Large-scale genomic investigation of the gut microbiome on Parkinson's disease etiology - Code

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Preprocessing GWAS Summary Datasets

The first step of this pipeline was to identify datasets we'd be performing our association studies on. Go to https://mibiogen.gcc.rug.nl/ to find the repository of datasets provided by Kurilshikov et. al, 2021 (https://pubmed.ncbi.nlm.nih.gov/33462485/). On this website, you'll find 6 links:

- 1. MBG.allHits.p1e4.txt -> top hit variants (p < 1e-4) for any level of the bacterial hierarchy
- 2. MiBioGen_QmbQTL_summary_phylum.zip (2.4 GB) -> summary statistics of bacterial phyla
- 3. MiBioGen_QmbQTL_summary_class.zip (4.4 GB) -> summary statistics of bacterial classes
- 4. MiBioGen QmbQTL summary order.zip (5.4 GB) ->summary statistics of bacterial orders
- 5. MiBioGen_QmbQTL_summary_family.zip (9.5 GB) -> summary statistics of bacterial families
- 6. MiBioGen_QmbQTL_summary_genus.zip (35.0 GB) -> summary statistics of bacterial genera

In our case, we wanted explore the bacterial genera taxanomy. In latter analyses, we hope to run family, order, class, and phylum taxanomies of bacteria. When you unzip the MiBioGen_QmbQTL_summary_genus.zip, it will look something like this:

- MiBioGen_QmbQTL_summary_genus:
 - genus.Clostridiuminnocuumgroup.id.14397.summary.txt.gz
 - $-\,$ genus. Eubacteriumbrachygroup.
id.11296.summary.txt.gz

Unzip the subfolders and you get a table like this:

bac	chr	bp	rsID	ref.allele	eff.allele	beta	SE	Z weighted	P weighted	N	Cohorts
name	5	71186626	$\mathrm{rs}6890185$	С	Τ	0.113	0.023	4.868	1.122e-06	4166	20

Now perform these data pre-processing steps (on sample data):

Import "reticulate" to incorporate python functionalities.

```
knitr::opts_chunk$set(tidy.opts=list(width.cutoff=80), tidy=TRUE)
library(reticulate)
use_virtualenv("base")
use_python("/Volumes/T7Touch/Applications/anaconda3/bin/python")
```

Each .gz file of the sum stats will give a txt file, convert these into csv files.

```
knitr::opts_chunk$set(tidy.opts=list(width.cutoff=80), tidy=TRUE)
sumstats_root <- "/Users/rodrigosandon/Documents/GitHub/ParkisonsMicrobiomeStudy/MB-PD_Association_Stud
source_python("Utilities.py")
out_paths <- txt_to_csv_files_in_root(sumstats_root)
# If this doesn't work, run the py file itself with the process execution</pre>
```

Add a "p" column to each sum stat csv file.

```
source python("Utilities.py")
knitr::opts_chunk$set(tidy.opts=list(width.cutoff=80), tidy=TRUE)
add_p_col_to_df <- function(bac_sumstat_path, new_col_name, out_path) {</pre>
  df <- read.csv(bac_sumstat_path, header = TRUE, sep = ",")</pre>
  df[new_col_name] <- 2*pnorm(-abs(df$beta/df$SE))</pre>
  eles_of_bac_path <- strsplit(bac_sumstat_path, "/")</pre>
  name_of_bac_sumstat <- eles_of_bac_path[[1]][length(eles_of_bac_path[[1]])]</pre>
  \# \hat{f}inding the name original name of the bac sumstat file
  new_name <- paste("addedP_", name_of_bac_sumstat, sep = "")</pre>
  new_path <- paste(out_path, new_name, sep = "")</pre>
  #print(new_path)
  # Export
  write.csv(df, new_path, row.names = FALSE, quote = FALSE)
}
create_lst_of_file_paths <- function(root_path, files_endswith_str) {</pre>
  files <- list.files(path = root_path, pattern = files_endswith_str, full.names = TRUE,
                       recursive = FALSE)
 return (files)
add_p_col_to_csvs_in_root <- function(root_path, out_path, files_endswith_str,
                                        new_col_name, omit_csvs_that_contain) {
  dir.create(out_path, showWarnings = FALSE) #uncomment if dir hasn't been created yet, else, keep comm
  files <- create_lst_of_file_paths(root_path, files_endswith_str)</pre>
  # Reminder: current files in the list should be in csv format
  for (bac_sumstat_path in files) {
    print(paste("Working on ...", bac_sumstat_path))
    start.time <- Sys.time()</pre>
    if (grepl(omit_csvs_that_contain, bac_sumstat_path, fixed = TRUE) == FALSE) {
      add_p_col_to_df(bac_sumstat_path, new_col_name, out_path)
    }
    end.time <- Sys.time()</pre>
    print(paste("Time to process: ", end.time - start.time))
 }
```

