SimpleViolins

Alex Badea

10/29/2021

Contents

Data sources 1
Plots for CPu 1

Data sources

We evaluate genotype and sex differences in mouse models of Alzheimer's disease risk, based on APOE alleles: APOE2 is protective, APOE3 is control, and APOE4 represents genetic risk for AD. One allele increases risk by 3 times, and two APOE4 alleles increase risk by 10 times (ref) in relation to APOE3. We use multiple data sources:

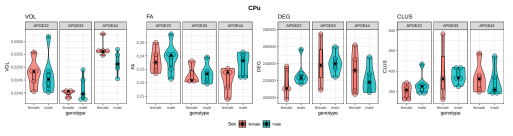
Morris Water Maze - testing spatial navigation memory Segmentation: regional volumes Microstructural properties: FA Tractography Based Connectivity: Degree, and Clustering Coefficient

We read the data and normalize volumes to the whole brain.

Plots for CPu

We embed plots for day d8 probe trials:

```
## Bin width defaults to 1/30 of the range of the data. Pick better value with `binwidth`. ## Bin width defaults to 1/30 of the range of the data. Pick better value with `binwidth`. ## Bin width defaults to 1/30 of the range of the data. Pick better value with `binwidth`. ## Bin width defaults to 1/30 of the range of the data. Pick better value with `binwidth`. ## Bin width defaults to 1/30 of the range of the data. Pick better value with `binwidth`.
```



```
print('CPu')
```

```
## [1] "CPu"
```

```
lm_CPu_VOL <- lm(CPu ~ Genotype*Sex, geno_combined_VOL)
summary(lm_CPu_VOL)</pre>
```

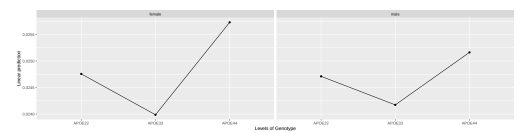
```
##
## lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
##
## Residuals:
##
         Min
                      1Q
                            Median
                                           30
                                                     Max
## -7.026e-04 -3.106e-04 -6.856e-05 1.250e-04 1.486e-03
## Coefficients:
##
                           Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          2.476e-02 2.420e-04 102.283 < 2e-16 ***
## GenotypeAPOE33
                         -7.692e-04 3.952e-04 -1.946 0.06395 .
## GenotypeAPOE44
                          9.720e-04 3.423e-04
                                                 2.840 0.00928 **
## Sexmale
                         -4.568e-05 3.277e-04
                                                -0.139 0.89036
## GenotypeAPOE33:Sexmale 2.317e-04 5.134e-04
                                                0.451 0.65601
## GenotypeAPOE44:Sexmale -5.205e-04 4.739e-04 -1.098 0.28336
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.0005412 on 23 degrees of freedom
## Multiple R-squared: 0.5711, Adjusted R-squared: 0.4778
## F-statistic: 6.124 on 5 and 23 DF, p-value: 0.0009559
lm_CPu_VOL_f <- lm(CPu ~ Genotype, combo_f_FA)</pre>
#summary(lm_CPu_VOL_f)
lm CPu VOL m <- lm(CPu ~ Genotype, combo m FA)</pre>
#summary(lm_CPu_VOL_f)
#omnibus testing
#https://cran.r-project.org/web/packages/emmeans/vignettes/confidence-intervals.html#adjust
sink(paste(outpath, "CPu_VOL_SimpleStats.txt",sep=''))
'omnibus testing'
## [1] "omnibus testing"
joint_tests(ref_grid(lm_CPu_VOL))
## model term
                df1 df2 F.ratio p.value
## Genotype
                   2
                     23 13.818 0.0001
                     23
                          0.476 0.4970
##
   Sex
                   1
                   2
                          1.148 0.3349
## Genotype:Sex
                     23
joint_tests(ref_grid(lm_CPu_VOL), by = "Sex", adjust = "sidak")
## Sex = female:
## model term df1 df2 F.ratio p.value
##
   Genotype
                2 23 10.239 0.0007
##
## Sex = male:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23 4.186 0.0282
```

```
joint_tests(ref_grid(lm_CPu_VOL), by = "Genotype",adjust = "sidak" )
## Genotype = APOE22:
## model term df1 df2 F.ratio p.value
                1 23
## Sex
                       0.019 0.8904
##
## Genotype = APOE33:
## model term df1 df2 F.ratio p.value
## Sex
                1 23
                       0.222 0.6423
##
## Genotype = APOE44:
## model term df1 df2 F.ratio p.value
## Sex
                1 23
                        2.736 0.1117
'anova'
## [1] "anova"
anova(lm CPu VOL)
## Analysis of Variance Table
##
## Response: CPu
##
                      Sum Sq
                               Mean Sq F value
## Genotype
                2 8.0970e-06 4.0485e-06 13.8218 0.0001142 ***
                1 1.9980e-07 1.9980e-07 0.6822 0.4173028
## Genotype:Sex 2 6.7230e-07 3.3610e-07 1.1476 0.3349205
## Residuals
             23 6.7368e-06 2.9290e-07
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#anova(lm_CPu_VOL_f)
#anova(lm_CPu_VOL_m)
#post hoc comparison
CPu_VOL.emm <- emmeans(lm_CPu_VOL , ~ Genotype | Sex, adjust = "sidak")
contrast(CPu_VOL.emm, simple = c("Genotype", "Sex")) #FDR
## contrast
                         estimate
                                       SE df t.ratio p.value
## APOE22 female effect 3.37e-06 0.000223 23
                                              0.015 0.9881
## APOE33 female effect -7.66e-04 0.000275 23 -2.784 0.0316
## APOE44 female effect 9.75e-04 0.000223 23
                                              4.378 0.0013
## APOE22 male effect -4.23e-05 0.000208 23 -0.204 0.9881
## APOE33 male effect -5.80e-04 0.000223 23 -2.603 0.0318
## APOE44 male effect 4.09e-04 0.000223 23
                                               1.837 0.1188
##
## P value adjustment: fdr method for 6 tests
contrast(CPu VOL.emm, simple = list("Genotype", "Sex"))
## $`simple contrasts for Genotype`
## Sex = female:
## contrast
                  estimate
                                 SE df t.ratio p.value
## APOE22 effect -6.76e-05 0.000208 23 -0.325 0.7484
## APOE33 effect -8.37e-04 0.000238 23 -3.523 0.0027
## APOE44 effect 9.04e-04 0.000208 23
                                       4.342 0.0007
##
```

