Analysis Illumina data

Heleen

8/6/2020

1. Pre blast

Load table and check read numbers

Functions:

```
getData <- function(){</pre>
  otu <- fread("~/Documents/derep_illum/changedheader/otu.about")</pre>
  before <- nrow(otu)
  # Remove X. or X from colnames
  names(otu) <- sub("#", "", names(otu))</pre>
  otu <- column_to_rownames(otu, var = "OTU ID")}</pre>
# optional
remove_chimeras <- function(){</pre>
  chimeras <- read.csv("~/Documents/IlluminaAdaptertrimmedAllreps/thingremoved", header=FALSE, sep=";")
  otu <- column_to_rownames(otu, var = "OTU.ID")</pre>
  otu <- otu[ ! sub("^.*?:", "", otu$OTU.ID) %in% chimeras$V1,] ##remove all chimeras
  after <- nrow(otu)
  cat(paste("removed", before-after, "chimeric sequences\n\n"))
  rownames(otu) <- NULL
  return(otu)}
# optional chimera removal: otu <- remove_chimeras()</pre>
# print results nicely:
myOTUcat <- function(){</pre>
  #total read sum in all clusters
  total_reads <- sum(rowSums(otu))</pre>
  cat(paste('total reads (grand total with which clustering was done):\n',
             total reads))
  cat("\n\n", 'Summary statistics of number of reads per OTU:\n')
  print(summary(rowSums(otu)))
  cat(paste("\n\n",
             'Total number of OTUs (including singletons):\n', nrow(otu)))
```

Execution:

```
otu <- getData()</pre>
myOTUcat()
## total reads (grand total with which clustering was done):
##
    2512237
##
##
    Summary statistics of number of reads per OTU:
##
       Min. 1st Qu.
                       Median
                                   Mean 3rd Qu.
##
       1.00
                1.00
                         1.00
                                   8.46
                                            2.00 86074.00
##
##
## Total number of OTUs (including singletons):
## 296884
```

Prepare Saba location data

Functions:

```
#function to change decimal to comma in one
decimal_to_comma <- function(data, column){</pre>
  data[,column] <- sub(",", ".",
                        data[,column],
                        fixed = TRUE)}
prepLocSaba <- function(){</pre>
  ## load the Saba sample location data
  locdata_saba <- read.delim("~/Downloads/NICO5-eDNA-64PE432-Metadata-MinIon - DataFilterSaba.txt")
  ## Change samplenames, colnames in metadatafile so they match the OTU file making merging is possible
  ## Change decimal to comma for computation.
  locdata_saba[,1] <- gsub("(?<![0-9])0+", "", locdata_saba[,1], perl = TRUE)
  locdata_saba[,1] <- gsub("\\.", "_", locdata_saba[,1], perl = TRUE)</pre>
  locdata_saba[,1] <- tolower(locdata_saba[,1])</pre>
  ## change long colnams to lat, long, altitude
  names(locdata_saba)[names(locdata_saba)=="geo_lat..in.decimalen..WGS84."] <- "lat"</pre>
  names(locdata_saba)[names(locdata_saba)=="geo_lon..in.decimalen..WGS84."] <- "long"</pre>
  names(locdata_saba)[names(locdata_saba)=="altitude..in.meters.aasl."] <- "altitude"</pre>
  names(locdata_saba)[1] <- "sample"</pre>
  ## change decimals to commas
  for (col in c("lat", "long")){
    locdata_saba[, col] <- as.numeric(decimal_to_comma(locdata_saba, col))}</pre>
  return(locdata_saba)}
```

Execution:

load and prepare Statia data

select only relevant columns and rows, and setnames, and change the numbers to depth

```
# take relevant columns, take out the samples that are not in the OTU table, and set the colnames to sa
locdata_statia <- read.delim("~/statia_location.txt") %>%
    select(Field.nr., lat, long, Average.depth) %>% # select relevant columns
    filter(!Field.nr. %in% c(528, 529)) %>% # discard irrelevant rows
    setNames(c("sample", "lat", "long", "altitude")) %>% # change column names
    mutate(altitude = as.numeric(gsub('[+]', '', altitude)) * -1) # mutate altidue column to negative
```