Polygenic Risk Score Analysis

do

Make a txt file of all of the genera you will be performing PRS on

```
#Identify the path where you've modified the sumstats
sumstats_path <- "/Users/rodrigosandon/Documents/GitHub/ParkisonsMicrobiomeStudy/MB-PD_Association_Study
#Define the group of files to identify within ^ this root path
look_for <- "addedP_"

#Identify path where you'll locate the txt file
txt_out <- "/Users/rodrigosandon/Documents/GitHub/ParkisonsMicrobiomeStudy/MB-PD_Association_Study_Pipe
source_python("Utilities.py")

files_lst <- find_paths_startswith(sumstats_path, look_for)
listdir_to_txt_file(files_lst, txt_out)</pre>
```

Using PRSice.R, perform PRS and acquire the output file

```
### Polygenic risk score analyses of 119 bacteria genuses versus PD risk

## Make a list of summary stats file
ls *csv > /Users/rodrigosandon/Documents/GitHub/ParkisonsMicrobiomeStudy/MB-PD_Association_Study_Pipeli

## Format these files
cat genuses.txt | while read LINE

do
    echo $LINE
    sed 's/\"//g' $LINE | sed 's/, / /g' > temp.txt
    awk '{print $0"\t"$2":"$3}' temp.txt | sed 's/chr\:bp/ID/' > $LINE.temp_formatted.txt
    rm temp.txt

done

## Identify independent risk SNPs using our in-house LD reference data for European populations (/data/
Rscript /data/LNG/pdMeta5v2/leaveOneOutPrsice/PRSice_linux/PRSice.R --cov-file /data/LNG/saraB/WGS/noag
Rscript /data/LNG/pdMeta5v2/leaveOneOutPrsice/PRSice_linux/PRSice.R --cov-file /data/LNG/saraB/WGS/noag
## Remove NeuroX individuals & extract nominated variants
cat genuses_formatted_list.txt | while read LINE
```

```
plink --bfile /data/LNG/saraB/concat_HARDCALLS_PD_september_2018_no_cousins --remove-fam NeuroX.txt --e.

## Make score files
cat genuses_formatted_list.txt | while read LINE
do
awk '{print $14, $6, $7}' addedPgenus.$LINE.summary.txt.csv.temp_formatted.txt | sed '1d' > $LINE.tosco
done

## Make sure score files have 3 expected fields rather than 2
cat genuses_formatted_list.txt | while read LINE
do
grep ":" $LINE.toscore.txt > true_$LINE.toscore.txt
done

## Calculate scores
cat genuses_formatted_list.txt | while read LINE
do
plink --bfile pruned_$LINE --score $LINE.toscore.txt --make-bed --out pruned_$LINE
done
```