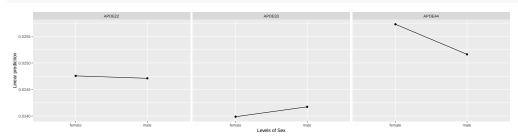
```
## Sex = male:
   contrast
                                 SE df t.ratio p.value
                  estimate
## APOE22 effect 2.87e-05 0.000186 23
                                        0.154 0.8791
## APOE33 effect -5.09e-04 0.000195 23 -2.611 0.0324
   APOE44 effect 4.80e-04 0.000195 23
                                        2.464 0.0324
##
## P value adjustment: fdr method for 3 tests
##
## $`simple contrasts for Sex`
## Genotype = APOE22:
## contrast
                                 SE df t.ratio p.value
                  estimate
## female effect 2.28e-05 0.000164 23
                                        0.139 0.8904
## male effect -2.28e-05 0.000164 23 -0.139 0.8904
##
## Genotype = APOE33:
## contrast
                  estimate
                                 SE df t.ratio p.value
## female effect -9.30e-05 0.000198 23 -0.471 0.6423
## male effect 9.30e-05 0.000198 23
                                        0.471 0.6423
##
## Genotype = APOE44:
## contrast
                  estimate
                                 SE df t.ratio p.value
## female effect 2.83e-04 0.000171 23
                                        1.654 0.1117
## male effect -2.83e-04 0.000171 23 -1.654 0.1117
## P value adjustment: fdr method for 2 tests
#post hoc tests
summary(glht(lm_CPu_VOL, emm(pairwise ~ Genotype*Sex, contr="sidak"))) #consider save to file
##
    Simultaneous Tests for General Linear Hypotheses
##
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
## Linear Hypotheses:
                                      Estimate Std. Error t value Pr(>|t|)
## APOE22 female - APOE33 female == 0 7.692e-04 3.952e-04
                                                           1.946 0.39957
## APOE22 female - APOE44 female == 0 -9.720e-04 3.423e-04 -2.840 0.08533 .
## APOE22 female - APOE22 male == 0
                                      4.568e-05 3.277e-04
                                                            0.139 0.99999
## APOE22 female - APOE33 male == 0
                                      5.832e-04 3.423e-04
                                                            1.704 0.54112
## APOE22 female - APOE44 male == 0 -4.058e-04 3.423e-04
                                                           -1.186 0.83794
## APOE33 female - APOE44 female == 0 -1.741e-03 3.952e-04
                                                           -4.406 0.00247 **
## APOE33 female - APOE22 male == 0 -7.235e-04 3.827e-04
                                                           -1.891 0.43058
## APOE33 female - APOE33 male == 0
                                   -1.860e-04 3.952e-04
                                                           -0.471 0.99671
                                    -1.175e-03 3.952e-04 -2.973 0.06505.
## APOE33 female - APOE44 male == 0
## APOE44 female - APOE22 male == 0
                                     1.018e-03
                                                3.277e-04
                                                            3.105 0.04942 *
## APOE44 female - APOE33 male == 0
                                     1.555e-03 3.423e-04
                                                            4.544 0.00180 **
## APOE44 female - APOE44 male == 0
                                      5.662e-04 3.423e-04
                                                            1.654 0.57153
## APOE22 male - APOE33 male == 0
                                      5.375e-04 3.277e-04
                                                            1.640 0.58018
## APOE22 male - APOE44 male == 0
                                                           -1.378 0.73784
                                     -4.515e-04 3.277e-04
## APOE33 male - APOE44 male == 0
                                     -9.890e-04 3.423e-04 -2.889 0.07745 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
```

```
res<-summary(glht(lm_CPu_VOL, emm(pairwise ~ Genotype*Sex))) #consider save to file
#summary(as.qlht(pairs(lm_CPu_VOL.emm), by = NULL))
summary(glht(lm_CPu_VOL, emm(pairwise ~ Genotype | Sex)))
## $`Sex = female`
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
## Linear Hypotheses:
                         Estimate Std. Error t value Pr(>|t|)
##
## APOE22 - APOE33 == 0 0.0007692 0.0003952
                                               1.946
                                                       0.1480
## APOE22 - APOE44 == 0 -0.0009720 0.0003423 -2.840
                                                       0.0241 *
## APOE33 - APOE44 == 0 -0.0017413 0.0003952 -4.406
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
##
##
## $`Sex = male`
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
## Linear Hypotheses:
                         Estimate Std. Error t value Pr(>|t|)
## APOE22 - APOE33 == 0 0.0005375 0.0003277
                                               1.640
                                                       0.2494
## APOE22 - APOE44 == 0 -0.0004515 0.0003277 -1.378
                                                        0.3686
## APOE33 - APOE44 == 0 -0.0009890 0.0003423 -2.889
                                                       0.0216 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
summary(glht(lm_CPu_VOL, emm(pairwise ~ Sex | Genotype)))
## $`Genotype = APOE22`
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
##
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
## female - male == 0 4.568e-05 3.277e-04
                                            0.139
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE33`
##
##
     Simultaneous Tests for General Linear Hypotheses
##
```

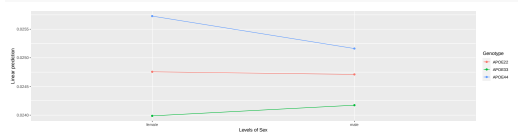
```
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
##
## Linear Hypotheses:
##
                       Estimate Std. Error t value Pr(>|t|)
## female - male == 0 - 0.0001860 0.0003952 - 0.471
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE44`
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_VOL)
##
## Linear Hypotheses:
##
                       Estimate Std. Error t value Pr(>|t|)
## female - male == 0 0.0005662 0.0003423 1.654
## (Adjusted p values reported -- single-step method)
mypairs<-pairs(CPu_VOL.emm) #_consider save to file</pre>
pairs(CPu_VOL.emm, by="Genotype")
## Genotype = APOE22:
## contrast
                  estimate
                                 SE df t.ratio p.value
## female - male 4.57e-05 0.000328 23 0.139 0.8904
##
## Genotype = APOE33:
## contrast
                  estimate
                                SE df t.ratio p.value
## female - male -1.86e-04 0.000395 23 -0.471 0.6423
## Genotype = APOE44:
## contrast
                                 SE df t.ratio p.value
                  estimate
                                         1.654 0.1117
## female - male 5.66e-04 0.000342 23
pairs(CPu VOL.emm, by="Sex")
## Sex = female:
## contrast
                    estimate
                                   SE df t.ratio p.value
## APOE22 - APOE33 0.000769 0.000395 23
                                          1.946 0.1486
## APOE22 - APOE44 -0.000972 0.000342 23 -2.840 0.0242
## APOE33 - APOE44 -0.001741 0.000395 23 -4.406 0.0006
##
## Sex = male:
## contrast
                    estimate
                                   SE df t.ratio p.value
## APOE22 - APOE33 0.000538 0.000328 23
                                          1.640 0.2496
## APOE22 - APOE44 -0.000451 0.000328 23 -1.378 0.3687
## APOE33 - APOE44 -0.000989 0.000342 23 -2.889 0.0217
## P value adjustment: tukey method for comparing a family of 3 estimates
print('CPu VOL')
## [1] "CPu VOL"
emmip(lm_CPu_VOL, ~ Genotype | Sex )
```



