Test with genus abundance data

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5/14/2020

Real data

From https://github.com/chvlyl/PLEASE

Inflammation, Antibiotics, and Diet as Environmental Stressors of the Gut Microbiome in Pediatric Crohn's Disease. Cell Host & Microbe. 2015

Load the raw data

The "unclassfied" taxa were removed and the total relative abundance in each sample were normalized to be one. "P", "F", "G", "S" at the beginning of each file indicate taxonomic levels "phylum", "family", "genus", "species".

We use genus abundance data.

```
PLEASE.raw =
 read excel("./real data/G Remove unclassfied Renormalized Merge Rel MetaPhlAn Result.xlsx",
           col names = T) %>%
 as.data.frame()
row.names(PLEASE.raw) = PLEASE.raw[,1]
PLEASE.raw = PLEASE.raw[,-1]
colname = as.numeric(colnames(PLEASE.raw))
samplepoint = as.Date(colname, origin = "1900-01-01")
head(PLEASE.raw[,1:5])
##
                      1132619
                               1132650
                                           1132678
                                                      1132709
                                                               1132984
## g__Bacteroides
                    ## g__Ruminococcus
                    0.0000000 \quad 0.0000000 \quad 0.006130341 \quad 0.000000000 \quad 4.9631993
## g__Akkermansia
                    ## g_Bifidobacterium 0.3756509 2.0725290 1.033117545 0.602090120
                                                             1.3608252
## g__Escherichia
                   38.3172402 21.4206238 41.259008130 36.090417220
taxa.raw <- data.frame(t(PLEASE.raw),</pre>
                   row.names = strtrim(samplepoint,7))
### Make sure you load the data correctly
cat('samples', 'taxa', dim(taxa.raw), '\n')
## samples taxa 335 105
taxa.raw[1:3,1:3]
         g_Bacteroides g_Ruminococcus g_Faecalibacterium
## 5001-01
            0.42722310
                         0.00000000
            0.23188470
## 5001-02
                         0.000000000
                                                   0
```

Each row represent a sample.

Load total non-human read counts

```
human.read.file <- './real data/please_combo_human_reads.xlsx'</pre>
human.read <-
  read_excel(human.read.file, col_names = T) %>%
  as.data.frame() %>%
  mutate(
    Sample = strtrim(as.Date(Sample,origin = "1900-01-01"),7)
head(human.read)
##
     Sample NonHumanReads TotalReads HumanReads
                                                    HumanPer GroupFcp GroupPcdai
## 1 1910-12
                  17525422
                            19515697
                                         1990275 10.19832959
                                                                Combo
                                                                           Combo
                                                                           Combo
## 2 1910-12
                  18089762
                                           96168 0.52880444
                                                                Combo
                           18185930
## 3 1910-12
                  27311061 27338002
                                           26941 0.09854781
                                                                Combo
                                                                           Combo
## 4 1910-12
                 11051439 11092808
                                           41369 0.37293536
                                                                Combo
                                                                           Combo
## 5 1910-12
                  9434025
                                           58171 0.61282980
                                                                Combo
                                                                           Combo
                            9492196
## 6 1910-12
                 23327496
                            23634581
                                          307085 1.29930382
                                                                Combo
                                                                           Combo
## first column: sample id
### Filter low depth samples (low non human reads)
low.depth.samples <- subset(human.read,NonHumanReads<10000)</pre>
head(low.depth.samples[,1:5])
        Sample NonHumanReads TotalReads HumanReads
                                                      HumanPer
## 56 5010-02
                       1014
                                   1104
                                          90 8.15217391
## 85 5018-03
                        6101
                                   6121
                                               20 0.32674400
## 98 5023-04
                        1954
                                   2679
                                              725 27.06233669
## 257 6007-01
                        1809
                                   4249
                                              2440 57.42527654
## 309 7001-02
                        9965
                                   9971
                                                 6 0.06017451
## 310 7001-03
                        566
                                    613
                                                47 7.66721044
### Delete these samples from PLEASE data.
# row.names(taxa.raw)
# row.names(low.depth.samples)
row.names(taxa.raw) [which(row.names(taxa.raw) %in% low.depth.samples$Sample)]
## [1] "5018-03" "7001-02" "7003-02" "7003-03" "7009-03" "7010-03"
### Before deletion
dim(taxa.raw)
## [1] 335 105
### After deletion
taxa.raw <- taxa.raw[-which(rownames(taxa.raw) %in% low.depth.samples$Sample),]
dim(taxa.raw)
## [1] 329 105
```