Run PRS (logistic regression) in R

```
#install.packages("data.table")
library("data.table")
listOfProfiles <- read.table("genera_formatted_list.txt", header = T)</pre>
names(listOfProfiles) <- c("id")</pre>
covs1 <- fread("/data/LNG/saraB/WGS/noage_toPRSice_phenosAndCovs_renamed.tab", header = T)</pre>
covs2 <- fread("/data/LNG/saraB/concat_HARDCALLS_PD_september_2018_no_cousins.fam", header = F)</pre>
colnames(covs2) <- c("FID", "IID", "MAT", "PAT", "SEX", "PHENO")</pre>
covsfinal <- merge (covs1, covs2, by ="FID")</pre>
covsfinal$CASE <- covsfinal$PHENO.x - 1</pre>
outPut <- matrix(ncol = 4, nrow = length(listOfProfiles$id), NA)</pre>
colnames(outPut) <- c("genus", "b", "se", "p")</pre>
for(i in 1:length(listOfProfiles$id))
{
    profileName <- as.character(listOfProfiles$id[i])</pre>
    profile <- fread(file = paste(profileName, ".profile", sep = ""), header = T)</pre>
    profile$index <- paste(profile$FID, profile$IID, sep = "")</pre>
    data <- merge(covsfinal, profile, by = "index")</pre>
    meanControls <- mean(data$SCORE[data$CASE == 0])</pre>
    sdControls <- sd(data$SCORE[data$CASE == 0])</pre>
    data$zSCORE <- (data$SCORE - meanControls)/sdControls</pre>
    grsTest <- glm(CASE ~ zSCORE + SEX + PC1 + PC2 + PC3 + PC4 + PC5 + PC6 + PC7 + PC8 + PC9 + PC10 + D
    beta <- summary(grsTest)$coefficients["zSCORE","Estimate"]</pre>
    se <- summary(grsTest)$coefficients["zSCORE","Std. Error"]</pre>
    p <- summary(grsTest)$coefficients["zSCORE","Pr(>|z|)"]
    outPut[i,1] <- profileName</pre>
    outPut[i,2] <- beta
    outPut[i,3] <- se</pre>
    outPut[i,4] <- p
}
write.table(outPut, "Genus_PRS.tab", quote = F, sep = "\t", row.names = F)
```

```
## SAMPLE RESULT:
# Call:
# qlm(formula = CASE ~ zSCORE + SEX + PC1 + PC2 + PC3 + PC4 + PC5 +
   PC6 + PC7 + PC8 + PC9 + PC10, family = "binomial", data = data)
# Deviance Residuals:
# Min 1Q Median
                        3Q
                              Max
# -1.919 -1.003 -0.810 1.278 1.796
# Coefficients:
           Estimate Std. Error z value Pr(>|z|)
# (Intercept) 0.43040 0.03974 10.831 < 2e-16 ***
# zSCORE 0.03021 0.01424
                              2.121 0.0339 *
# SEX
           -35.38600 2.73911 -12.919 < 2e-16 ***
# PC1
           50.17593 2.86877 17.490 < 2e-16 ***
# PC2
           10.63757 2.68575 3.961 7.47e-05 ***
# PC3
# PC4
            0.63048 2.65991 0.237 0.8126
# PC5
           16.19265 2.70187 5.993 2.06e-09 ***
           -24.20283 2.74525 -8.816 < 2e-16 ***
# PC6
# PC7
            1.61607 2.62627 0.615 0.5383
# PC8
           12.66905 2.70987 4.675 2.94e-06 ***
           -5.96044 2.64009 -2.258 0.0240 *
# PC9
         -16.18338 2.65955 -6.085 1.16e-09 ***
# PC10
# ---
# Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
# (Dispersion parameter for binomial family taken to be 1)
   Null deviance: 35275 on 26385 degrees of freedom
# Residual deviance: 34124 on 26373 degrees of freedom
# AIC: 34150
# Number of Fisher Scoring iterations: 4
```

Linkage Disequilibirum Score Regression (LDSC) Analysis

Data Preprocessing

```
pre_ldsc_chrPosRs <- source_python("/Users/rodrigosandon/Documents/GitHub/ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-ParkisonsMicrobiomeStudy/MB-Parkiso
```

