Monte_carlo_pip

Cecilia Wang

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```
library(ggplot2)
library(phyloseq)
library(plyr)
library(vegan)
## Loading required package: permute
## Loading required package: lattice
## This is vegan 2.5-2
library("biom")
NZGL_taxonomy<-import_qiime_sample_data("~/PIP2018/primary_data/taxonomy.tsv")
  # the imported taxonomy data should have each sample as a row and each variable or taxonomy as a colu
Taxonomy_filter_file<-NZGL_taxonomy # make a copy
  #First make a plot of unfiltered taxonomy data, showing E coli abundance for each age group.
NZGL_taxonomy$time<-as.factor(NZGL_taxonomy$time) # to separate boxplot by different age category, type
### SEE PLOT 1: Supplemental 1: Abundance of E. coli x stratified by age in unfiltered data (chunk3)
# Then filter those samples out of all data, and use these data for every downstream analysis.
# select the useful part and find the interquartile range for E. coli, filter out samples that E. coli
Taxonomy_filter_file$E_coli<-NZGL_taxonomy$k__Bacteria.p__Proteobacteria.c__Gammaproteobacteria.o__Ente
# Split the dataset by time/age
E_coli_abundance_AtBirth<-subset(Taxonomy_filter_file, time==0)</pre>
E_coli_abundance_3_month<-subset(Taxonomy_filter_file, time==3)</pre>
E_coli_abundance_12_month<-subset(Taxonomy_filter_file, time==12)</pre>
E_coli_abundance_24_month<-subset(Taxonomy_filter_file, time==24)
# Calculate IQR by each time
E_coli_abundance_IOR_AtBirth<-IQR(E_coli_abundance_AtBirth$E_coli)
E_coli_abundance_IQR_3_month<-IQR(E_coli_abundance_3_month$E_coli)
E_coli_abundance_IQR_12_month<-IQR(E_coli_abundance_12_month$E_coli)
E_coli_abundance_IQR_24_month<-IQR(E_coli_abundance_24_month$E_coli)
# Filter the whole dataset at each time on E.coli > 1.5IQR. Except don't really.
Taxonomy_filtered_AtBirth<-subset(E_coli_abundance_AtBirth, E_coli<=100)
Taxonomy_filtered_3_month<-subset(E_coli_abundance_3_month,E_coli<=100)</pre>
Taxonomy_filtered_12_month<-subset(E_coli_abundance_12_month,E_coli<=100)
Taxonomy_filtered_24_month<-subset(E_coli_abundance_24_month,E_coli<=100)</pre>
Taxonomy_filtered<-rbind(Taxonomy_filtered_AtBirth, Taxonomy_filtered_3_month, Taxonomy_filtered_12_month
```

Plot4: Our power to detect effects