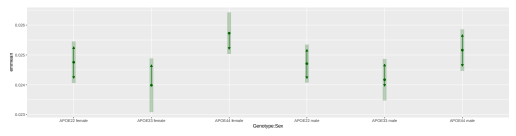
emmip(lm_CPu_VOL, ~ Sex | Genotype)



emmip(lm_CPu_VOL, Genotype~ Sex)



```
plot(CPu_VOL.emm, by = NULL, comparisons = TRUE, adjust = "mvt",
horizontal = FALSE, colors = "darkgreen", main = "CPu VOL")
```



ggsave(paste(outpath,'CPu_VOL_emmeans.pdf',sep=''), plot = last_plot(), device='pdf', scale=1, width=16

```
'simple stats'
```

[1] "simple stats"

print('CPu')

[1] "CPu"

lm_CPu_FA <- lm(CPu ~ Genotype*Sex, geno_combined_FA)
summary(lm_CPu_FA)</pre>

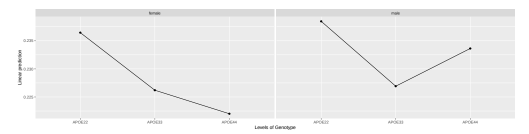
```
##
## Call:
## lm(formula = CPu ~ Genotype * Sex, data = geno combined FA)
## Residuals:
                            Median
                                           3Q
##
         Min
                     1Q
                                                     Max
## -0.0157390 -0.0071250 0.0007236 0.0057157 0.0178542
##
## Coefficients:
##
                          Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                          0.236449 0.004266 55.423
                                                       <2e-16 ***
                         -0.010244 0.006967 -1.470
## GenotypeAPOE33
                                                        0.1550
## GenotypeAPOE44
                         -0.014434 0.006033 -2.392
                                                        0.0253 *
## Sexmale
                          0.002017 0.005777
                                                0.349
                                                        0.7302
## GenotypeAPOE33:Sexmale -0.001301
                                     0.009050 -0.144
                                                        0.8869
## GenotypeAPOE44:Sexmale 0.009600 0.008353
                                               1.149
                                                        0.2623
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.00954 on 23 degrees of freedom
## Multiple R-squared: 0.3388, Adjusted R-squared: 0.195
## F-statistic: 2.357 on 5 and 23 DF, p-value: 0.07251
lm_CPu_FA_f <- lm(CPu ~ Genotype, combo_f_FA)</pre>
#summary(lm_CPu_FA_f)
lm_CPu_FA_m <- lm(CPu ~ Genotype, combo_m_FA)</pre>
#summary(lm_CPu_FA_f)
#omnibus testing
#https://cran.r-project.org/web/packages/emmeans/vignettes/confidence-intervals.html#adjust
sink(paste(outpath, "CPu_FA_SimpleStats.txt",sep=''))
'omnibus testing'
## [1] "omnibus testing"
joint_tests(ref_grid(lm_CPu_FA))
## model term df1 df2 F.ratio p.value
## Genotype
                  2 23
                          3.860 0.0359
## Sex
                  1
                     23
                          1.740 0.2001
                          0.924 0.4113
## Genotype:Sex
                  2 23
joint_tests(ref_grid(lm_CPu_FA), by = "Sex", adjust = "sidak")
## Sex = female:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23
                        2.978 0.0708
##
## Sex = male:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23
                        2.001 0.1581
joint_tests(ref_grid(lm_CPu_FA), by = "Genotype",adjust = "sidak" )
## Genotype = APOE22:
```

```
## model term df1 df2 F.ratio p.value
## Sex
                1 23 0.122 0.7302
##
## Genotype = APOE33:
## model term df1 df2 F.ratio p.value
## Sex
                1 23
                       0.011 0.9191
## Genotype = APOE44:
## model term df1 df2 F.ratio p.value
## Sex
                1 23 3.707 0.0666
'anova'
## [1] "anova"
anova(lm_CPu_FA)
## Analysis of Variance Table
## Response: CPu
##
               Df
                      Sum Sq
                                Mean Sq F value Pr(>F)
## Genotype
                2 0.00072293 0.00036146 3.9719 0.03298 *
                1 0.00018127 0.00018127 1.9919 0.17153
## Sex
## Genotype:Sex 2 0.00016811 0.00008405 0.9236 0.41131
## Residuals
             23 0.00209312 0.00009101
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#anova(lm_CPu_FA_f)
#anova(lm_CPu_FA_m)
#post hoc comparison
CPu_FA.emm <- emmeans(lm_CPu_FA , ~ Genotype | Sex, adjust = "sidak")
contrast(CPu_FA.emm, simple = c("Genotype", "Sex")) #FDR
## contrast
                        estimate
                                     SE df t.ratio p.value
## APOE22 female effect 0.00583 0.00393 23
                                            1.486 0.3018
## APOE33 female effect -0.00441 0.00485 23 -0.909 0.4471
## APOE44 female effect -0.00860 0.00393 23 -2.190 0.1282
## APOE22 male effect 0.00785 0.00366 23
                                            2.145 0.1282
## APOE33 male effect -0.00369 0.00393 23 -0.941 0.4471
## APOE44 male effect
                       0.00302 0.00393 23
                                            0.768 0.4502
##
## P value adjustment: fdr method for 6 tests
contrast(CPu_FA.emm, simple = list("Genotype", "Sex"))
## $`simple contrasts for Genotype`
## Sex = female:
## contrast
                  estimate
                                SE df t.ratio p.value
## APOE22 effect 0.008226 0.00367 23
                                      2.240 0.1051
## APOE33 effect -0.002018 0.00419 23 -0.482 0.6343
## APOE44 effect -0.006208 0.00367 23 -1.691 0.1566
##
## Sex = male:
## contrast
                  estimate
                                SE df t.ratio p.value
## APOE22 effect 0.005460 0.00328 23
                                      1.663 0.1650
```

