# 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.

# Association analysis on disease severity against GoldenGate SNP panels with several potentional covariates.

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.

Table 1: Genotype case I

Disease-level	AA	AB	BB
Severe	SNPCnt	SNPCnt	SNPCnt
Mild	SNPCnt	SNPCnt	SNPCnt

Table 2: Genotype case II

Disease-level	AA	AB/BB
Severe	SNPCnt	SNPCnt
Mild	SNPCnt	SNPCnt

Table 3: Genotype case III

Disease-level	AA/AB	BB
Severe	SNPCnt	SNPCnt
Mild	SNPCnt	SNPCnt

# Analysis set up and load necessary functions

#### Load up preprocessed data

Analysis results on disease severity – logistic regression model

```
## $`Top Tier IRF7 to TLR4`
## $`Top Tier IRF7 to TLR4`$`Case-I`
## SNP pval
## 1 seq.rs3806496 0.03249843
```

```
## 2 seq.rs2066843 0.03161207
## 3 seq.rs17244587 0.04618051
## $`Top Tier IRF7 to TLR4`$`Case-II`
                SNP
                          pval
## 1 seq.rs12987402 0.03525483
## 2 seq.rs469390 0.02409298
## 3 seq.rs3806265 0.03776518
## 4 seq.rs2066843 0.03371025
## $`Top Tier IRF7 to TLR4`$`Case-III`
                SNP
## 1 seq.rs17189298 0.04594427
## 2 seq.rs2770146 0.01637843
##
##
## $`BPIIFA1(PLUNC) to MYD88`
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
               SNP
## 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
## $`BPIIFA1(PLUNC) to MYD88`$`Case-II`
                SNP
                           pval
## 1 seq.rs2243592 0.003018021
## 2 seq.rs2257167 0.006062165
## 3 seq.rs2243599 0.006208847
## 4 seq.rs2254180 0.021865717
## 5 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
## $`BPIIFA1(PLUNC) to MYD88`$`Case-III`
                           pval
               SNP
## 1 seq.rs2280789 0.026207189
## 2 seq.rs2252930 0.010498389
## 3
     seq.rs2243592 0.006836451
     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
##
```

```
##
## $ OAS2 to TNFR
## $ OAS2 to TNFR $ Case-I
             SNP pval
## 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
##
## $`OAS2 to TNFR`$`Case-II`
              SNP
## 1 seq.rs7966314 0.02788702
## 2 seq.rs2072133 0.01039301
## $`OAS2 to TNFR`$`Case-III`
##
               SNP
## 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
##
##
## $`Additional_CASP8 to VDR`
## $`Additional_CASP8 to VDR`$`Case-I`
     SNP
                       pval
## 1 seq.rs3769823 0.0402697
## $`Additional_CASP8 to VDR`$`Case-II`
## NULL
##
## $ Additional_CASP8 to VDR $ Case-III
             SNP
                       pval
## 1 seq.rs3769823 0.01005097
## 2 seq.rs4844590 0.03257378
##
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
         SNP
                 pval
## 1 rs3761624 0.02619375
## $`Genotyped prior`$`Case-II`
          SNP
                    pval
## 1 rs3761624 0.02858216
## $`Genotyped prior`$`Case-III`
## NULL
```

Analysis results on disease severity – logistic with gender as covariate

```
## $`Top Tier IRF7 to TLR4`
```

```
## $`Top Tier IRF7 to TLR4`$`Case-I`
              SNP
## 1 seq.rs3738447 0.03095243
## 2 seq.rs2806117 0.02231901
## $`Top Tier IRF7 to TLR4`$`Case-II`
              SNP
                    pval
## 1 seq.rs2806117 0.02550568
##
## $`Top Tier IRF7 to TLR4`$`Case-III`
              SNP
                   pval
## 1 seq.rs2066842 0.02627682
## 2 seq.rs2917666 0.04877192
##
##
## $`BPIIFA1(PLUNC) to MYD88`
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
              SNP
                    pval
## 1 seq.rs2107538 0.01438484
## 2 seq.rs2280789 0.03260841
## 3 seq.rs3024492 0.02204524
## $`BPIIFA1(PLUNC) to MYD88`$`Case-II`
              SNP
## 1 seq.rs2107538 0.01714037
## 2 seq.rs4845623 0.01818976
## 3 seq.rs4129267 0.02053515
## 4 seq.rs3024492 0.01755576
## 5 seq.rs3848722 0.02404389
## 6 seq.rs443099 0.04551007
## $`BPIIFA1(PLUNC) to MYD88`$`Case-III`
              SNP
## 1 seq.rs2280789 0.04637065
## 2 seq.rs6694817 0.01024136
##
## $`OAS2 to TNFR`
## $`OAS2 to TNFR`$`Case-I`
               SNP
## 1 seq.rs1293749 0.04924504
## 2 seq.rs12367468 0.04644929
## 3 seq.rs6581568 0.01465038
##
## $ OAS2 to TNFR $ Case-II
##
               SNP
## 1 seq.rs1293749 0.0165843840
## 2 seq.rs15895 0.0482663324
## 3 seq.rs12367468 0.0022164286
## 4 seq.rs6581568 0.0009284449
## $`OAS2 to TNFR`$`Case-III`
## NULL
##
```

```
##
## $`Additional_CASP8 to VDR`
## $ Additional CASP8 to VDR \ Case-I
              SNP
                     pval
## 1 seq.rs2229095 0.04752476
##
## $`Additional_CASP8 to VDR`$`Case-II`
## NULL
##
## $ Additional_CASP8 to VDR $ Case-III
##
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
##
          SNP
                    pval
## 1 rs6721961 0.04200231
## $`Genotyped prior`$`Case-II`
   SNP
                    pval
## 1 rs2569190 0.04132563
## $`Genotyped prior`$`Case-III`
## NULL
```

