## 60.5374609 0.2471694 1.4949021 0.3819562 0.7898097 0.2033245
```

2. Mediation analyses

(1) No Effect modification by any covariates

```
fit0.0 <- regmedint(data = anemia3,
                    yvar = "anemiayrs",
                    eventvar = "anemiaev",
                    avar = "canakinumab",
                    mvar = "diff_logCRP",
                    cvar = cvar,
                    ## EMM
                    emm_ac_mreg = NULL,
                    emm_ac_yreg = NULL,
                    emm_mc_yreg = NULL,
                    ##
                    mreg = "linear",
                    yreg = "survAFT_weibull",
                    interaction = TRUE,
                    casecontrol = FALSE,
                    a0 = 0,
                    a1 = 1,
                    m_cde = mean(anemia3$diff_logCRP),
                    c_cond = c_cond0)
summary(fit0.0)
```

```
## ### Mediator model
##
## Call:
## lm(formula = diff_logCRP ~ canakinumab + age + female + base_logCRP +
      chf + htn + dm, data = data)
##
##
## Residuals:
               1Q Median
                              3Q
                                     Max
## -4.4422 -0.5246 -0.0366 0.4550 5.6302
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.4554245 0.0624193 7.296 3.23e-13 ***
## canakinumab -0.6504159 0.0194098 -33.510 < 2e-16 ***
       -0.0006661 0.0009536 -0.698 0.4849
## age
```

```
## female
              0.0124121 0.0214418
                                      0.579
## base_logCRP -1.4574217  0.0135301 -107.717  < 2e-16 ***
              0.0928082 0.0191562
                                      4.845 1.29e-06 ***
## htn
               0.0161637
                          0.0230710
                                      0.701
                                              0.4836
## dm
               0.0427669 0.0229648
                                      1.862
                                              0.0626 .
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.8336 on 8294 degrees of freedom
## Multiple R-squared: 0.6082, Adjusted R-squared: 0.6078
## F-statistic: 1839 on 7 and 8294 DF, p-value: < 2.2e-16
## ### Outcome model
##
## Call:
## survival::survreg(formula = Surv(anemiayrs, anemiaev) ~ canakinumab +
      diff_logCRP + canakinumab:diff_logCRP + age + female + base_logCRP +
##
##
      chf + htn + dm, data = data, dist = "weibull")
##
                             Value Std. Error
## (Intercept)
                           7.95064
                                     0.33587 23.67 < 2e-16
                                     0.14936
## canakinumab
                           0.08959
                                              0.60 0.54860
## diff logCRP
                          -0.23037
                                     0.06386 -3.61 0.00031
                                     0.00446 - 12.61 < 2e-16
## age
                          -0.05624
## female
                                     0.09005 -0.08 0.93985
                          -0.00679
## base_logCRP
                          -0.73261
                                     0.08947 -8.19 2.6e-16
## chf
                          -0.63685
                                     0.08163 -7.80 6.1e-15
## htn
                          -0.06529
                                     0.10624 -0.61 0.53883
                          -0.31877
                                     0.09297 -3.43 0.00061
## Log(scale)
                           0.52382
                                     0.02211 23.69 < 2e-16
##
## Scale= 1.69
##
## Weibull distribution
## Loglik(model) = -6073.9 Loglik(intercept only) = -6258.5
## Chisq= 369.15 on 9 degrees of freedom, p= 5.2e-74
## Number of Newton-Raphson Iterations: 7
## n= 8302
##
## ### Mediation analysis
             est
                         se
                                  Z
                                                        lower
                                               р
## cde 0.1788398 0.08779440 2.037030 4.164704e-02 0.0067659649 0.3509137
## pnde 0.1698583 0.08658498 1.961752 4.979133e-02 0.0001548423 0.3395617
## tnie 0.1769992 0.03295536 5.370878 7.835435e-08 0.1124078731 0.2415905
## tnde 0.1970205 0.09481515 2.077943 3.771461e-02 0.0111862011 0.3828547
## pnie 0.1498370 0.04177719 3.586575 3.350502e-04 0.0679552181 0.2317188
       0.3468575 0.08320654 4.168632 3.064328e-05 0.1837756542 0.5099393
## pm
       0.5534728 0.13876434 3.988581 6.646960e-05 0.2814997466 0.8254460
##
## Evaluated at:
## avar: canakinumab
## a1 (intervened value of avar) = 1
## a0 (reference value of avar) = 0
## mvar: diff_logCRP
```

```
## m_cde (intervend value of mvar for cde) = -2.137102
## cvar: age female base_logCRP chf htn dm
## c_cond (covariate vector value) = 60.53746 0 1.494902 0 1 0
##
## Note that effect estimates can vary over m_cde and c_cond values when interaction = TRUE.
```

- (2) Effect modification by baseline logCRP in mediator model, and age & diabetes in outcome model
- a. Specify low and high risk groups' covariate levels

b. Fit regmedint function