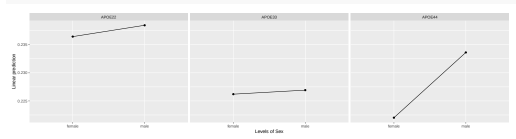
```
## APOE33 effect -0.006085 0.00343 23 -1.772 0.1650
## APOE44 effect 0.000625 0.00343 23
                                      0.182 0.8571
##
## P value adjustment: fdr method for 3 tests
##
## $`simple contrasts for Sex`
## Genotype = APOE22:
   contrast
                  estimate
                                SE df t.ratio p.value
   female effect -0.001008 0.00289 23
                                      -0.349 0.7302
                  0.001008 0.00289 23
##
   male effect
                                       0.349 0.7302
##
## Genotype = APOE33:
## contrast
                  estimate
                                SE df t.ratio p.value
## female effect -0.000358 0.00348 23 -0.103 0.9191
                  0.000358 0.00348 23
## male effect
                                      0.103 0.9191
##
## Genotype = APOE44:
## contrast
                  estimate
                                SE df t.ratio p.value
## female effect -0.005808 0.00302 23
                                      -1.925 0.0666
## male effect
                  0.005808 0.00302 23
                                        1.925
                                              0.0666
##
## P value adjustment: fdr method for 2 tests
#post hoc tests
summary(glht(lm_CPu_FA, emm(pairwise ~ Genotype*Sex, contr="sidak"))) #consider save to file
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
##
## Linear Hypotheses:
                                       Estimate Std. Error t value Pr(>|t|)
## APOE22 female - APOE33 female == 0 0.0102443 0.0069668
                                                             1.470
                                                                    0.6837
## APOE22 female - APOE44 female == 0 0.0144341 0.0060334
                                                             2.392
                                                                     0.1988
## APOE22 female - APOE22 male == 0 -0.0020165 0.0057766 -0.349
                                                                    0.9992
## APOE22 female - APOE33 male == 0
                                      0.0095289 0.0060334
                                                            1.579
                                                                     0.6176
## APOE22 female - APOE44 male == 0
                                                            0.467
                                      0.0028180 0.0060334
                                                                     0.9968
## APOE33 female - APOE44 female == 0 0.0041898 0.0069668
                                                            0.601
                                                                     0.9898
## APOE33 female - APOE22 male == 0 -0.0122608 0.0067456 -1.818
                                                                     0.4726
## APOE33 female - APOE33 male == 0 -0.0007154 0.0069668 -0.103
                                                                     1.0000
## APOE33 female - APOE44 male == 0
                                     -0.0074263 0.0069668
                                                           -1.066
                                                                     0.8889
## APOE44 female - APOE22 male == 0
                                    -0.0164506 0.0057766
                                                           -2.848
                                                                    0.0841
## APOE44 female - APOE33 male == 0
                                     -0.0049051 0.0060334
                                                           -0.813
                                                                     0.9618
## APOE44 female - APOE44 male == 0
                                                           -1.925
                                     -0.0116161 0.0060334
                                                                     0.4111
## APOE22 male - APOE33 male == 0
                                     0.0115455 0.0057766
                                                             1.999
                                                                     0.3712
## APOE22 male - APOE44 male == 0
                                     0.0048346 0.0057766
                                                            0.837
                                                                     0.9569
## APOE33 male - APOE44 male == 0
                                     -0.0067109 0.0060334 -1.112
                                                                    0.8704
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
res<-summary(glht(lm_CPu_FA, emm(pairwise ~ Genotype*Sex))) #consider save to file
\#summary(as.glht(pairs(lm_CPu_FA.emm), by = NULL))
```

```
summary(glht(lm_CPu_FA, emm(pairwise ~ Genotype | Sex)))
## $`Sex = female`
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
##
## Linear Hypotheses:
                        Estimate Std. Error t value Pr(>|t|)
## APOE22 - APOE33 == 0 0.010244 0.006967
                                              1.470
                                                      0.3222
## APOE22 - APOE44 == 0 0.014434 0.006033
                                              2.392
                                                      0.0628 .
## APOE33 - APOE44 == 0 0.004190 0.006967
                                              0.601
                                                      0.8202
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
##
##
## $`Sex = male`
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
## Linear Hypotheses:
                         Estimate Std. Error t value Pr(>|t|)
##
## APOE22 - APOE33 == 0 0.011545
                                    0.005777
                                              1.999
                                                        0.135
## APOE22 - APOE44 == 0 0.004835
                                    0.005777
                                               0.837
                                                        0.684
## APOE33 - APOE44 == 0 -0.006711
                                    0.006033 -1.112
                                                        0.516
## (Adjusted p values reported -- single-step method)
summary(glht(lm_CPu_FA, emm(pairwise ~ Sex | Genotype)))
## $`Genotype = APOE22`
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
## Linear Hypotheses:
                       Estimate Std. Error t value Pr(>|t|)
                                 0.005777 -0.349
## female - male == 0 - 0.002017
                                                       0.73
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE33`
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
##
## Linear Hypotheses:
                        Estimate Std. Error t value Pr(>|t|)
## female - male == 0 - 0.0007154 0.0069668 - 0.103
```