Bind Saba and statia data by row (to get merged data frame (mdf))

```
mdf <- plyr::rbind.fill(locdata_saba, locdata_statia)</pre>
```

estimate boxcore altitude data

boxcore altitude data is missing, so it's estimated by taking the nearest point geographically of which altitude data is available

```
#fill in missing boxcore altitude data wth nearestby latitude, the lowest value of that
mdf <- mdf %>%
  group_by(lat) %>%
  # arrange the groups by descending altitude within the groups
  arrange(desc(altitude), .by_group = TRUE) %>%
  # make new column with lowest altitude of group if the value is missing
  mutate(altitude = ifelse(is.na(altitude), min(altitude, na.rm = TRUE), altitude)) %>%
  # because for some boxcore samples it was taken at a slgihty different latitude, it does not belong t
  # Thus, R introduces infinite values which this command changes to NA values
  mutate(altitude = ifelse(is.infinite(altitude), NA, altitude)) %>%
  # needs to be ungrouped to fill it with the nearest & lowest altitude
  ungroup() %>%
  fill(altitude, .direction = 'down')
```

```
## Warning in min(altitude, na.rm = TRUE): no non-missing arguments to min;
## returning Inf
## Warning in min(altitude, na.rm = TRUE): no non-missing arguments to min;
## returning Inf
```

Put every sample in north, south or statia catogery based on latitude, to enable exchange testing. Add a tag indication what region the sampling location is in: Saba north, Saba south, or Statia.

Control reads

investigate control reads

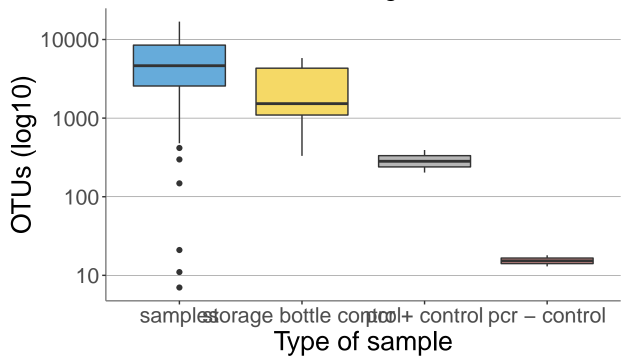
####Functions:

```
#prep data
controlDf <- function(){</pre>
  copy <- otu
  copy[copy>0] <- 1
  per_type <- copy %>% colSums() %>%
    as.data.frame() %>% rownames_to_column(var = "sample")%>%
    mutate(type = ifelse(sample %in% bottlecontrol, "storage bottle control",
                              ifelse(grepl("unicon", sample), "pcr + control",
                                     ifelse(sample %in% c("0", "neg_controle"),
                                             "pcr - control", "samples")))) %>%
    mutate(type = fct_reorder(type, desc(.)))
  return(per_type)}
plot_pertype <- function(df){</pre>
  control_plotOTU <-</pre>
    ggplot(df, aes(x = type,
                   y = .,
                   fill = type)) +
    geom_boxplot() +
    labs(title = "Number of OTUs per sample type",
         subtitle = "before abundance filterig",
       x = "Type of sample",
       y = "OTUs (log10)") +
    scale_fill_jco(alpha = 0.6) +
    scale_y_log10() +
  # edit lines and background
    theme(text = element text(size = 20),
        panel.grid.major.x = element_blank(),
        panel.grid.major.y = element_line("gray50", size = 0.2),
        panel.background = element_blank(),
        axis.line = element_line("gray50"),
```

```
legend.position = "none")
control_plotOTU}
```

Execution:

Number of OTUs per sample type before abundance filterig



anova of storage bottle control

```
res.aov <- aov(d = controlDf, . ~ type)
summary(res.aov)</pre>
```