```
#Monte Carlo for BUG and MODULE beta diversity: C-section, time, antibiotics, treatment #bf
# source the functions for power test use Monte-carlo simulations
source("~/PIP2018/src/Monte_carlo_power_test.R")
## Loading required package: tcltk
Meta_pip<-as.data.frame(Taxonomy_filtered[,c(1:28)])</pre>
# NOTE: any category that tested here need to be factor. Therefore, transform every category as factors
Meta_pip[] <- lapply(Meta_pip, function(x){as.factor(x)})</pre>
# load metadata (pcl file does not have the info)
Metadata_pip<-read.csv("~/PIP2018/primary_data/metadata.csv")</pre>
# assign BF info to Meta pip and reset the subsets
Meta_pip$ageanyformula<-Metadata_pip$ageanyformula[match(Meta_pip$Studyid,Metadata_pip$Studyid)]
####=======Taxonomy test========####
## separate relative abundance data based on their associated metadata by time
t<-as.data.frame(t(Taxonomy_filtered[,c(-1:-28)]))
# subset metadata for further use of time stratefication
Meta_at_birth<-subset(Meta_pip,time==0)</pre>
Meta_3_month<-subset(Meta_pip, time ==3)</pre>
Meta_12_month<-subset(Meta_pip, time ==12)</pre>
Meta_24_month<-subset(Meta_pip, time ==24)</pre>
Monte_carlo_time<-Power_CI_figure(t,seq(5,50,5), Meta_pip, "time", 30, 10)
pd <- position_dodge(0.1)
time_tax<-ggplot(Monte_carlo_time,aes(Sampling_depth,mean)) + geom_errorbar(aes(ymin=CI_low, ymax=CI_hi
\#ggsave("\sim/pip-resubmit/results/unfiltered-time\_tax.pdf", plot = last\_plot(), device = \#NULL, path = NULL, 
\# scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
## caesar
  # figure out maximum sampling depth
summary(Meta_at_birth$caesar)
##
         0
                  1
## 123 53
summary(Meta_3_month$caesar)
##
                  1
## 107 48
summary(Meta_12_month$caesar)
##
          0
## 138 56
summary(Meta_24_month$caesar)
## 0 1
## 93 28
caeser_AB<-Power_CI_figure(t, seq(5,45,5), Meta_at_birth, "caesar", 30, 10 )</pre>
caeser_AB$age<-"0"
caeser_3month<-Power_CI_figure(t, seq(5,30,5), Meta_3_month, "caesar", 30, 10 )</pre>
caeser 3month$age<-"3"</pre>
```

```
caeser_12month<-Power_CI_figure(t, seq(5,45,5), Meta_12_month, "caesar", 30, 10 )</pre>
caeser_12month$age<-"12"
caeser_24month<-Power_CI_figure(t, seq(5,20,5), Meta_24_month, "caesar", 30, 10)
caeser_24month$age<-"24"
Caesar_all_time<-rbind(caeser_AB, caeser_3month, caeser_12month, caeser_24month)</pre>
Caesar_all_time$age<-as.factor(Caesar_all_time$age)</pre>
caesar_tax<-ggplot(Caesar_all_time,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(ymin=CI_lour))</pre>
\#ggsave("\sim/pip-resubmit/results/unfiltered-caesartax.pdf", plot = last\_plot(), device = \#NULL, path = Note = Not
# scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
## Antibiotics before 3 months
    # figure out maximum sampling depth
summary(Meta_at_birth$Antibiotics_before_6_months)
       0
               1
##
## 151 25
summary(Meta_3_month$Antibiotics_before_6_months)
##
         0
                1
## 135 20
summary(Meta_12_month$Antibiotics_before_6_months)
##
         0
                 1
## 168 26
summary(Meta_24_month$Antibiotics_before_6_months)
## 105 16
Anti6_AB<-Power_CI_figure(t,seq(5,20,5),Meta_at_birth, "Antibiotics_before_6_months", 30, 10)
Anti6 AB$age<-"0"
Anti6_3month<-Power_CI_figure(t,seq(5,15,5),Meta_3_month, "Antibiotics_before_6_months", 30, 10)
Anti6 3month$age<-"3"
Anti6_12month<-Power_CI_figure(t,seq(5,20,5),Meta_12_month, "Antibiotics_before_6_months", 30, 10)
Anti6_12month$age<-"12"
Anti6_24month<-Power_CI_figure(t,seq(5,10,5),Meta_24_month, "Antibiotics_before_6_months", 30, 10)
Anti6_24month$age<-"24"
Anti6_all_time<-rbind(Anti6_AB,Anti6_3month,Anti6_12month,Anti6_24month)
Anti6_all_time$age<-as.factor(Anti6_all_time$age)</pre>
abxtax<-ggplot(Anti6_all_time,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(ymin=CI_low, ym
\#ggsave("\sim/pip-resubmit/results/unfiltered-abx-tax.pdf", plot = last\_plot(), device = NULL, path = NULL
 \# scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
## Treatment/Studygroup
    # figure out maximum sampling depth
summary(Meta_at_birth$studygroup)
## bifido DR10 lactob DR20
                                                               placeb
```