Filter low abundant bacterial data

```
### Filter low abundant bacterial data
filter.index1 <- apply(taxa.raw,2,function(X){sum(X>0)>0.4*length(X)})
filter.index2 <- apply(taxa.raw,2,function(X){quantile(X,0.9)>1})
taxa.filter <- taxa.raw[,filter.index1 & filter.index2]</pre>
taxa.filter <- 100*sweep(taxa.filter, 1, rowSums(taxa.filter), FUN="/")
cat('after filter:','samples','taxa',dim(taxa.filter),'\n')
## after filter: samples taxa 329 18
cat(colnames(taxa.filter),'\n')
## g_Bacteroides g_Ruminococcus g_Faecalibacterium g_Bifidobacterium g_Escherichia g_Clostridium ,
head(rowSums(taxa.filter))
## 5001-01 5001-02 5001-03 5001-04 5002-01 5002-02
               100
                       100
                                100
       100
                                        100
After filter, there remains 18 bacteria in the taxa table.
taxa.data <- taxa.filter
dim(taxa.data)
## [1] 329 18
```

Load sample information

```
head(sample.info)
```

```
##
     Sample Subject Species.Cluster
                                       Cluster Treatment FCPResponse
                                                                        Type Time
## 1
               4000
                           cluster 1 cluster 1
                                                                    NA COMBO
                                                       NA
                                                                    NA COMBO
## 2
       4001
               4001
                           cluster 1 cluster 1
                                                       NA
                                                                               NA
## 3
       4002
               4002
                           cluster 1 cluster 1
                                                       NA
                                                                    NA COMBO
                                                                               NA
## 4
       4004
               4004
                           cluster 1 cluster 1
                                                       NA
                                                                    NA COMBO
                                                                               NA
       4005
## 5
               4005
                           cluster 1 cluster 1
                                                       NA
                                                                    NA COMBO
                                                                               NA
## 6
       4006
                           cluster 1 cluster 1
                                                                    NA COMBO
               4006
                                                       NA
                                                                               NA
##
     BristolScore FCP PCDAI PUCAI log.FCP Group Response Antibiotics.visit
## 1
                                        NA COMBO
               NA NA
                          NA
                                                        NA
                                                                      Not.Use
## 2
               NA
                   NA
                          NA
                                NA
                                        NA COMBO
                                                                      Not.Use
                                                        NΑ
                                        NA COMBO
## 3
               NA
                   NA
                          NA
                                NA
                                                                      Not.Use
## 4
               NA
                   NA
                          NA
                                NA
                                        NA COMBO
                                                                      Not.Use
                                                        NΑ
## 5
                   NA
                          NA
                                NA
                                        NA COMBO
                                                                      Not.Use
## 6
               NA NA
                          NΑ
                                NA
                                        NA COMBO
                                                                      Not.Use
                                                        NΑ
##
     Steroids Treatment.Specific Disease NonHumanReads
                                                                      Human.Per
## 1
                               NA Control
                                                17525422
           NA
                                                                    10.19832959
## 2
           NA
                               NA Control
                                                18089762
                                                           0.52880444000000004
## 3
           NA
                               NA Control
                                                27311061 9.8547805000000002E-2
## 4
           NA
                               NA Control