# Analysis results on disease severity – logistic with region as covariate

```
## $`Top Tier IRF7 to TLR4`
## $`Top Tier IRF7 to TLR4`$`Case-I`
               SNP
## 1 seq.rs5743291 0.007589368
## 2 seq.rs17244587 0.029565288
## 3 seq.rs4986790 0.037814172
## 4 seq.rs10759930 0.024301793
## $`Top Tier IRF7 to TLR4`$`Case-II`
               SNP
## 1 seq.rs6722472 0.033356496
## 2 seq.rs3731612 0.023233536
## 3 seq.rs5743291 0.007166732
## 4 seq.rs17244587 0.024779377
## 5 seq.rs75059843 0.013924264
## 6 seq.rs4986790 0.037448234
## 7 seq.rs4986791 0.025379831
## $`Top Tier IRF7 to TLR4`$`Case-III`
              SNP
                         pval
## 1 seq.rs2917666 0.01499783
## 2 seq.rs3936292 0.01410505
## 3 seq.rs7022323 0.02726623
##
## $`BPIIFA1(PLUNC) to MYD88`
```

```
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
##
               SNP
                          pval
## 1 seq.rs7296163 0.046356411
## 2 seq.rs9272346 0.033613178
## 3 seq.rs2252930 0.007116561
## 4 seq.rs2243592 0.018099573
## 5 seq.rs2243599 0.049535474
## 6 seq.rs6517154 0.029281023
## 7 seq.rs1554606 0.039698733
##
## $`BPIIFA1(PLUNC) to MYD88`$`Case-II`
                SNP
                           pval
## 1 seq.rs13069802 0.048974384
## 2 seq.rs2252930 0.006260596
## 3 seq.rs2243592 0.006443205
## 4 seq.rs2243599 0.042282970
## 5 seq.rs6517154 0.021185432
## 6 seq.rs1554606 0.010554988
## $`BPIIFA1(PLUNC) to MYD88`$`Case-III`
##
               SNP
                         pval
## 1 seq.rs1046161 0.02513435
## 2 seq.rs9272346 0.03415691
## 3 seq.rs2250226 0.04812289
##
## $ OAS2 to TNFR
## $`OAS2 to TNFR`$`Case-I`
              SNP
## 1 seq.rs1293745 0.03445199
## 2 seq.rs1800630 0.04190544
## 3 seq.rs1041981 0.03180766
## $`OAS2 to TNFR`$`Case-II`
              SNP
                          pval
## 1 seq.rs2072137 0.007765022
## 2 seq.rs7953402 0.046495039
## 3 seq.rs1799964 0.044793060
## 4 seq.rs1800630 0.047761580
## 5 seq.rs1041981 0.045486371
## $ OAS2 to TNFR $ Case-III
             SNP
## 1 seq.rs1293745 0.04176694
## 2 seq.rs2280232 0.03598239
##
## $`Additional_CASP8 to VDR`
## $`Additional_CASP8 to VDR`$`Case-I`
              SNP
                          pval
## 1 seq.rs3890733 0.007209354
## $`Additional_CASP8 to VDR`$`Case-II`
##
                SNP
                          pval
```

```
## 1 seq.rs11208545 0.03722931
## 2 seq.rs6580642 0.04844294
##
## $ Additional_CASP8 to VDR $ Case-III
##
                SNP
                           pval
## 1 seq.rs3890733 0.003656025
## 2 seq.rs11168293 0.004391315
## 3 seq.rs4516035 0.011011403
## 4 seq.rs7139166 0.004782714
##
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
           SNP
                     pval
## 1 rs2569190 0.03129927
## 2 rs6721961 0.02408053
##
## $`Genotyped prior`$`Case-II`
           SNP
                     pval
## 1 rs2569190 0.01943644
## 2 rs6721961 0.01451079
## $`Genotyped prior`$`Case-III`
## NULL
```

