This is a note for Golden Gate Association analysis

R. Markdown

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This project is led by Ms. Jacqui Marzec. Our objectives are to find any significant association between SNPs (Golden Gate SNP candidate pool) in RSV positivie patients.

GWAS on mitochondrial variants

In this analysis, we tried three different cases as following. Each setting will help to answer different hypothesis. The Analysis will be done with each case independently.

Case1, all three genotypes are kept: AA/AB/BB Case2, two genotypes are kept: AA AB/BB Case3, tow genotypes are kept: AA/AB BB

Analysis set up and load necessary functions

```
library(HardyWeinberg)
library(XLConnect)
source("helperScripts/analysisFuncs.R")
```

Load up preprocessed data

```
## Phenotype data
dt <- read.table("data/RSV_cohort_data_ASCII.txt",header = TRUE, sep= "\t")

## ONLY use for RSV positive candidates
row2use <- which (dt[,5] == 1)

## This is for reading Excel file
## We need to set memeory at 24 GB
options(java.parameters = "-Xmx24g")

snpFiles <- "data/ASCII_genotype_data_GoldenGate_for_JYL_07122015.xlsx"

## There are five sheets of data
sheets <- c (seq (1,5))</pre>
```

Analysis results on disease severity – logistic regression model

```
sig.asso.SAT <- c (list())
results.logit <- logisticAnalysis (sig.asso.SAT, snpFiles, sheets, dt, 6, row2use)
results.logit</pre>
```

```
## [[1]]
## [[1]]$`Case-I`
                SNP
## 1 seq.rs3806496 0.03249843
## 2 seq.rs2066843 0.03161207
## 3 seq.rs17244587 0.04618051
## [[1]]$`Case-II`
                          pval
##
                SNP
## 1 seq.rs12987402 0.03525483
      seq.rs469390 0.02409298
## 3 seq.rs3806265 0.03776518
## 4 seq.rs2066843 0.03371025
## [[1]]$`Case-III`
##
                SNP
                          pval
## 1 seq.rs17189298 0.04594427
## 2 seq.rs2770146 0.01637843
##
##
## [[2]]
## [[2]]$`Case-I`
##
              SNP
                           pval
## 1 seq.rs2252930 0.0123745940
## 2 seq.rs2243592 0.0005072162
## 3 seq.rs2257167 0.0016167643
## 4 seq.rs2243599 0.0006175882
## 5 seq.rs2254180 0.0054136510
## 6 seq.rs2254315 0.0042648958
## 7 seq.rs1474347 0.0233376514
## 8 seq.rs8193036 0.0138688712
##
## [[2]]$`Case-II`
                SNP
                           pval
## 1 seq.rs2243592 0.003018021
## 2 seq.rs2257167 0.006062165
## 3 seq.rs2243599 0.006208847
## 4 seq.rs2254180 0.021865717
     seq.rs2254315 0.015360578
## 6 seq.rs1800795 0.025821513
## 7 seq.rs2069832 0.012473056
## 8 seq.rs1474347 0.011935878
## 9 seq.rs1800893 0.048201983
## 10 seq.rs8193036 0.019022066
## [[2]]$`Case-III`
                SNP
                           pval
## 1 seq.rs2280789 0.026207189
## 2 seq.rs2252930 0.010498389
     seq.rs2243592 0.006836451
## 4 seq.rs2257167 0.011030534
## 5 seq.rs2243599 0.004429888
## 6 seq.rs2254180 0.019988072
## 7 seq.rs2254315 0.022174971
```

```
## 8 seq.rs9376268 0.042321508
## 9 seq.rs9376267 0.029287461
## 10 seq.rs369908 0.023580640
##
##
## [[3]]
## [[3]]$`Case-I`
               SNP
## 1 seq.rs1293749 0.041542545
## 2 seq.rs15895 0.034259581
## 3 seq.rs7966314 0.030823898
## 4 seq.rs2072133 0.007942976
## 5 seq.rs1041981 0.024407218
## [[3]]$`Case-II`
##
               SNP
                         pval
## 1 seq.rs7966314 0.02788702
## 2 seq.rs2072133 0.01039301
## [[3]]$`Case-III`
##
                SNP
                          pval
## 1 seq.rs1293755 0.04087340
## 2
      seq.rs15895 0.04041382
## 3 seq.rs12693591 0.01505629
## 4 seq.rs3771300 0.03111144
## 5 seq.rs1800693 0.01013580
##
## [[4]]
## [[4]]$`Case-I`
##
                        pval
## 1 seq.rs3769823 0.0402697
## [[4]]$`Case-II`
## NULL
##
## [[4]]$`Case-III`
##
               SNP
## 1 seq.rs3769823 0.01005097
## 2 seq.rs4844590 0.03257378
##
##
## [[5]]
## [[5]]$`Case-I`
           SNP
                     pval
## 1 rs3761624 0.02619375
##
## [[5]]$`Case-II`
           SNP
                     pval
## 1 rs3761624 0.02858216
## [[5]]$`Case-III`
## NULL
```