                                                11051439
                                                           0.37293536300000002
## 5
           NA
                               NA Control
                                                 9434025
                                                           0.61282980399999998
## 6
           NA
                               NA Control
                                                23327496
                                                                    1.299303817
##
                 Fungi.Per
                                       Distance
                                                           Bact.Div
## 1
       1.02707940499236E-4
                              0.452380952380952 133.36064650458701
## 2 7.5180646378874396E-3 0.42857142857142899
                                                   173.757418771479
## 3 8.2750355249838195E-4 0.40476190476190499 90.479491742827904
## 4 3.5651465840783299E-3 0.35714285714285698
                                                   103.213875049946
## 5 1.5984693701786901E-2
                                             0.5 149.26361597807599
```

```
## 6 6.1986935931743403E-3 0.28571428571428598
                                                 112.488017364215
##
        Species.Distance
## 1 0.40449438202247201
## 2 0.33707865168539303
## 3 0.33707865168539303
       0.426966292134831
## 5 0.43820224719101097
## 6
       0.235955056179775
create\ covariates,\ Time,\ Treatment(antiTNF+EEN)
complete_subject =
  sample.info %>%
  filter(Sample %in% rownames(taxa.data)) %>%
  filter(Treatment.Specific!='PEN')%>%
  dplyr::select(Sample, Time, Subject, Response, Treatment. Specific) %>%
  group_by(Subject) %>%
  summarise(count = n()) %>%
  filter(count==4)
reg.cov =
  sample.info %>%
  filter(Subject %in%complete_subject$Subject) %>%
  mutate(Treat=ifelse(Treatment.Specific=='antiTNF',1,0)) %>%
  dplyr::mutate(Subject=paste('S',Subject,sep='')) %>%
  dplyr::mutate(Time=ifelse(Time=='1',0,ifelse(Time=='2',1,ifelse(Time=='3',4,ifelse(Time=='4',8,NA))))
  dplyr::mutate(Time.X.Treatment=Time*Treat) %>%
  dplyr::select(Sample,Subject,Time,Response,Treat,Time.X.Treatment,everything())
take out first time point
             <- subset(reg.cov,Time==0)
reg.cov.t1
rownames(reg.cov.t1) <- reg.cov.t1$Subject</pre>
reg.cov.t234 <- subset(reg.cov,Time!=0)
reg.cov.t234 <- data.frame(
  baseline.sample=reg.cov.t1[reg.cov.t234$Subject, 'Sample'],
  baseline.subject=reg.cov.t1[reg.cov.t234$Subject,'Subject'],
  reg.cov.t234,
  stringsAsFactors = FALSE)
head(reg.cov.t234)
##
     baseline.sample baseline.subject Sample Subject Time
                                                                Response Treat
## 2
             5001-01
                                S5001 5001-02
                                                S5001
                                                         1 Non.Response
                                                                             1
## 3
             5001-01
                                S5001 5001-03
                                                S5001
                                                          4 Non.Response
                                                                             1
## 4
             5001-01
                                S5001 5001-04
                                                S5001
                                                          8 Non.Response
                                                                             1
## 6
             5002-01
                                S5002 5002-02
                                                S5002
                                                         1 Non.Response
                                                                             1
## 7
             5002-01
                                S5002 5002-03
                                                S5002
                                                          4 Non.Response
                                                                             1
## 8
             5002-01
                                S5002 5002-04
                                                S5002
                                                          8 Non.Response
                                                                             1
##
     Time.X.Treatment Species.Cluster
                                        Cluster Treatment FCPResponse
                                                                            Type
## 2
                                                                     0 PLEASE-T2
                   1
                            cluster 2 cluster 2
                                                  antiTNF
## 3
                    4
                            cluster 2 cluster 2
                                                  antiTNF
                                                                     O PLEASE-T3
## 4
                            cluster 2 cluster 2 antiTNF
                                                                     0 PLEASE-T4
```

```
## 6
                            cluster 2 cluster 1