```
##
            14
                        75
                                    87
summary(Meta_3_month$studygroup)
                                placeb
## bifido DR10 lactob DR20
##
                        70
                                    76
summary(Meta_12_month$studygroup)
## bifido DR10 lactob DR20
                                placeb
##
            19
                                    91
summary(Meta_24_month$studygroup)
## bifido DR10 lactob DR20
                                placeb
# bifido group has very limited number of samples
  # test with lactob and placeb groups
Studygroup_meta<-subset(Meta_pip, studygroup != "bifido DR10")</pre>
# reset the factor category studygroup
Studygroup_meta\studygroup<-as.factor(as.character(Studygroup_meta\studygroup))
SGmeta_AB<-subset(Studygroup_meta, time =="0")</pre>
SGmeta_3month<-subset(Studygroup_meta, time == "3")
SGmeta_12month<-subset(Studygroup_meta, time == "12")
SGmeta_24month<-subset(Studygroup_meta, time =="24")
SG_AB<-Power_CI_figure(t, seq(5,55,5), SGmeta_AB, "studygroup", 30, 10)
SG_AB$age<-"0"
SG_3month<-Power_CI_figure(t, seq(5,50,5), SGmeta_3month, "studygroup", 30, 10)
SG_3month$age<-"3"
SG_12month<-Power_CI_figure(t, seq(5,65,5), SGmeta_12month, "studygroup", 30, 10)
SG 12month$age<-"12"
SG_24month<-Power_CI_figure(t, seq(5,40,5), SGmeta_24month, "studygroup", 30, 10)
SG_24month$age<-"24"
SG_all_time<-rbind(SG_AB,SG_3month,SG_12month,SG_24month)
SG_all_time$age<-as.factor(as.character(SG_all_time$age))
tax_studygroup<-ggplot(SG_all_time,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(ymin=CI_lo
#ggsave("~/pip-resubmit/results/unfiltered-tax-studygroup.pdf", plot = last_plot(), device #= NULL, pat
# scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
####============####
# read the filtered modules file and modify it to fit the requirement for power test
Module <- import_qiime_sample_data("~/PIP/primary_data/modules.pcl")
Module<-as.data.frame(t(Module))</pre>
## Warning in class(X) <- NULL: Setting class(x) to NULL; result will no
## longer be an S4 object
Module_filtered<-Module[rownames(Module)%in%Taxonomy_filtered$Sample,]
t_modules<-Module_filtered[,c(-1:-27)] # remove metadata on top of the datset
t_modules<-as.data.frame(t(t_modules))</pre>
t_modules[] <- lapply(t_modules, function(x){as.numeric(as.character(x))})</pre>
t_modules[is.na(t_modules)]<-0 # replaced all the NA element in the dataset to 0
# Time test for modules
```