LDSC

```
Note: Make sure you are running in a Python 2 environment.
"""
import pandas as pd
```

```
import csv
import os
import time
import os.path
from os import path
# the PD file for LD SR run is : /Volumes/T7Touch/NIHSummer2021/Data/LDSR_analysis/nalls_onlyRsIDs.sums
# Ex: /Volumes/Passport/119Bacs_addedP/addedPgenus..Clostridiuminnocuumgroup.id.14397.summary.txt.csv
def formatSumStatsForBac(csvPath):
   newFileName = "formatted." + csvPath.split("/")[4].replace(
        ".txt", "").replace("..", ".").replace("addedPgenus", "addedP.genus")
   newFilePath = "/Volumes/Passport/formatted119Bacs_addedP/%s" % (
       newFileName)
    if path.exists(newFilePath) == False:
       df = pd.read_csv(csvPath)
       newDf = df[['rsID', 'eff.allele', 'ref.allele',
                    'beta', 'SE', 'N', 'pDerived']].copy()
       newDf.insert(3, 'Zscore', newDf['beta'] / newDf['SE'], True)
       newDf = newDf.drop(['beta', 'SE'], axis=1)
       newDf = newDf.rename(columns={
                             'rsID': 'snpid', 'pDerived': 'P-value', 'eff.allele': 'A1', 'ref.allele':
        # Now formatted.addedP.genus.Clostridiuminnocuumgroup.id.14397.summary.csv
       newDf.to_csv(newFilePath, index=False)
   else:
       print("File %s already exists" % (newFilePath))
    return newFilePath
def CSVtoTXT(csv_file, txtFile):
    with open(txtFile, "w") as my_output_file:
        with open(csv_file, "r") as my_input_file:
            [my output file.write(" ".join(row)+'\n')
            for row in csv.reader(my_input_file)]
       my_output_file.close()
# ex csv file now: /Volumes/Passport/formatted119Bacs_addedP/formatted.addedP.genus.Clostridiuminnocuum
def mungeDataCall(csv_file):
   txtFilePath = csv_file.replace(".csv", ".txt")
    \# /Volumes/Passport/formatted119Bacs_addedP/formatted.addedP.genus.Clostridiuminnocuumgroup.id.1439
   newFilePath = txtFilePath.replace(
        "formatted119Bacs_addedP", "munge119Bacs_output").replace(".txt", "")
    if path.exists(txtFilePath) == False:
```

```
CSVtoTXT(csv_file, txtFilePath)
   else:
       print("File %s already munged" % (newFilePath))
    # os.chdir("/Users/rodrigosandon/ldsc")
   cmd = "./munge_sumstats.py \
       --sumstats %s \
       --out %s \
       --merge-alleles /Volumes/T7Touch/NIHSummer2021/Data/LDSR_analysis/ldsc/w_hm3.snplist" % (txtFil
   if path.exists(newFilePath + ".sumstats.gz") == False: # only if .gz file don't exist
       os.system(cmd)
   return newFilePath
# ex gx bac name munged: /Volumes/Passport/munge119Bacs_output/formatted.addedP.genus.Clostridiuminnocu
def LDscore_regression(munged_bac_output):
   print("munged_bac_output: ", munged_bac_output)
   LDSR_outName = "/Volumes/T7Touch/NIHSummer2021/Code/LDSC_analysis2/results/%s_ldscResults" % (
       munged_bac_output.split("/")[4].split(".")[3]) # <--only bac name</pre>
   # os.chdir("/Users/rodrigosandon/ldsc")
   cmd = "./ldsc.py \
       --rg %s,/Volumes/T7Touch/NIHSummer2021/Code/LDSC_analysis2/munged_META5_all_with_rsid.sumstats.
       --ref-ld-chr /Volumes/T7Touch/NIHSummer2021/Data/LDSR_analysis/ldsc/eur_w_ld_chr/ \
       --out %s " % (munged_bac_output + ".sumstats.gz", LDSR_outName)
   os.system(cmd)
   return LDSR_outName
###MAIN###
masterDir = "/Volumes/Passport/119Bacs_addedP/"
for root, dirs, files in os.walk(masterDir):
   for name in files:
       start = time.time()
       bacPathToProcess = os.path.join(root, name)
       print("Processing", bacPathToProcess)
       newFilePath1 = formatSumStatsForBac(bacPathToProcess)
       newFilePath2 = mungeDataCall(newFilePath1)
       LDSR_outName = LDscore_regression(newFilePath2)
```