```
fit1.1 <- regmedint(data = anemia3,</pre>
                    yvar = "anemiayrs",
                    eventvar = "anemiaev",
                    avar = "canakinumab",
                    mvar = "diff_logCRP",
                    cvar = cvar,
                    ## EMM
                    emm_ac_mreg = c("base_logCRP"),
                    emm_ac_yreg = NULL,
                    emm_mc_yreg = c("age", "dm"),
                    ##
                    mreg = "linear",
                    yreg = "survAFT_weibull",
                    interaction = TRUE,
                    casecontrol = FALSE,
                    a0 = 0,
                    a1 = 1,
                    m_cde = mean(anemia3$diff_logCRP),
                    c_cond = c_cond_low_all)
summary(fit1.1)
```

```
## ### Mediator model
##
## Call:
## lm(formula = diff_logCRP ~ canakinumab + age + female + base_logCRP +
```

```
chf + htn + dm + canakinumab:base_logCRP, data = data)
##
##
## Residuals:
##
                               3Q
      Min
                1Q Median
                                      Max
## -4.4475 -0.5258 -0.0350 0.4525 5.6261
##
## Coefficients:
##
                            Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                           0.4228295 0.0690623
                                                  6.122 9.63e-10 ***
## canakinumab
                          ## age
                          -0.0006421 0.0009539 -0.673
                                                          0.5008
                                                  0.588
## female
                           0.0126025 0.0214422
                                                          0.5567
                          -1.4365841 0.0232393 -61.817 < 2e-16 ***
## base_logCRP
## chf
                           0.0928445 0.0191560
                                                  4.847 1.28e-06 ***
## htn
                                                  0.708
                           0.0163260 0.0230711
                                                          0.4792
## dm
                            0.0427971
                                      0.0229645
                                                  1.864
                                                          0.0624 .
                                                          0.2701
## canakinumab:base_logCRP -0.0314079  0.0284792  -1.103
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.8336 on 8293 degrees of freedom
## Multiple R-squared: 0.6082, Adjusted R-squared: 0.6078
## F-statistic: 1609 on 8 and 8293 DF, p-value: < 2.2e-16
## ### Outcome model
##
## Call:
## survival::survreg(formula = Surv(anemiayrs, anemiaev) ~ canakinumab +
       diff_logCRP + canakinumab:diff_logCRP + age + female + base_logCRP +
       chf + htn + dm + diff_logCRP:age + diff_logCRP:dm, data = data,
##
##
       dist = "weibull")
##
                             Value Std. Error
                                                  z
## (Intercept)
                           7.40059
                                      0.53565 13.82 < 2e-16
                           0.08224
                                      0.14929 0.55
## canakinumab
                                                      0.582
## diff_logCRP
                          -0.48373
                                      0.20525 - 2.36
                                                      0.018
                                      0.00802 -5.98 2.2e-09
## age
                          -0.04799
## female
                          -0.00885
                                      0.09005 -0.10
## base_logCRP
                          -0.73354
                                      0.08943 -8.20 2.4e-16
## chf
                          -0.63861
                                      0.08165 -7.82 5.2e-15
## htn
                          -0.06670
                                      0.10627 -0.63
                                                      0.530
                                      0.17003 -1.26
                          -0.21359
                                                      0.209
## canakinumab:diff_logCRP -0.04483
                                      0.06241 - 0.72
                                                      0.473
## diff_logCRP:age
                           0.00376
                                      0.00307 1.23
                                                      0.220
## diff_logCRP:dm
                                      0.06412 0.74
                           0.04716
                                                      0.462
## Log(scale)
                           0.52366
                                      0.02211 23.69 < 2e-16
##
## Scale= 1.69
##
## Weibull distribution
## Loglik(model) = -6072.8 Loglik(intercept only) = -6258.5
## Chisq= 371.37 on 11 degrees of freedom, p= 7.2e-73
## Number of Newton-Raphson Iterations: 7
## n= 8302
##
```