Df Sum Sq Mean Sq F value Pr(>F)

```
3 1.763e+08 58764044
                                      3.444 0.0198 *
## type
## Residuals 96 1.638e+09 17060989
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
TukeyHSD (res.aov)
    Tukey multiple comparisons of means
##
##
      95% family-wise confidence level
##
## Fit: aov(formula = . ~ type, data = controlDf)
## $type
##
                                            diff
                                                       lwr
                                                                  upr
                                                                          p adj
## storage bottle control-samples
                                     -3137.205 -7376.563 1102.152 0.2204976
                                       -5496.348 -13218.159 2225.463 0.2517563
## pcr + control-samples
                                  -5778.848 -13500.659 1942.963 0.2117674
## pcr - control-samples
## pcr + control-storage bottle control -2359.143 -11018.102 6299.817 0.8919758
## pcr - control-storage bottle control -2641.643 -11300.602 6017.317 0.8553299
## pcr - control-pcr + control
                                       -282.500 -11082.120 10517.120 0.9998844
# check for assumptions
check_assumption <- function(){</pre>
 plot(res.aov, 1) # homogeneity of variances
 plot(res.aov, 2) # normality of residuals
 shapiro.test(residuals(res.aov))} # shapiro wilk of anova residuals
```

Post-blast

investigate storage bottle control identifications

```
`lca storage` <- read.delim("~/Documents/derep_illum/controls/underep/taxadded/lca") ## load lca file of
species <- table(`lca storage`$X.genus) %>%
    data.frame() %>%
    mutate(Var1 = ifelse(Freq < 1000, "Other", as.character(Var1))) %>%
    filter(!Var1=="no identification") %>%
    group_by(Var1) %>%
    dplyr::summarise(Freq = sum(Freq)) %>%
    mutate(Prop = (Freq/sum(Freq))*100) %>%
    ungroup() %>%
    mutate(Var1 = fct_reorder(Var1, desc(Freq))) %>%
    mutate(Var1 = fct_relevel(Var1, "Other", after = Inf))

## `summarise()` ungrouping output (override with `.groups` argument)

get_col <- function(){
    colorcount <- length(genuscount$Var1)
    qual_col <- brewer.pal.info[brewer.pal.info$category == "qual",]</pre>
```

```
col_vector <- unlist(mapply(brewer.pal,</pre>
                               qual_col$maxcolors, rownames(qual_col)))
  mycol <- sample(col_vector, colorcount)}</pre>
ggplot(species, aes(x = Var1, y = Freq, fill = Var1)) +
  geom_bar(stat = "identity", color = "black") +
  theme(text = element_text(size = 20),
        axis.text.x = element_text(angle = 45, hjust =1, size = 15),
        legend.position = "none") +
  scale_fill_jco() +
  labs(title = "Genera represented in storage bottles",
       x = "Genus \n",
       y = "Number of identifications (log10)") +
  scale_y_log10()
```

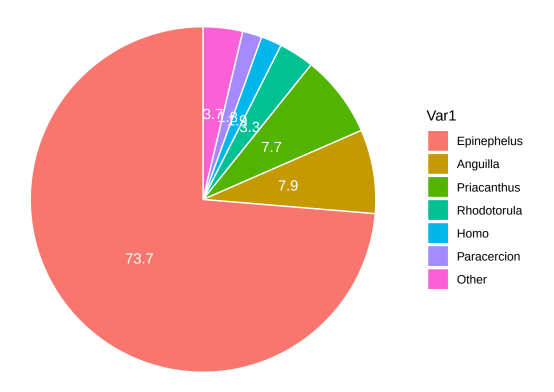
Number of identifications (log Genera represented in storage bottle 10000-1000-100-10-Homo

Epinephelus Anglilla Priacanthus Anodotorula

```
# Pie Chart
# add position of label
count.data <- species %>%
  arrange(desc(Var1)) %>%
  mutate(lab.ypos = cumsum(Prop) - 0.5*Prop)
ggplot(count.data, aes(x = "", y = Prop, fill = Var1)) +
     geom bar(width = 1, stat = "identity", color = "white") +
     coord_polar("y", start = 0)+
```

Genus

```
geom_text(aes(y = lab.ypos, label = round(Prop,1)), color = "white")+
theme_void()
```