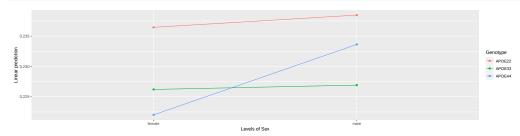
```
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE44`
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_FA)
##
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
## female - male == 0 -0.011616  0.006033 -1.925  0.0666 .
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
mypairs<-pairs(CPu_FA.emm) #_consider save to file</pre>
pairs(CPu_FA.emm, by="Genotype")
## Genotype = APOE22:
## contrast
                                SE df t.ratio p.value
                 estimate
## female - male -0.002017 0.00578 23 -0.349 0.7302
##
## Genotype = APOE33:
## contrast
                 estimate
                                SE df t.ratio p.value
## female - male -0.000715 0.00697 23 -0.103 0.9191
##
## Genotype = APOE44:
## contrast
                  estimate
                                SE df t.ratio p.value
## female - male -0.011616 0.00603 23 -1.925 0.0666
pairs(CPu_FA.emm, by="Sex")
## Sex = female:
## contrast
             estimate
                                 SE df t.ratio p.value
## APOE22 - APOE33 0.01024 0.00697 23 1.470 0.3232
## APOE22 - APOE44 0.01443 0.00603 23
                                        2.392 0.0630
## APOE33 - APOE44 0.00419 0.00697 23 0.601 0.8208
##
## Sex = male:
## contrast
                                 SE df t.ratio p.value
                   estimate
## APOE22 - APOE33 0.01155 0.00578 23 1.999 0.1351
## APOE22 - APOE44 0.00483 0.00578 23 0.837 0.6844
## APOE33 - APOE44 -0.00671 0.00603 23 -1.112 0.5164
## P value adjustment: tukey method for comparing a family of 3 estimates
print('CPu FA')
## [1] "CPu FA"
emmip(lm_CPu_FA, ~ Genotype | Sex )
```



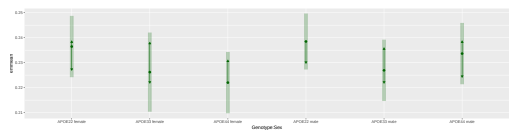
emmip(lm_CPu_FA, ~ Sex | Genotype)



emmip(lm_CPu_FA, Genotype~ Sex)



```
sink()
plot(CPu_FA.emm, by = NULL, comparisons = TRUE, adjust = "mvt",
horizontal = FALSE, colors = "darkgreen", main = "CPu FA")
```



ggsave(paste(outpath, 'CPu_FA_emmeans.pdf',sep=''), plot = last_plot(), device='pdf', scale=1, width=16,

```
'simple stats'
```

[1] "simple stats"

print('CPu')

[1] "CP11"

lm_CPu_DEG <- lm(CPu ~ Genotype*Sex, geno_combined_DEG)
summary(lm_CPu_DEG)</pre>

##

Call:

```
## lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
##
## Residuals:
##
     Min
              1Q Median
                            3Q
                                  Max
## -31667 -11917 -3800 14800
                                34333
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                            218000
                                        8706 25.040
                                                       <2e-16 ***
## GenotypeAPOE33
                             22667
                                        14217
                                                1.594
                                                         0.125
## GenotypeAPOE44
                             12200
                                       12312
                                              0.991
                                                         0.332
## Sexmale
                              9917
                                        11788
                                              0.841
                                                         0.409
## GenotypeAPOE33:Sexmale
                             -9783
                                       18468 -0.530
                                                         0.601
                                       17045 -1.075
                            -18317
                                                         0.294
## GenotypeAPOE44:Sexmale
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 19470 on 23 degrees of freedom
## Multiple R-squared: 0.1853, Adjusted R-squared: 0.008217
## F-statistic: 1.046 on 5 and 23 DF, p-value: 0.4148
lm_CPu_DEG_f <- lm(CPu ~ Genotype, combo_f_FA)</pre>
#summary(lm_CPu_DEG_f)
lm_CPu_DEG_m <- lm(CPu ~ Genotype, combo_m_FA)</pre>
\#summary(lm\_CPu\_DEG\_f)
#omnibus testing
#https://cran.r-project.org/web/packages/emmeans/vignettes/confidence-intervals.html#adjust
sink(paste(outpath, "CPu_DEG_SimpleStats.txt",sep=''))
'omnibus testing'
## [1] "omnibus testing"
joint_tests(ref_grid(lm_CPu_DEG))
## model term
                 df1 df2 F.ratio p.value
## Genotype
                   2
                     23
                           2.004 0.1577
                     23
                           0.006 0.9414
## Sex
                   1
## Genotype:Sex
                   2 23
                           0.579 0.5683
joint_tests(ref_grid(lm_CPu_DEG), by = "Sex", adjust = "sidak")
## Sex = female:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23
                       1.327 0.2849
##
## Sex = male:
## model term df1 df2 F.ratio p.value
                 2 23
                        1.247 0.3060
## Genotype
joint_tests(ref_grid(lm_CPu_DEG), by = "Genotype",adjust = "sidak" )
## Genotype = APOE22:
## model term df1 df2 F.ratio p.value
                 1 23 0.708 0.4089
## Sex
```

```
##
## Genotype = APOE33:
## model term df1 df2 F.ratio p.value
                1 23 0.000 0.9926
## Sex
## Genotype = APOE44:
## model term df1 df2 F.ratio p.value
                        0.465 0.5019
## Sex
                1 23
'anova'
## [1] "anova"
anova(lm_CPu_DEG)
## Analysis of Variance Table
##
## Response: CPu
##
               Df
                      Sum Sq Mean Sq F value Pr(>F)
                2 1538125392 769062696 2.0294 0.1543
## Genotype
                     5580291
                              5580291 0.0147 0.9045
## Sex
                1
## Genotype:Sex 2 439053800 219526900 0.5793 0.5683
## Residuals
               23 8716275000 378968478
#anova(lm CPu DEG f)
#anova(lm_CPu_DEG_m)
#post hoc comparison
CPu_DEG.emm <- emmeans(lm_CPu_DEG , ~ Genotype | Sex, adjust = "sidak")</pre>
contrast(CPu_DEG.emm, simple = c("Genotype", "Sex")) #FDR
  contrast
                        estimate
                                  SE df t.ratio p.value
## APOE22 female effect -11897 8013 23 -1.485 0.4842
                        10769 9894 23
## APOE33 female effect
                                          1.088 0.4842
## APOE44 female effect
                           303 8013 23
                                         0.038 0.9702
                        -1981 7469 23 -0.265 0.9519
## APOE22 male effect
                        10903 8013 23
## APOE33 male effect
                                         1.361 0.4842
## APOE44 male effect
                          -8097 8013 23 -1.010 0.4842
##
## P value adjustment: fdr method for 6 tests
contrast(CPu DEG.emm, simple = list("Genotype", "Sex"))
## $`simple contrasts for Genotype`
## Sex = female:
## contrast
                            SE df t.ratio p.value
                 estimate
## APOE22 effect -11622 7493 23 -1.551 0.3134
## APOE33 effect 11044 8543 23
                                   1.293 0.3134
## APOE44 effect
                    578 7493 23
                                  0.077 0.9392
##
## Sex = male:
## contrast
                 estimate
                            SE df t.ratio p.value
## APOE22 effect
                   -2256 6702 23 -0.337 0.7395
## APOE33 effect
                   10628 7009 23
                                  1.516 0.3667
## APOE44 effect
                    -8372 7009 23 -1.195 0.3667
##
## P value adjustment: fdr method for 3 tests
```

```
##
## $`simple contrasts for Sex`
## Genotype = APOE22:
## contrast
                 estimate
                            SE df t.ratio p.value
## female effect -4958.3 5894 23 -0.841 0.4089
## male effect
                   4958.3 5894 23
                                    0.841 0.4089
## Genotype = APOE33:
   contrast
                 estimate
                            SE df t.ratio p.value
## female effect
                    -66.7 7108 23 -0.009 0.9926
## male effect
                     66.7 7108 23
                                    0.009 0.9926
##
## Genotype = APOE44:
                  estimate
                            SE df t.ratio p.value
## contrast
## female effect
                  4200.0 6156 23
                                    0.682 0.5019
## male effect
                  -4200.0 6156 23 -0.682 0.5019
##
## P value adjustment: fdr method for 2 tests
#post hoc tests
summary(glht(lm_CPu_DEG, emm(pairwise ~ Genotype*Sex, contr="sidak"))) #consider save to file
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
## Linear Hypotheses:
                                     Estimate Std. Error t value Pr(>|t|)
                                                 14216.8 -1.594
## APOE22 female - APOE33 female == 0 -22666.7
                                                                    0.608
## APOE22 female - APOE44 female == 0 -12200.0
                                                 12312.1 -0.991
                                                                    0.916
## APOE22 female - APOE22 male == 0
                                      -9916.7
                                                 11787.9 -0.841
                                                                    0.956
## APOE22 female - APOE33 male == 0
                                    -22800.0
                                                 12312.1 -1.852
                                                                    0.453
## APOE22 female - APOE44 male == 0
                                      -3800.0
                                                 12312.1 -0.309
                                                                    1.000
## APOE33 female - APOE44 female == 0 10466.7
                                                 14216.8 0.736
                                                                    0.975
## APOE33 female - APOE22 male == 0
                                      12750.0
                                                 13765.3 0.926
                                                                    0.935
## APOE33 female - APOE33 male == 0
                                       -133.3
                                                 14216.8 -0.009
                                                                    1.000
## APOE33 female - APOE44 male == 0
                                                          1.327
                                      18866.7
                                                 14216.8
                                                                    0.766
## APOE44 female - APOE22 male == 0
                                       2283.3
                                                 11787.9 0.194
                                                                    1.000
## APOE44 female - APOE33 male == 0
                                     -10600.0
                                                 12312.1 -0.861
                                                                    0.952
## APOE44 female - APOE44 male == 0
                                       8400.0
                                                 12312.1 0.682
                                                                    0.982
## APOE22 male - APOE33 male == 0
                                     -12883.3
                                                 11787.9 -1.093
                                                                    0.878
## APOE22 male - APOE44 male == 0
                                       6116.7
                                                 11787.9 0.519
                                                                    0.995
## APOE33 male - APOE44 male == 0
                                      19000.0
                                                 12312.1
                                                           1.543
                                                                    0.640
## (Adjusted p values reported -- single-step method)
res<-summary(glht(lm_CPu_DEG, emm(pairwise ~ Genotype*Sex))) #consider save to file
#summary(as.glht(pairs(lm_CPu_DEG.emm), by = NULL))
summary(glht(lm_CPu_DEG, emm(pairwise ~ Genotype | Sex)))
## $`Sex = female`
##
##
     Simultaneous Tests for General Linear Hypotheses
```