Analysis results on disease severity - chi-square test based model

```
<- c (list())
sig.asso.SAT
results.chisqure <- AssoAnalysis
                                      (sig.asso.SAT, snpFiles, sheets, dt, 6, row2use)
results.chisqure
## [[1]]
## [[1]]$`Case-I`
               SNP
                         pval
## 1 seq.rs469390 0.03553412
## 2 seq.rs2770146 0.03031926
## [[1]]$`Case-II`
##
                SNP
                          pval
## 1 seq.rs12987402 0.04484957
      seq.rs469390 0.03136834
## 3 seq.rs3806265 0.04864552
## 4 seq.rs2066843 0.04292077
##
## [[1]]$`Case-III`
##
               SNP
                         pval
## 1 seq.rs2770146 0.02269727
##
##
## [[2]]
## [[2]]$`Case-I`
                           pval
                SNP
## 1
     seq.rs2252930 0.032963115
## 2
     seq.rs2243592 0.002065785
     seq.rs2257167 0.004097110
## 4 seq.rs2243599 0.002391277
     seq.rs2254180 0.016497081
## 6
     seq.rs2254315 0.014203726
     seq.rs2069832 0.026046287
## 8 seq.rs1474347 0.040922019
     seq.rs8193036 0.046061333
## 10 seq.rs8193038 0.049507445
##
## [[2]]$`Case-II`
               SNP
                          pval
## 1 seq.rs2243592 0.003973553
## 2 seq.rs2257167 0.007940527
## 3 seq.rs2243599 0.008163886
## 4 seq.rs2254180 0.028029801
## 5 seq.rs2254315 0.019873260
## 6 seq.rs1800795 0.033217808
## 7 seq.rs2069832 0.016223152
## 8 seq.rs1474347 0.015568224
## 9 seq.rs8193036 0.024515805
## [[2]]$`Case-III`
##
                SNP
                           pval
## 1 seq.rs2280789 0.037358791
## 2 seq.rs2252930 0.013686847
```

```
## 3 seq.rs2243592 0.009214950
## 4 seq.rs2257167 0.014382366
## 5 seq.rs2243599 0.005967165
## 6 seq.rs2254180 0.027873558
## 7 seq.rs2254315 0.030957836
## 8 seq.rs9376268 0.033497757
## 9 seq.rs9376267 0.035090018
## 10 seq.rs369908 0.033457931
##
##
## [[3]]
## [[3]]$`Case-I`
                SNP
                          pval
## 1 seq.rs2072133 0.02734773
## 2 seq.rs12693591 0.01358428
## 3 seq.rs361525 0.02179245
## 4 seq.rs1800693 0.01481453
## [[3]]$`Case-II`
                         pval
               SNP
## 1 seq.rs7966314 0.03569512
## 2 seq.rs2072133 0.01355917
##
## [[3]]$`Case-III`
                SNP
                          pval
## 1 seq.rs12693591 0.02004173
## 2 seq.rs3771300 0.03990646
     seq.rs361525 0.02179672
## 4 seq.rs1800693 0.01382160
##
##
## [[4]]
## [[4]]$`Case-I`
               SNP
                         pval
## 1 seq.rs3769823 0.03369201
## [[4]]$`Case-II`
## NULL
##
## [[4]]$`Case-III`
               SNP
## 1 seq.rs3769823 0.01319883
## 2 seq.rs4844590 0.04305651
##
## [[5]]
## [[5]]$`Case-I`
## NULL
## [[5]]$`Case-II`
           SNP
                     pval
## 1 rs3761624 0.03637326
##
## [[5]]$`Case-III`
```

NULL