# Analysis results on disease severity – logistic with social as covariate

```
sig.asso.SAT <- c(list())</pre>
results.logit <- logisticAnalysisWCorrV(sig.asso.SAT, snpFiles,
    sheets, dt, 6, row2use, params$sesStatus)
results.logit
## $`Top Tier IRF7 to TLR4`
## $`Top Tier IRF7 to TLR4`$`Case-I`
                SNP
## 1 seq.rs2193035 0.034177766
## 2 seq.rs5743291 0.004372155
## 3 seq.rs4986790 0.005361488
## 4 seq.rs4986791 0.027518773
## 5 seq.rs10759930 0.015191125
##
## $`Top Tier IRF7 to TLR4`$`Case-II`
##
                SNP
                           pval
## 1 seq.rs2193035 0.036090953
## 2 seq.rs5743291 0.003114019
## 3 seq.rs75059843 0.049610011
     seq.rs1927911 0.017150741
      seq.rs4986790 0.003999410
## 6 seq.rs4986791 0.006687538
## $`Top Tier IRF7 to TLR4`$`Case-III`
                SNP
                          pval
## 1 seq.rs3936292 0.02484239
```

```
## 2 seq.rs7022323 0.02953915
## 3 seq.rs10759930 0.02168704
##
##
## $`BPIIFA1(PLUNC) to MYD88`
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
                          pval
               SNP
## 1 seq.rs7296163 0.016802503
## 2 seq.rs2252930 0.009035721
## 3 seq.rs2243592 0.023568224
## 4 seq.rs2243599 0.038480553
## 5 seq.rs6517154 0.037488503
## 6 seq.rs3024496 0.045334709
## $`BPIIFA1(PLUNC) to MYD88`$`Case-II`
##
                SNP
                           pval
## 1 seq.rs7296163 0.021309670
## 2 seq.rs2252930 0.003443911
## 3 seq.rs2243592 0.019517942
## 4 seq.rs6517154 0.020560818
## 5 seq.rs10211925 0.043408749
## 6 seq.rs9376268 0.046019117
##
## $`BPIIFA1(PLUNC) to MYD88`$`Case-III`
               SNP
                        pval
## 1 seq.rs1046161 0.02900484
##
## $`OAS2 to TNFR`
## $ OAS2 to TNFR $ Case-I
               SNP
## 1 seq.rs1800630 0.03458350
## 2 seq.rs1041981 0.01775615
## $ OAS2 to TNFR $ Case-II
             SNP
## 1 seq.rs2072137 0.03815757
## 2 seq.rs7953402 0.03677857
## 3 seq.rs1800630 0.03404981
## 4 seq.rs1041981 0.02103300
## $ OAS2 to TNFR $ Case-III
## NULL
##
## $`Additional_CASP8 to VDR`
## $`Additional_CASP8 to VDR`$`Case-I`
                SNP
                         pval
## 1 seq.rs11574663 0.04743326
## 2 seq.rs11208545 0.04210803
## 3 seq.rs3890733 0.00734571
## $`Additional_CASP8 to VDR`$`Case-II`
##
                SNP
                         pval
```

```
## 1 seq.rs3769823 0.04331380
## 2 seq.rs11208545 0.01539316
## $`Additional_CASP8 to VDR`$`Case-III`
               SNP
                          pval
## 1 seq.rs2734648 0.046999929
## 2 seq.rs3890733 0.008839898
## 3 seq.rs11168293 0.009520885
## 4 seq.rs4516035 0.012611197
## 5 seq.rs7139166 0.006271190
##
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
## NULL
##
## $`Genotyped prior`$`Case-II`
     SNP
                    pval
## 1 rs2569190 0.02675012
## $`Genotyped prior`$`Case-III`
## NULL
```