Bayesian Colocalization

```
# addedPgenus.CandidatusSoleaferrea.id.11350.summary.txt.csv on request
# META5_all_with_rsid.txt upon request
# nallsEtAl2019_excluding23andMe_allVariants.tab from Nalls et. al 2019.
library("data.table")
install.packages("coloc")
source("https://bioconductor.org/biocLite.R")
biocLite("snpStats")
install.packages("robustbase")
library("robustbase")
library("coloc")
library("tidyverse")
#The file that contains SNPs with significance of 5E-N for both disease and phenotype
masterDir <- "/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/p smr multi mrQTL belowFDR"
files <- list.files(path=masterDir, pattern="*.tab", full.names = TRUE, recursive = FALSE)
#Run this same pipeline for other bacterial taxa
Candida_instruments <- fread(file="/Volumes/Passport/119Bacs_addedP/addedPgenus.CandidatusSoleaferrea.in
for (k in files) {
  TOCOLOC <- read.table(k, header = T, sep="\t")
  nameOfTest <- str_replace(str_split(k, "/", n = Inf, simplify = TRUE)[[8]], ".tab", "")
  #Just need a subset of file that contains SNPs with significance of 5E-N for both disease and phenoty
  SNPlist <- subset(TOCOLOC, select=c(topSNP, topSNP_bp, topSNP_chr))</pre>
  #For each SNP, search through phenotype summary stats to get 1MB of variants b4 and after SNP, this i
  for(i in 1:length(SNPlist$topSNP)) {
   thisSNP <- SNPlist$topSNP[i]</pre>
   thisChr <- SNPlist$topSNP_chr[i]</pre>
   thisBP <- SNPlist$topSNP_bp[i]</pre>
   thisBpLow <- thisBP - 1000000
   thisBpHigh <- thisBP + 1000000
   getRegions <- subset(Candida_instruments, chr == thisChr & bp > thisBpLow & bp < thisBpHigh)
   print(paste("Count is:", i))
   print(paste("Size of region for ",thisSNP," in ",thisChr,"_",thisBP, " : ", nrow(getRegions), sep="
   dir.create(paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/coloc_mrQTL_regions/",nameOfTe
    write.table(getRegions, paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/coloc_mrQTL_region
```

```
}
#(PD vs Candida)
##Bayesian colocalisation analysis
#######Excluding some columns and changing name of columns of region files#########
library("dplyr")
masterDir <- "/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/regionsV2"</pre>
files <- list.files(path=masterDir, pattern="*.tab", full.names = TRUE, recursive = FALSE)
for (i in files) {
 \#/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/regions/rs489567\_region.tab
 data <- fread(file=i,header=T,sep="\t")</pre>
 df <- select(data, `chr:pos`, rsID, eff.allele, ref.allele, `beta.x`, SE, N, Ncohorts, pDerived)
 names(df) <- c("SNP","rsID", "A1", "A2", "beta", "SE", "N", "Ncohorts","P-value")</pre>
 write.table(df, i, quote = F, sep = "\t", row.names = F)
}
#########reading one type of Dx sum stats and renaming col so that merging is possible###########
dataPD <- fread(file="/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/META5 all with rsid.txt", heade
names(dataPD)[17] <- "rsID"</pre>
write.table(dataPD, "/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/META5 all with rsid.tab", quote
#########reading one type of Dx sum stats but no change of col bc rsID column name already exists fo