Filter out out controls

If a OTU also contains control reads, these need to be filtered out of the samples contain them in frequencies that are close to the control frequencies. This could be contamination from the bottles the sample was stored in, or PCR contamination.

```
rate <- posContamination()

## Contamination percentage of positive control in other samples: 0.00671

negContamination()</pre>
```

Contamination percentage in negative samples: 0.0612

Low abundance filter

the rate of contamination in the positive control was used as low abundance filter rate.

```
lowAbuncanceFilter <- function(rate){</pre>
  before <- nrow(otu)</pre>
  colsum <- colSums(otu)</pre>
  min_read <- colsum * rate # if OTU contains less than this many reads, filter out
    mapply(col = otu, min = min_read, function(col, min){
    col[col < min] <- 0</pre>
    col}) %>%
    as.data.frame () %>%
    `rownames<-`(rownames(otu)) %>% filter(!rowSums(.[samples]) == 0) # take out "empty" otus
  after <- nrow(otu)
  percenage_ret <- ((before-after)/before)*100</pre>
  cat(paste("filtered out ", before-after, " OTUs, which is ",
            round(percenage_ret, 2), "% of original OTUs
            n'n',
            after, " OTUs were retained", sep = ""))
 return(otu)
}
controls <- c("sxm_2018_62", "sxm_2018_63", "sxm_2018_64",
              "sxm_2018_65", "sxm_2018_66", "sxm_2018_70",
              "sxm_2018_71", "0", "unicon1",
              "unicon1A", "neg_controle")
bottlecontrol <- c("sxm_2018_62", "sxm_2018_63", "sxm_2018_64",
              "sxm_2018_65", "sxm_2018_66", "sxm_2018_70",
              "sxm_2018_71")
samples <- names(otu) [-which(names(otu) %in% controls)]</pre>
otu <- lowAbuncanceFilter(rate = rate)</pre>
## filtered out 244221 OTUs, which is 82.26% of original OTUs
##
##
## 52663 OTUs were retained
```

remove singleton OTUs

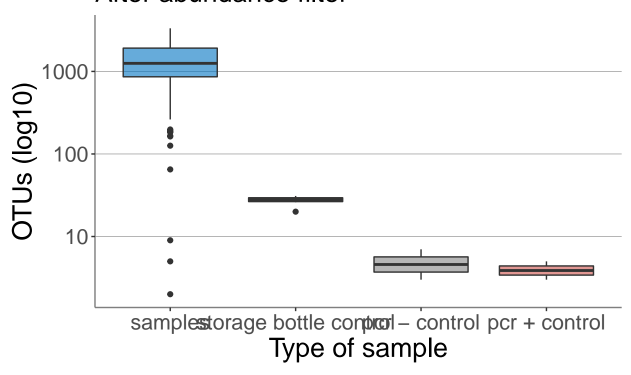
plot number of otus per sample

```
saba <- samples[grepl("sxm", samples)]</pre>
copy <- otu
copy[copy>0] <- 1
copy <- copy %>% colSums() %>%
  as.data.frame() %>% rownames_to_column(var = "sample")%>%
 mutate(type = ifelse(sample %in% bottlecontrol, "storage bottle control",
                             ifelse(grepl("unicon", sample), "pcr + control",
                                     ifelse(sample %in% c("0", "neg_controle"),
                                            "pcr - control", "samples")))) %>%
  mutate(type = fct_reorder(type, desc(.)))
control_plotOTU <-</pre>
  ggplot(copy, aes(x = type,
                            y = .,
                            fill = type)) +
  geom_boxplot() +
  labs(title = "Number of OTUs per sample type",
       subtitle = "After abundance filter",
       x = "Type of sample",
       y = "OTUs (log10)") +
  scale_fill_jco(alpha = 0.6) +
  scale_y_log10() +
  # edit lines and background
  theme(text = element text(size = 20),
        panel.grid.major.x = element_blank(),
        panel.grid.major.y = element_line("gray50", size = 0.2),
        panel.background = element_blank(),
        axis.line = element_line("gray50"),
        legend.position = "none")
```

control_plotOTU

- ## Warning: Transformation introduced infinite values in continuous y-axis
- ## Warning: Removed 1 rows containing non-finite values (stat_boxplot).