##

```
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
##
## Linear Hypotheses:
                        Estimate Std. Error t value Pr(>|t|)
##
## APOE22 - APOE33 == 0
                          -22667
                                      14217 -1.594
                        -12200
## APOE22 - APOE44 == 0
                                      12312 -0.991
                                                       0.589
## APOE33 - APOE44 == 0
                        10467
                                      14217 0.736
## (Adjusted p values reported -- single-step method)
##
##
## $`Sex = male`
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
## Linear Hypotheses:
##
                        Estimate Std. Error t value Pr(>|t|)
## APOE22 - APOE33 == 0
                                      11788 -1.093
                          -12883
                                                       0.528
## APOE22 - APOE44 == 0
                            6117
                                      11788
                                             0.519
                                                       0.863
## APOE33 - APOE44 == 0
                           19000
                                      12312
                                              1.543
                                                       0.290
## (Adjusted p values reported -- single-step method)
summary(glht(lm_CPu_DEG, emm(pairwise ~ Sex | Genotype)))
## $`Genotype = APOE22`
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
##
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
##
## female - male == 0
                         -9917
                                    11788 -0.841
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE33`
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
##
## Linear Hypotheses:
##
                      Estimate Std. Error t value Pr(>|t|)
## female - male == 0 -133.3
                                  14216.8 -0.009
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE44`
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_DEG)
##
```

```
## Linear Hypotheses:
##
                    Estimate Std. Error t value Pr(>|t|)
## female - male == 0
                       8400
                                  12312
                                        0.682
## (Adjusted p values reported -- single-step method)
mypairs<-pairs(CPu_DEG.emm) #_consider save to file</pre>
pairs(CPu_DEG.emm, by="Genotype")
## Genotype = APOE22:
## contrast estimate SE df t.ratio p.value
## female - male -9917 11788 23 -0.841 0.4089
##
## Genotype = APOE33:
## contrast
                          SE df t.ratio p.value
                estimate
## female - male -133 14217 23 -0.009 0.9926
##
## Genotype = APOE44:
## contrast
                estimate
                            SE df t.ratio p.value
                    8400 12312 23 0.682 0.5019
## female - male
pairs(CPu_DEG.emm, by="Sex")
## Sex = female:
## contrast
                  estimate
                              SE df t.ratio p.value
## APOE22 - APOE33 -22667 14217 23 -1.594 0.2682
## APOE22 - APOE44 -12200 12312 23 -0.991 0.5899
## APOE33 - APOE44 10467 14217 23 0.736 0.7448
##
## Sex = male:
## contrast
               estimate
                              SE df t.ratio p.value
## APOE22 - APOE33 -12883 11788 23 -1.093 0.5279
## APOE22 - APOE44
                     6117 11788 23 0.519 0.8629
## APOE33 - APOE44
                   19000 12312 23
                                    1.543 0.2901
##
## P value adjustment: tukey method for comparing a family of 3 estimates
print('CPu DEG')
## [1] "CPu DEG"
emmip(lm_CPu_DEG, ~ Genotype | Sex )
```

