Ecological dioversity of human microbiome

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This script investigates ecological diversity in human microbiome.

Publicly available data of Human Microbiome Project are used in this exercise.

Reference: Lloyd-Price, J., Mahurkar, A., Rahnavard, G. et al. Strains, functions and dynamics in the expanded Human Microbiome Project. Nature 550, 61–66 (2017). https://doi.org/10.1038/nature23889

Load libraries

```
library(vegan)

## Loading required package: permute

## Loading required package: lattice

## This is vegan 2.5-7

library(ggplot2)
library(ape)
library(rmarkdown)
library(knitr)
```

Set working directory

Read data

Load sample info, a tab-delimited file

Load microbial species abundances, , a tab-delimited file

[1] 2355 1091

Calculate similarity between samples based on microbial species abundance

```
veg_dist <- as.matrix(vegdist(Microbial_Species, method="bray"))</pre>
```

Write the distance matrix in you computer as a tab-delimited file

Ecological diversity

Create a data frame to hold statistics

```
diversity_data <-
  setNames(data.frame(matrix(
    nrow = dim(Microbial_Species)[1],    ncol = 4,
  )), c("Sample", "Group", "Alpha_diversity", "Beta_diversity"))
rownames(diversity_data) <- rownames(Microbial_Species)</pre>
```

Alpha diversity (within sample diversity)

```
shannoon <- diversity(Microbial_Species, "shannon")

diversity_data$Alpha_diversity <- shannoon
diversity_data$Body_area <- sample_info[rownames(Microbial_Species), "Body_area"]</pre>
```

Kruska Wallis test

Test hypothesis: microbial alpha diversity is different among human body areas

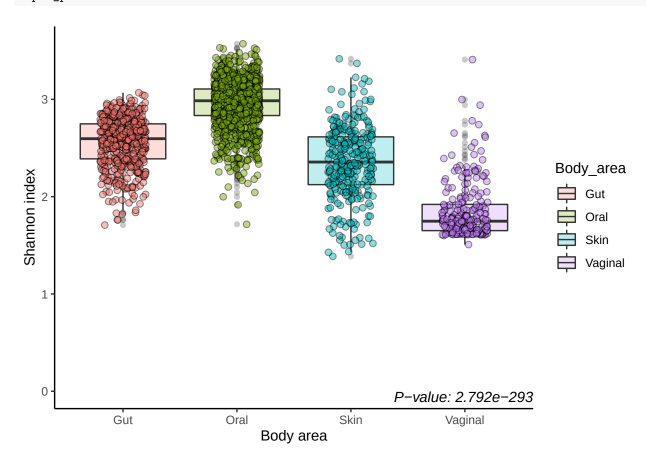
```
res_kruskal.test <- kruskal.test(diversity_data$Alpha_diversity, sample_info$Body_area)
pvalue <- res_kruskal.test$p.value</pre>
```

Visulization

ggplot2 for manuscript quality figures

```
alpha_plot <- ggplot2::ggplot(</pre>
  data = diversity_data, ggplot2::aes(factor(Body_area), y=Alpha_diversity)) +
  ggplot2::geom_boxplot(ggplot2::aes(fill = Body_area), alpha=0.25) +
  ggplot2::geom_point(ggplot2::aes(fill = Body_area), size=2, alpha=0.5,
                      shape = 21, colour = "black",
                      position = ggplot2::position_jitterdodge(),
                      show.legend = F) +
 ggplot2::annotate(
   geom = "text",
   x = Inf,
   y = 0,
   hjust = 1,
   vjust = 1,
   label = sprintf(
      "P-value: %1.3e",
     pvalue
   ) ,
   color = "black",
   size = 4,
   fontface = "italic"
 )+
  xlab("Body area") + ylab("Shannon index")+
  theme_classic()
```

alpha_plot



Save the plot with specifics

```
ggsave("analysis/alpha_diversity.pdf",alpha_plot, width = 5.6, height = 3.6,
    units = "in", dpi = 350 )
```