```
## ### Mediation analysis
##
              est
                                    Ζ
                          se
                                                           lower
                                                                     upper
## cde 0.1780520 0.08776683 2.028693 4.248956e-02 0.006032135 0.3500718
## pnde 0.1387042 0.09665386 1.435061 1.512698e-01 -0.050733912 0.3281422
## tnie 0.2068696 0.03998078 5.174225 2.288590e-07 0.128508667 0.2852305
## tnde 0.1671901 0.08655933 1.931508 5.342023e-02 -0.002463110 0.3368432
## pnie 0.1783837 0.04636945 3.847009 1.195687e-04 0.087501223 0.2692661
       0.3455737 0.09103109 3.796216 1.469213e-04 0.167156063 0.5239914
## pm
       0.6395734 0.17300163 3.696921 2.182302e-04 0.300496404 0.9786503
##
## Evaluated at:
## avar: canakinumab
## a1 (intervened value of avar) = 1
## a0 (reference value of avar) = 0
## mvar: diff_logCRP
## m_{cde} (intervend value of mvar for cde) = -2.137102
## cvar: age female base_logCRP chf htn dm
## c_cond (covariate vector value) = 54 0 1.0116 0 1 0
## Note that effect estimates can vary over m_cde and c_cond values when interaction = TRUE.
fit1.2 <- regmedint(data = anemia3,</pre>
                    yvar = "anemiayrs",
                    eventvar = "anemiaev",
                    avar = "canakinumab",
                    mvar = "diff_logCRP",
                    cvar = cvar,
                    ## EMM
                    emm_ac_mreg = c("base_logCRP"),
                    emm_ac_yreg = NULL,
                    emm_mc_yreg = c("age", "dm"),
                    ##
                    mreg = "linear",
                    yreg = "survAFT_weibull",
                    interaction = TRUE,
                    casecontrol = FALSE,
                    a0 = 0,
                    a1 = 1,
                    m_cde = mean(anemia3$diff_logCRP),
                    c_cond = c_cond_high_all)
summary(fit1.2)
## ### Mediator model
##
## lm(formula = diff_logCRP ~ canakinumab + age + female + base_logCRP +
       chf + htn + dm + canakinumab:base_logCRP, data = data)
##
## Residuals:
       Min
                1Q Median
                                3Q
## -4.4475 -0.5258 -0.0350 0.4525 5.6261
##
```

```
## Coefficients:
##
                            Estimate Std. Error t value Pr(>|t|)
                           0.4228295 0.0690623
                                                  6.122 9.63e-10 ***
## (Intercept)
## canakinumab
                          ## age
                          -0.0006421 0.0009539 -0.673
                                                          0.5008
                           0.0126025 0.0214422
## female
                                                  0.588
                                                          0.5567
## base logCRP
                          -1.4365841 0.0232393 -61.817 < 2e-16 ***
## chf
                           0.0928445 0.0191560
                                                  4.847 1.28e-06 ***
## htn
                           0.0163260 0.0230711
                                                  0.708
                                                          0.4792
## dm
                           0.0427971 0.0229645
                                                  1.864
                                                          0.0624 .
## canakinumab:base_logCRP -0.0314079 0.0284792 -1.103
                                                          0.2701
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
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## F-statistic: 1609 on 8 and 8293 DF, p-value: < 2.2e-16
##
## ### Outcome model
##
## Call:
## survival::survreg(formula = Surv(anemiayrs, anemiaev) ~ canakinumab +
      diff_logCRP + canakinumab:diff_logCRP + age + female + base_logCRP +
##
      chf + htn + dm + diff logCRP:age + diff logCRP:dm, data = data,
##
##
      dist = "weibull")
                             Value Std. Error
                                                  7.
## (Intercept)
                           7.40059
                                      0.53565 13.82 < 2e-16
## canakinumab
                           0.08224
                                      0.14929 0.55
                                                      0.582
                                      0.20525 -2.36
## diff_logCRP
                          -0.48373
                                                      0.018
                          -0.04799
                                      0.00802 -5.98 2.2e-09
## age
## female
                          -0.00885
                                      0.09005 - 0.10
                                                      0.922
## base_logCRP
                          -0.73354
                                      0.08943 -8.20 2.4e-16
## chf
                          -0.63861
                                      0.08165 -7.82 5.2e-15
## htn
                                      0.10627 -0.63
                          -0.06670
                                                      0.530
                          -0.21359
                                      0.17003 -1.26
                                                      0.209
## canakinumab:diff_logCRP -0.04483
                                      0.06241 -0.72
                                                      0.473
## diff logCRP:age
                           0.00376
                                      0.00307 1.23
                                                      0.220
## diff_logCRP:dm
                           0.04716
                                      0.06412 0.74
                                                      0.462
## Log(scale)
                           0.52366
                                      0.02211 23.69 < 2e-16
##
## Scale= 1.69
##
## Weibull distribution
## Loglik(model) = -6072.8 Loglik(intercept only) = -6258.5
## Chisq= 371.37 on 11 degrees of freedom, p= 7.2e-73
## Number of Newton-Raphson Iterations: 7
## n= 8302
##
## ### Mediation analysis
                         se
                                                        lower
                                                p
## cde 0.1780520 0.08776683 2.028693 0.0424895616 0.006032135 0.3500718
## pnde 0.1905550 0.09129110 2.087334 0.0368579399 0.011627762 0.3694823
## tnie 0.1521701 0.04534835 3.355582 0.0007919818 0.063288972 0.2410512
## tnde 0.2202737 0.11154304 1.974787 0.0482923802 0.001653350 0.4388940
```