# Analysis results on disease severity – logistic with breastfeeding as covariate

```
## $`Top Tier IRF7 to TLR4`
## $`Top Tier IRF7 to TLR4`$`Case-I`
               SNP
                         pval
## 1 seq.rs4925659 0.04857201
## 2 seq.rs7502875 0.02966858
## 3 seq.rs5030725 0.03166908
##
## $`Top Tier IRF7 to TLR4`$`Case-II`
               SNP
                          pval
## 1 seq.rs11795343 0.03356908
## 2 seq.rs10813831 0.03897348
## 3 seq.rs7502875 0.04425052
## 4 seq.rs5030725 0.03166908
## $`Top Tier IRF7 to TLR4`$`Case-III`
              SNP
                        pval
## 1 seq.rs469012 0.02343009
## 2 seq.rs469390 0.01958672
##
##
## $`BPIIFA1(PLUNC) to MYD88`
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
                SNP
## 1 seq.rs11063084 0.045638507
## 2 seq.rs2107538 0.009151977
## 3 seq.rs2257167 0.017594107
## 4 seq.rs2275913 0.013270682
## 5 seq.rs8193038 0.014041868
```

```
seq.rs7744 0.026364135
##
## $`BPIIFA1(PLUNC) to MYD88`$`Case-II`
## SNP
                    pval
## 1 seq.rs2107538 0.018246758
## 2 seq.rs2275913 0.008774505
## 3 seq.rs8193038 0.006582946
##
## $`BPIIFA1(PLUNC) to MYD88`$`Case-III`
##
              SNP
                        pval
## 1 seq.rs2257167 0.02022729
## 2 seq.rs2250889 0.04664627
##
## $ OAS2 to TNFR
## $`OAS2 to TNFR`$`Case-I`
               SNP
                          pval
## 1 seq.rs16824035 0.005067530
## 2 seq.rs1800692 0.003280719
## $`OAS2 to TNFR`$`Case-II`
              SNP
## 1 seq.rs10199181 0.042982420
## 2 seq.rs16824035 0.006314110
## 3 seq.rs1800692 0.009707555
## $`OAS2 to TNFR`$`Case-III`
## NULL
##
##
## $ Additional_CASP8 to VDR
## $`Additional_CASP8 to VDR`$`Case-I`
               SNP
                         pval
## 1 seq.rs11208545 0.03037701
## $`Additional_CASP8 to VDR`$`Case-II`
               SNP
## 1 seq.rs310202 0.01642526
## 2 seq.rs11208545 0.02557307
##
## $ Additional_CASP8 to VDR $ Case-III
##
              SNP
                         pval
## 1 seq.rs310247 0.03893488
## 2 seq.rs12743599 0.01310665
##
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
## NULL
##
## $`Genotyped prior`$`Case-II`
## NULL
##
## $`Genotyped prior`$`Case-III`
```

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

```
## $`Top Tier IRF7 to TLR4`
## $`Top Tier IRF7 to TLR4`$`Case-I`
              SNP
                        pval
## 1 seq.rs469390 0.03553412
## 2 seq.rs2770146 0.03031926
##
## $`Top Tier IRF7 to TLR4`$`Case-II`
               SNP
                     pval
## 1 seq.rs12987402 0.04484957
## 2 seq.rs469390 0.03136834
## 3 seq.rs3806265 0.04864552
## 4 seq.rs2066843 0.04292077
##
## $`Top Tier IRF7 to TLR4`$`Case-III`
             SNP
                       pval
## 1 seq.rs2770146 0.02269727
##
##
## $`BPIIFA1(PLUNC) to MYD88`
## $`BPIIFA1(PLUNC) to MYD88`$`Case-I`
##
               SNP
## 1 seq.rs2252930 0.032963115
## 2 seq.rs2243592 0.002065785
## 3 seq.rs2257167 0.004097110
## 4 seq.rs2243599 0.002391277
## 5 seq.rs2254180 0.016497081
## 6
     seq.rs2254315 0.014203726
## 7
     seq.rs2069832 0.026046287
## 8 seq.rs1474347 0.040922019
## 9 seq.rs8193036 0.046061333
## 10 seq.rs8193038 0.049507445
## $`BPIIFA1(PLUNC) to MYD88`$`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
##
## $`BPIIFA1(PLUNC) to MYD88`$`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
##
##
## $ OAS2 to TNFR
## $`OAS2 to TNFR`$`Case-I`
               SNP
## 1 seq.rs2072133 0.02734773
## 2 seq.rs12693591 0.01358428
## 3 seq.rs361525 0.02179245
## 4 seq.rs1800693 0.01481453
##
## $ OAS2 to TNFR $ Case-II
              SNP
## 1 seq.rs7966314 0.03569512
## 2 seq.rs2072133 0.01355917
## $`OAS2 to TNFR`$`Case-III`
               SNP
## 1 seq.rs12693591 0.02004173
## 2 seq.rs3771300 0.03990646
     seq.rs361525 0.02179672
## 4 seq.rs1800693 0.01382160
##
##
## $ Additional_CASP8 to VDR
## $`Additional_CASP8 to VDR`$`Case-I`
              SNP
                        pval
## 1 seq.rs3769823 0.03369201
## $`Additional_CASP8 to VDR`$`Case-II`
## NULL
##
## $`Additional_CASP8 to VDR`$`Case-III`
##
              SNP
## 1 seq.rs3769823 0.01319883
## 2 seq.rs4844590 0.04305651
##
## $`Genotyped prior`
## $`Genotyped prior`$`Case-I`
## NULL
##
## $`Genotyped prior`$`Case-II`
          SNP
                    pval
## 1 rs3761624 0.03637326
## $`Genotyped prior`$`Case-III`
## NULL
```