```
Monte_carlo_time_module<-Power_CI_figure(t_modules,seq(5,50,5), Meta_pip, "time", 30, 10)
pd <- position_dodge(0.1)</pre>
modtime<-ggplot(Monte_carlo_time_module,aes(Sampling_depth,mean)) + geom_errorbar(aes(ymin=CI_low, ymax
\#ggsave("\sim/pip-resubmit/results/unfiltered-mod-time.pdf", plot = last_plot(), device = NULL, path = NULL
 \# scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
## caesar
caeser_AB_module<-Power_CI_figure(t_modules, seq(5,45,5), Meta_at_birth, "caesar", 30, 10 )</pre>
caeser_AB_module$age<-"0"</pre>
caeser_3month_module<-Power_CI_figure(t_modules, seq(5,30,5), Meta_3_month, "caesar", 30, 10 )</pre>
caeser 3month module$age<-"3"</pre>
caeser_12month_module<-Power_CI_figure(t_modules, seq(5,45,5), Meta_12_month, "caesar", 30, 10 )</pre>
caeser_12month_module$age<-"12"</pre>
caeser_24month_module<-Power_CI_figure(t_modules, seq(5,20,5), Meta_24_month, "caesar", 30, 10)
caeser_24month_module$age<-"24"</pre>
Caesar_all_time_module<-rbind(caeser_AB_module, caeser_3month_module, caeser_12month_module, caeser_24mont
Caesar_all_time_module$age<-as.factor(Caesar_all_time_module$age)</pre>
mod_caesar<-ggplot(Caesar_all_time_module,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(yminestymean)) + geom
\#ggsave("\sim/pip-resubmit/results/unfiltered-mod-caesar.pdf", plot = last\_plot(), device = NULL, path = Null = Nul
  # scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
    #dpi = 300, limitsize = FALSE)
## Antibiotics before 6 months
Anti6_AB_module<-Power_CI_figure(t_modules,seq(5,20,5),Meta_at_birth, "Antibiotics_before_6_months", 30
Anti6_AB_module$age<-"0"</pre>
Anti6_3month_module<-Power_CI_figure(t_modules, seq(5,15,5), Meta_3_month, "Antibiotics_before_6_months",
Anti6_3month_module$age<-"3"</pre>
Anti6_12month_module<-Power_CI_figure(t_modules, seq(5,20,5), Meta_12_month, "Antibiotics_before_6_months
Anti6_12month_module$age<-"12"
Anti6_24month_module<-Power_CI_figure(t_modules, seq(5,10,5), Meta_24_month, "Antibiotics_before_6_months
Anti6_24month_module$age<-"24"</pre>
Anti6_all_time_module<-rbind(Anti6_AB_module,Anti6_3month_module,Anti6_12month_module,Anti6_24month_mod
Anti6_all_time_module$age<-as.factor(Anti6_all_time_module$age)</pre>
mod_anti<-ggplot(Anti6_all_time_module,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(ymin=C
\#ggsave("\sim/pip-resubmit/results/unfiltered-mod-anti.pdf", plot = last\_plot(), device = NULL, path = NULL
    \#scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
  # dpi = 300, limitsize = FALSE)
## Studygroup
SG AB module <- Power CI figure (t modules, seq (5,55,5), SGmeta AB, "studygroup", 30, 10)
SG_AB_module$age<-"0"
SG_3month_module<-Power_CI_figure(t_modules, seq(5,50,5), SGmeta_3month, "studygroup", 30, 10)
SG_3month_module$age<-"3"
SG_12month_module<-Power_CI_figure(t_modules, seq(5,65,5), SGmeta_12month, "studygroup", 30, 10)
SG_12month_module$age<-"12"
SG_24month_module<-Power_CI_figure(t_modules, seq(5,40,5), SGmeta_24month, "studygroup", 30, 10)
SG_24month_module$age<-"24"
```

```
SG_all_time_module<-rbind(SG_AB_module,SG_3month_module,SG_12month_module,SG_24month_module)
SG_all_time_module$age<-as.factor(as.character(SG_all_time_module$age))
mod_studygroup<-ggplot(SG_all_time_module,aes(Sampling_depth,mean, colour=age)) + geom_errorbar(aes(yminestudygroup) + geom_errorbar(aes(yminestudygroup)) + geom_errorbar(aes(yminestud
 \#ggsave("\sim/pip-resubmit/results/unfiltered-mod-studygroup.pdf", plot = last\_plot(), device \#= NULL, patients \#= NULL + 
 # scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
# dpi = 300, limitsize = FALSE)
library(ggpubr)
## Loading required package: magrittr
cfig<-ggarrange(caesar_tax, mod_caesar, labels = c("A", "B"), common.legend = TRUE, legend="bottom" )</pre>
ggsave("~/PIP2018/results/unfiltered-fig4.pdf", plot = last_plot(), device = NULL, path = NULL,
         scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
         dpi = 300, limitsize = FALSE)
supfig5<-ggarrange(time_tax, modtime, tax_studygroup, mod_studygroup, abxtax, mod_anti, labels=c("A", "
ggsave("~/PIP2018/results/unfiltered-supfig5.pdf", plot = last_plot(), device = NULL, path = NULL,
         scale = 1, width = 9, height = 6, units = c("in", "cm", "mm"),
         dpi = 300, limitsize = FALSE)
```