emmip(lm_CPu_DEG, ~ Sex | Genotype)

```
emmip(lm_CPu_DEG, Genotype~ Sex )
                                  Levels of Sex
sink()
plot(CPu_DEG.emm, by = NULL, comparisons = TRUE, adjust = "mvt",
horizontal = FALSE, colors = "darkgreen", main = "CPu DEG")
ggsave(paste(outpath,'CPu_DEG_emmeans.pdf',sep=''), plot = last_plot(), device='pdf', scale=1, width=16
'simple stats'
## [1] "simple stats"
print('CPu')
## [1] "CPu"
lm_CPu_CLUS <- lm(CPu ~ Genotype*Sex, geno_combined_CLUS)</pre>
summary(lm_CPu_CLUS)
##
## Call:
## lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
##
## Residuals:
##
                1Q Median
                                 ЗQ
                                        Max
## -70.017 -20.873 -5.233 11.696 89.933
##
## Coefficients:
##
                           Estimate Std. Error t value Pr(>|t|)
                             350.33
                                         16.50 21.229
## (Intercept)
                                                          <2e-16 ***
```

1.892

0.0711 .

26.95

50.99

GenotypeAPOE33

```
23.34 1.449
                                                      0.1607
## GenotypeAPOE44
                            33.83
## Sexmale
                           17.17
                                      22.34 0.768
                                                    0.4501
## GenotypeAPOE33:Sexmale -29.15
                                      35.01 -0.833 0.4135
## GenotypeAPOE44:Sexmale -27.20
                                      32.31 -0.842 0.4086
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 36.9 on 23 degrees of freedom
## Multiple R-squared: 0.1803, Adjusted R-squared: 0.002162
## F-statistic: 1.012 on 5 and 23 DF, p-value: 0.4332
lm_CPu_CLUS_f <- lm(CPu ~ Genotype, combo_f_FA)</pre>
#summary(lm_CPu_CLUS_f)
lm_CPu_CLUS_m <- lm(CPu ~ Genotype, combo_m_FA)</pre>
#summary(lm_CPu_CLUS_f)
#omnibus testing
#https://cran.r-project.org/web/packages/emmeans/vignettes/confidence-intervals.html#adjust
sink(paste(outpath, "CPu_CLUS_SimpleStats.txt",sep=''))
'omnibus testing'
## [1] "omnibus testing"
joint_tests(ref_grid(lm_CPu_CLUS))
## model term
                df1 df2 F.ratio p.value
## Genotype
                  2 23
                          2.233 0.1299
                  1 23
## Sex
                         0.013 0.9094
## Genotype:Sex
                 2 23
                          0.486 0.6210
joint_tests(ref_grid(lm_CPu_CLUS), by = "Sex", adjust = "sidak")
## Sex = female:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23 2.035 0.1536
##
## Sex = male:
## model term df1 df2 F.ratio p.value
## Genotype
                2 23
                       0.491 0.6181
joint_tests(ref_grid(lm_CPu_CLUS), by = "Genotype",adjust = "sidak")
## Genotype = APOE22:
## model term df1 df2 F.ratio p.value
                1 23 0.590 0.4501
## Sex
##
## Genotype = APOE33:
## model term df1 df2 F.ratio p.value
                1 23
                       0.198 0.6607
##
## Genotype = APOE44:
## model term df1 df2 F.ratio p.value
          1 23 0.185 0.6714
```