nalls_PD <-fread("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/nallsEtAl2019_excluding23andMe_all
nalls_PD$SNP <- gsub("chr","",nalls_PD$SNP) #substituing a pattern of characters in a specific column i
########Merging all info on candida data with even more info from the Dx sum stats by rsID#######
masterDir <- "/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/regionsFullStats"
files <- list.files(path=masterDir, pattern="*.tab", full.names = TRUE, recursive = FALSE)
for (i in files) {
 print(paste("Working on...",i))
 data <- fread(file=i,header=T,sep="\t")</pre>
 print("here")
 df <- merge(data, dataPD, by="rsID")</pre>
 print("here")
 write.table(df, i, quote = F, sep = "\t", row.names = F)
}
##########Running the colocalization##########
```

```
#We subset data frames and rename them and let the function coloc.abf do all the work
SNPlist <- fread(file = paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/hitsBelow5E-5ForPD/hi
regions <- c(SNPlist$ID)
for (i in regions) {
 #/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/regions
 print(i)
 data <- fread(file = paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/regionsFullStats/",thi
 #print(names(data))
 df2_Candida_coloc <- data[,c("SNP","beta","P-value.x")]</pre>
 df1_PD_coloc <- data[,c("SNP","b","p","Freq1")]</pre>
 df2_Candida_coloc$var <- var(df2_Candida_coloc$beta)</pre>
 df1_PD_coloc$var <- var(df1_PD_coloc$b)</pre>
 names(df1_PD_coloc) <- c("snp","beta", "pvalues","MAF","varbeta")</pre>
 names(df2_Candida_coloc) <- c("snp","beta", "pvalues","varbeta")</pre>
 #Remove duplicates in dataset 1
 df1_PD_coloc <- df1_PD_coloc[!duplicated(df1_PD_coloc$snp), ]</pre>
 df2_Candida_coloc <- df2_Candida_coloc[!duplicated(df2_Candida_coloc$snp), ]</pre>
 df1_PD_coloc$type <- "cc"
 df2_Candida_coloc$type <- "quant"</pre>
 df1_PD_coloc$s <- 0.0590704464
 df2_Candida_coloc$sdY <- 1
 results <- coloc.abf(df1_PD_coloc, df2_Candida_coloc, MAF = NULL, p1 = 1e-02, p2 = 1e-02, p12 = 1e-02
 \#results \leftarrow coloc.abf(df1\_PD\_coloc, df2\_Candida\_coloc, MAF = NULL, p1 = 1e-04, p2 = 1e-04, p12 = 1e-04
 sink(file = paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/PD_coloc/",thisRegion,"_results
 print(results$summary)
 write.table(results$results, file = paste("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/PD_colo
#####POST COLOCALIZATION: FINDING COLOCALIZED REGIONS THAT EXPLAIN PHENOTYPE AND DX ETIOLOGIES VIA SIMI
masterDir <- "/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/PD coloc"
files <- list.files(path=masterDir, pattern="*Summary.txt", full.names = TRUE, recursive = FALSE)
for (i in files) {
 #print(paste("Working on...",i))
 data <- fread(file=i,header=T,sep=" ")</pre>
 #print(paste("The PP.H4.abf for this region is:",data$`PP.H4.abf`))
 print(as.double(data$PP.H4.abf))
 if (as.double(data$PP.H0.abf) > 0.95) {
   print("PP.HO.abf HIT")
 if (as.double(data$PP.H1.abf) > 0.95) {
   print("PP.H1.abf HIT")
 }
 if (as.double(data$PP.H2.abf) > 0.95) {
   print("PP.H2.abf HIT")
```

```
if (as.double(data$PP.H3.abf) > 0.95) {
   print("PP.H3.abf HIT")
}
if (as.double(data$PP.H4.abf) > 0.95) {
   print("PP.H4.abf HIT")
}
```