                                                 antiTNF
                                                                    0 PLEASE-T2
## 7
                            cluster 2 cluster 1
                                                 antiTNF
                                                                    O PLEASE-T3
                            cluster 2 cluster 1
                                                 antiTNF
                                                                    O PLEASE-T4
## 8
                   8
    BristolScore FCP PCDAI PUCAI
##
                                             log.FCP Group Antibiotics.visit
## 2
               6
                  607
                                25 6.4085287910595001 PLEASE
                                                                       Not.Use
## 3
               6 867
                         NA
                                20 6.7650389767805397 PLEASE
                                                                       Not.Use
## 4
                               15 6.3225652399272798 PLEASE
                                                                       Not.Use
               6 557
                          5
                               10 6.8564619845945902 PLEASE
## 6
               6 950
                         NA
                                                                       Not.Use
## 7
               6 1947
                         NA
                               50 7.5740450053722004 PLEASE
                                                                       Not.Use
## A
                         35
                               40 7.5390270558239996 PLEASE
                                                                       Not.Use
               6 1880
     Steroids Treatment.Specific Disease NonHumanReads
                                                                 Human.Per
## 2 Not.Use
                                   Crohn
                                              1350309
                                                               89.31803171
                        antiTNF
     Not.Use
                                                       19.689170000000001
## 3
                        antiTNF
                                   Crohn
                                              10946591
## 4
     Not.Use
                                             14230882 0.8513367339999998
                        antiTNF
                                   Crohn
## 6
         Use
                                   Crohn
                                             12020377
                        antiTNF
                                                               17.08929796
## 7
         Use
                         antiTNF
                                   Crohn
                                               1910666
                                                       88.544540839999996
## 8
                                                606565
         Use
                        antiTNF
                                   Crohn
                                                                89.5498726
##
                Fungi.Per
                                      Distance
                                                        Bact.Div
## 2 7.0946724046125703E-2 0.69047619047619002 79.225334004595695
## 3 1.2168171808008501E-2 0.52272727272727304 66.915876889694502
## 4
          1.26499538117174 0.59090909090909105 53.865413747151202
       1.05154771767974E-2 0.42857142857142899 81.330542193164007
## 7 6.6678320543726605E-2 0.59523809523809501 89.862102084722594
##
        Species.Distance
## 2 0.82417582417582402
## 3 0.72527472527472503
## 4 0.7333333333333295
## 6 0.59550561797752799
## 7 0.75280898876404501
## 8 0.85393258426966301
taxa_all = colnames(taxa.data)
store = function(taxa){
# X: Baseline abundance time Treat
# Y: Response abundance at time 1 4 8
X <- data.frame(</pre>
      Baseline=taxa.data[reg.cov.t234$baseline.sample,taxa]/100,
      reg.cov.t234[,c('Time','Treat')])
rownames(X) <- reg.cov.t234$Sample</pre>
Y <- data.frame(
      Abundance=taxa.data[reg.cov.t234$Sample, taxa]/100,
      reg.cov.t234[,c('Time','Treat')])
return(list(X = X,
           Y = Y)
}
store_results = lapply(taxa_all, store)
names(store_results) = taxa_all
#example
```

head(store_results\$g__Bacteroides\$X) Baseline Time Treat ## ## 5001-02 0.004514608 ## 5001-03 0.004514608 ## 5001-04 0.004514608 1 ## 5002-02 0.732645555 ## 5002-03 0.732645555 1 ## 5002-04 0.732645555 all_set = cbind(reg.cov,taxa.data[reg.cov\$Sample,]) baseline_set = cbind(reg.cov.t1, taxa.data[reg.cov.t1\$Sample,]) visit_set = cbind(reg.cov.t234, taxa.data[reg.cov.t234\$Sample,]) test_genus_Eriz = all_set %>% pivot_longer(g_Bacteroides : g_Escherichia, names_to = "genus_type", names_prefix = "g__", values_to = "abundance" test_genus_Eriz %>% ggplot(aes(x = Time, y = abundance)) + geom_point(aes(x = Time, y = abundance,color = genus_type),alpha = 0.8) 100 -75 genus_type **Bacteroides** abundance Bifidobacterium 50 -Escherichia Faecalibacterium Ruminococcus 25 -0 -2 6

Time

max(taxa.data\$g__Bacteroides)

```
## [1] 99.07837
```

save.image(file = "Explore_EricZ.RData")