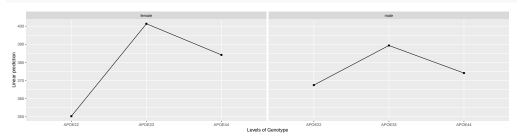
```
'anova'
## [1] "anova"
anova(lm_CPu_CLUS)
## Analysis of Variance Table
##
## Response: CPu
                   Sum Sq Mean Sq F value Pr(>F)
               Df
                2 5566.0 2783.01 2.0439 0.1524
## Genotype
## Sex
                1
                      0.1
                             0.07 0.0001 0.9943
## Genotype:Sex 2 1324.6 662.30 0.4864 0.6210
## Residuals
               23 31317.3 1361.62
#anova(lm_CPu_CLUS_f)
#anova(lm_CPu_CLUS_m)
#post hoc comparison
CPu_CLUS.emm <- emmeans(lm_CPu_CLUS , ~ Genotype | Sex, adjust = "sidak")</pre>
contrast(CPu_CLUS.emm, simple = c("Genotype", "Sex")) #FDR
## contrast
                        estimate
                                  SE df t.ratio p.value
## APOE22 female effect
                        -27.47 15.2 23 -1.808 0.5020
## APOE33 female effect
                          23.53 18.8 23
                                          1.254 0.6668
## APOE44 female effect
                           6.36 15.2 23
                                         0.419 0.8113
## APOE22 male effect -10.30 14.2 23 -0.727 0.7116
                        11.54 15.2 23
## APOE33 male effect
                                         0.760 0.7116
## APOE44 male effect
                          -3.67 15.2 23 -0.241 0.8113
##
## P value adjustment: fdr method for 6 tests
contrast(CPu_CLUS.emm, simple = list("Genotype", "Sex"))
## $`simple contrasts for Genotype`
## Sex = female:
## contrast
                 estimate
                            SE df t.ratio p.value
## APOE22 effect -28.27 14.2 23 -1.991 0.1756
                  22.72 16.2 23
## APOE33 effect
                                   1.403 0.2610
## APOE44 effect
                    5.55 14.2 23
                                   0.391 0.6994
##
## Sex = male:
## contrast
                estimate
                            SE df t.ratio p.value
## APOE22 effect -9.49 12.7 23 -0.747 0.6940
  APOE33 effect 12.35 13.3 23
                                  0.930 0.6940
  APOE44 effect
                   -2.86 13.3 23 -0.215 0.8314
##
## P value adjustment: fdr method for 3 tests
## $`simple contrasts for Sex`
## Genotype = APOE22:
## contrast
                 estimate
                            SE df t.ratio p.value
## female effect -8.58 11.2 23 -0.768 0.4501
## male effect
                     8.58 11.2 23
                                  0.768 0.4501
##
## Genotype = APOE33:
```

```
estimate SE df t.ratio p.value
## contrast
                    5.99 13.5 23
## female effect
                                     0.445 0.6607
   male effect
                     -5.99 13.5 23 -0.445 0.6607
##
## Genotype = APOE44:
##
   contrast
                  estimate
                             SE df t.ratio p.value
   female effect
                      5.01 11.7 23
                                     0.430 0.6714
## male effect
                     -5.01 11.7 23 -0.430 0.6714
## P value adjustment: fdr method for 2 tests
#post hoc tests
summary(glht(lm_CPu_CLUS, emm(pairwise ~ Genotype*Sex, contr="sidak"))) #consider save to file
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
## Linear Hypotheses:
                                      Estimate Std. Error t value Pr(>|t|)
## APOE22 female - APOE33 female == 0
                                      -50.993
                                                   26.948 -1.892
                                                                     0.430
## APOE22 female - APOE44 female == 0 -33.826
                                                   23.338 -1.449
                                                                     0.696
                                                   22.344 -0.768
## APOE22 female - APOE22 male == 0
                                       -17.169
                                                                     0.970
## APOE22 female - APOE33 male == 0
                                       -39.008
                                                   23.338 -1.671
                                                                     0.561
## APOE22 female - APOE44 male == 0
                                       -23.798
                                                   23.338 -1.020
                                                                     0.906
## APOE33 female - APOE44 female == 0
                                        17.167
                                                   26.948
                                                           0.637
                                                                     0.987
## APOE33 female - APOE22 male == 0
                                        33.823
                                                   26.092
                                                           1.296
                                                                     0.783
## APOE33 female - APOE33 male == 0
                                       11.985
                                                   26.948 0.445
                                                                     0.997
## APOE33 female - APOE44 male == 0
                                        27.195
                                                           1.009
                                                                     0.909
                                                   26.948
## APOE44 female - APOE22 male == 0
                                        16.657
                                                           0.745
                                                                     0.974
                                                   22.344
## APOE44 female - APOE33 male == 0
                                       -5.182
                                                   23.338 -0.222
                                                                     1.000
## APOE44 female - APOE44 male == 0
                                       10.028
                                                   23.338
                                                           0.430
                                                                     0.998
## APOE22 male - APOE33 male == 0
                                       -21.839
                                                   22.344 -0.977
                                                                     0.920
## APOE22 male - APOE44 male == 0
                                        -6.629
                                                   22.344 -0.297
                                                                     1.000
## APOE33 male - APOE44 male == 0
                                        15.210
                                                   23.338
                                                          0.652
                                                                     0.985
## (Adjusted p values reported -- single-step method)
res<-summary(glht(lm_CPu_CLUS, emm(pairwise ~ Genotype*Sex))) #consider save to file
\#summary(as.glht(pairs(lm_CPu_CLUS.emm), by = NULL))
summary(glht(lm CPu CLUS, emm(pairwise ~ Genotype | Sex)))
## $`Sex = female`
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
## Linear Hypotheses:
                        Estimate Std. Error t value Pr(>|t|)
## APOE22 - APOE33 == 0
                          -50.99
                                      26.95
                                            -1.892
                                                       0.163
## APOE22 - APOE44 == 0
                          -33.83
                                      23.34 -1.449
                                                       0.332
## APOE33 - APOE44 == 0
                          17.17
                                      26.95
                                                       0.801
## (Adjusted p values reported -- single-step method)
```

```
##
##
## $`Sex = male`
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
##
## Linear Hypotheses:
##
                        Estimate Std. Error t value Pr(>|t|)
## APOE22 - APOE33 == 0 -21.839
                                     22.344 -0.977
## APOE22 - APOE44 == 0
                         -6.629
                                     22.344 -0.297
                                                        0.953
## APOE33 - APOE44 == 0
                          15.210
                                     23.338
                                             0.652
                                                        0.793
## (Adjusted p values reported -- single-step method)
summary(glht(lm_CPu_CLUS, emm(pairwise ~ Sex | Genotype)))
## $`Genotype = APOE22`
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
##
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
## female - male == 0 -17.17
                                    22.34 -0.768
## (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE33`
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
##
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
##
## female - male == 0
                         11.98
                                    26.95
                                            0.445
                                                      0.661
  (Adjusted p values reported -- single-step method)
##
##
## $`Genotype = APOE44`
##
##
    Simultaneous Tests for General Linear Hypotheses
## Fit: lm(formula = CPu ~ Genotype * Sex, data = geno_combined_CLUS)
## Linear Hypotheses:
                      Estimate Std. Error t value Pr(>|t|)
## female - male == 0
                         10.03
                                    23.34
                                             0.43
                                                      0.671
## (Adjusted p values reported -- single-step method)
mypairs<-pairs(CPu_CLUS.emm) #_consider save to file</pre>
pairs(CPu_CLUS.emm, by="Genotype")
```

```
## Genotype = APOE22:
## contrast
             estimate SE df t.ratio p.value
## female - male -17.2 22.3 23 -0.768 0.4501
##
## Genotype = APOE33:
## contrast
                estimate SE df t.ratio p.value
## female - male 12.0 26.9 23 0.445 0.6607
##
## Genotype = APOE44:
## contrast
                estimate
                           SE df t.ratio p.value
## female - male
                    10.0 23.3 23
                                 0.430 0.6714
pairs(CPu_CLUS.emm, by="Sex")
## Sex = female:
## contrast
                            SE df t.ratio p.value
                  estimate
## APOE22 - APOE33 -50.99 26.9 23 -1.892 0.1636
## APOE22 - APOE44 -33.83 23.3 23 -1.449 0.3332
## APOE33 - APOE44 17.17 26.9 23 0.637 0.8014
##
## Sex = male:
## contrast
               estimate SE df t.ratio p.value
## APOE22 - APOE33 -21.84 22.3 23 -0.977 0.5981
## APOE22 - APOE44 -6.63 22.3 23 -0.297 0.9527
## APOE33 - APOE44 15.21 23.3 23 0.652 0.7933
##
## P value adjustment: tukey method for comparing a family of 3 estimates
print('CPu CLUS')
## [1] "CPu CLUS"
```

emmip(lm_CPu_CLUS, ~ Genotype | Sex)



emmip(lm_CPu_CLUS, ~ Sex | Genotype)