Generalised Summary-data-based Mendelian Randomisation

Analysis for each bacterial taxa hit

```
library("gsmr")
library("dplyr")
allo_sumstats <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/Alloprevo
cand_sumstats <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/Candidatu</pre>
meta5_pd <- fread("/Volumes/T7Touch/NIHSummer2021/Code/Colocolization/META5_all_with_rsid.tab", header=T
merge1 <- merge(allo_sumstats, meta5_pd, by="rsID")</pre>
hits_allo <- merge1[merge1$`P-value` < 0.0005, ]
hits_allo2 <- hits_allo[hits_allo$`p_bac` < 0.0005, ]
write.table(hits_allo2, file = "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/A
merge2 <- merge(cand_sumstats, meta5_pd, by="rsID")</pre>
hits_cand <- merge2[merge2$`P-value` < 0.0005, ]
hits_cand2 <- hits_cand[hits_cand$`p_bac` < 0.0005, ]</pre>
write.table(hits_cand2, file = "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/C
allo <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/Alloprevotella.id."
cand <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/CandidatusSoleafer.</pre>
allo_sub <- allo[,c("rsID", "eff.allele", "ref.allele", "beta.x", "SE", "p_bac", "N", "Effect", "StdErr
cand_sub <- cand[,c("rsID", "eff.allele", "ref.allele", "beta.x", "SE", "p_bac", "N", "Effect", "StdErr
allo_sub <- setNames(allo_sub, c("SNP","a1","a2","bzx","bzx_se","bzx_pval","bzx_n","bzy_","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bzy_se","bz
cand_sub <- setNames(cand_sub, c("SNP","a1","a2","bzx","bzx_se","bzx_pval","bzx_n","bzy_,"bzy_se","bzy_</pre>
write.table(allo_sub, file = "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/All
write.table(cand_sub, file = "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/ScoreFilesButMergedWSumstats/Cand
# Save the genetic variants and effect alleles in a text file using R
write.table(allo_sub[,c(1,2)], "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/allo_sub.allele", col.names=F,
write.table(cand_sub[,c(1,2)], "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/cand_sub.allele", col.names=F,
# Extract the genotype data from a GWAS dataset using GCTA
\#gcta64 --bfile gsmr\_example --extract gsmr\_example\_snps.allele --update-ref-allele gsmr\_example\_snps.a
#ALLOPREVOTELLA
```

```
bzx = allo_sub$std_bzx  # SNP effects on the risk factor
bzx_se = allo_sub$std_bzx_se # standard errors of bzx
bzx_pval = allo_sub$bzx_pval # p-values for bzx
bzy = allo_sub$bzy # SNP effects on the disease
bzy_se = allo_sub$bzy_se
                         # standard errors of bzy
bzy_pval = allo_sub$bzy_pval
                              # p-values for bzy
               # Sample size of the reference sample
n_ref = 7703
gwas thresh = 5e-8
                    # GWAS threshold to select SNPs as the instruments for the GSMR analysis
single_snp_heidi_thresh = 0.01  # p-value threshold for single-SNP-based HEIDI-outlier analysis
multi_snp_heidi_thresh = 0.01  # p-value threshold for multi-SNP-based HEIDI-outlier analysis
nsnps_thresh = 10  # the minimum number of instruments required for the GSMR analysis
heidi_outlier_flag = T  # flag for HEIDI-outlier analysis
ld_r2_thresh = 0.05  # LD r2 threshold to remove SNPs in high LD
ld_fdr_thresh = 0.05  # FDR threshold to remove the chance correlations between the SNP instruments
gsmr2_beta = 0 # 0 - the original HEIDI-outlier method; 1 - the new HEIDI-outlier method that is cu
gsmr_results = gsmr(bzx, bzx_se, bzx_pval, bzy, bzy_se, bzy_pval, ldrho, snp_coeff_id, n_ref, heidi_out
filtered_index=gsmr_results$used_index
cat("The estimated effect of the exposure on outcome: ",gsmr_results$bxy)
##Getting formatted summ stats
bac1 <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/bacSumStats_GCTA-COJO_format/Alloprevotella.id."
bac2 <- fread("/Volumes/T7Touch/NIHSummer2021/Code/GSMR/bacSumStats_GCTA-COJO_format/CandidatusSoleafer.
bac1 sub <- bac1[,c("rsID","eff.allele","ref.allele","beta.x","SE","p bac","N")]</pre>
bac2 sub <- bac2[,c("rsID","eff.allele","ref.allele","beta.x","SE","p bac","N")]</pre>
bac1_sub <- setNames(bac1_sub, c("SNP", "A1", "A2", "b", "se", "p", "N"))
bac2_sub <- setNames(bac2_sub, c("SNP", "A1", "A2", "b", "se", "p", "N"))
write.table(bac1_sub, "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/bacSumStats_GCTA-COJO_format/Alloprevot
write.table(bac2_sub, "/Volumes/T7Touch/NIHSummer2021/Code/GSMR/bacSumStats_GCTA-COJO_format/Candidatus
```

Merging SNPs for all bacterial taxa and disease

```
bacsMRQTLs_pdMRQTLs_mergeFDR <- function(bac_mrqtlfolder_paths,pd_mrqtl_root_path,results_rootpath) {
  for (i in bac mrqtlfolder paths){
    bac_mrqtls <- list.files(path=bac_mrqtlfolder_paths, pattern="*.msmr", full.names = TRUE, recursive
    nameBac <- str_split(str_split(i, "/", n = Inf, simplify = TRUE)[[7]], "_", n = Inf, simplify = TRU</pre>
    #print(paste("Working on",nameBac))
    dir.create(paste(results_rootpath,nameBac,"_PDmerged",sep=""))
    for (k in bac_mrqtls) { #going through each .msmr file
      #example name: /Volumes/T7Touch/NIHSummer2021/Data/MR-QTL_bacteria/SMR_results_CS/CS_chromatin_bl
      subMRQTL_name <- str_replace(str_split(k, "/", n = Inf, simplify = TRUE)[[8]], paste(nameBac,"_",</pre>
      pdMRQTL_filepath <- paste(pd_mrqtl_path_root_path, "PD_", subMRQTL_name, sep="") #so i dont need to
      #print(paste("MRQTL focused on:",pdMRQTL_filepath))
      \#possibleError \leftarrow tryCatch(fread(pdMRQTL_filepath, header=T, sep="\t"), error=function(e) {next})
      if (file.exists(pdMRQTL_filepath) == T) {
        #REAL WORK
        bac_mrqtl <- fread(k, header=T, sep="\t")</pre>
        pd_mrqtl <- fread(pdMRQTL_filepath, header=T, sep="\t") #now have both csvs opened
        #/Volumes/T7Touch/NIHSummer2021/Data/MR-QTL_bacteria/MR_QTL/PD_chromatin_blood_Bryois_SMR_allCh
        nameofMRQTL <- str_replace(paste(nameBac,"_",subMRQTL_name,sep=""),".msmr","")</pre>
        newdf <- merge(bac_mrqtl,pd_mrqtl,by="topSNP") #merging w/ mrqtl</pre>
        #new dir for just this mrqtl cuz will have p filtered and p unfiltered results
        dir.create(paste(results rootpath,nameBac," PDmerged","/",nameofMRQTL,sep=""))
        write.table(newdf,paste(results_rootpath,nameBac,"_PDmerged/",nameofMRQTL,"/","FDR_",nameofMRQT
        #do some p_SMR filtering
        \#newdf\$P\_fdr\_bac \leftarrow p.adjust(newdf\$`p\_SMR\_multi.x`, method = "fdr")
        \#newdf\$P_fdr_pd \leftarrow p.adjust(newdf\$^p_SMR_multti.y^n, method = "fdr")
        \#hits1 \leftarrow newdf[newdf$P_fdr_bac < 0.05, ]
        #hits2 <- hits1[hits1$P_fdr_pd < 0.05, ]
        #alpha <- 0.05
        #hits1 <- newdf[newdf$`p_SMR_multi.x` < alpha, ]</pre>
        #hits2 <- hits1[hits1$`p_SMR_multi.y` < alpha, ]</pre>
        print(paste(nameofMRQTL,"N Rows for MRQTL after p filtering:",nrow(hits2)))
        if (nrow(hits2) > 0) {
          print("########HIT#######")
        write.table(hits2,paste(results_rootpath,nameBac,"_PDmerged/",nameofMRQTL,"/",toString(alpha),"
   }
 }
bacsMRQTLs_pdMRQTLs_mergeFDR(bac_mrqtlfolder_paths,pd_mrqtl_root_path,results_rootpath)
```

Including Plots

You can also embed plots, for example:

Note that the echo = FALSE parameter was added to the code chunk to prevent printing of the R code that generated the plot.