Number of OTUs per sample type After abundance filter



```
ggsave("controlplot AFTER abundance filter", plot = control_plotOTU, device = "png", height = 7, width
```

- ## Warning: Transformation introduced infinite values in continuous y-axis
- ## Warning: Removed 1 rows containing non-finite values (stat_boxplot).

Additional bottle control contamination check for saba samples

And remove controls from otu table

```
"sxm_2018_65", "sxm_2018_66", "sxm_2018_70",
              "sxm_2018_71")
samples <- names(otu)[-which(names(otu) %in% controls)]</pre>
saba <- samples[grepl("sxm", samples)]</pre>
# go over the rows(OTUs) where there are control reads and change any reads to 0 if they contain less t
filter_controls <- function(){</pre>
 OTUbefore <- nrow(otu)
 mcr <- do.call(pmax, otu[bottlecontrol]) # max control value for each otu
                                       # control values > 0
  mcp <- mcr > 0
  otu[mcp, saba][otu[mcp, saba] < 2*mcr[mcp]] <- 0
  # discard controls, and OTUs that have no reads associated bc of control filter
  otu <- otu %>%
    select(all_of(samples)) %>% #only keep samples
    filter(!rowSums(.) == 0)# discard OTUs that have no reads because of filtering
  ncolbefore <- ncol(otu)</pre>
  OTUafter <- nrow(otu)
  cat(paste("Control filtering removed", OTUbefore-OTUafter, "OTUs, which is ",
            round(((OTUbefore-OTUafter)/OTUbefore)*100, 2)), "%")
  otu <- otu[,colSums(otu) > 2000]
  ncolafter <- ncol(otu)</pre>
  after2000 <- nrow(otu)
  cat(paste("\n\nanother", OTUafter-after2000, "OTUs, were removed by removing ", ncolbefore-ncolafter,
  return(otu)}
otu <- filter controls()</pre>
## Control filtering removed 33 OTUs, which is 0.08 \%
## another 0 OTUs, were removed by removing 3 samples that head read numbers below 2000 reads
# select only samples that have read counts of higher than two thousand
```

write sequences to blast to file

now that all control reads and singletons have been filtered out, the remaining OTUs can be blasted. For this, the OTU centroid sequences from filtering step at 98% are extracted and then blasted -> taxadded -> dummyadded(for LCA script to work) -> lca script. Then its back to R

[1] 40376

3. LCA

remove bacteria hits

The entire dataset was blasted to genbank in galaxy. The resulting lca file is imported here to remova any reads belonging to bacteria.

```
genbank <- read.delim("~/Downloads/Galaxy6-[filtOTUseqs40376.fasta_BLAST_original_taxonomy_lca].tabular
bact <- genbank %>% filter(X.kingdom == "Bacteria")

toremove <- bact$X.Query
length(toremove)

## [1] 22428

otu <- otu[!row.names(otu) %in% toremove,]
nrow(otu)

## [1] 17948</pre>
```

import and prepare lca data

Then, the lca files at bitscore 8 and 12 percent are imported and compared

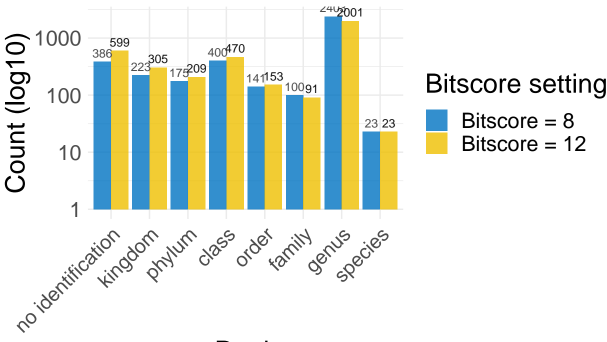
```
getLca <- function(){</pre>
  df <- read.delim("~/Documents/derep illum/changedheader/taxadded/bit8range")</pre>
  df2 <- read.delim("~/Documents/derep_illum/changedheader/taxadded/bit12range")
  dfs <- list(df, df2)
  # remove X. from cols and name the dfs
    lapply(dfs, function(x){setNames(x, sub("^X.", "", names(x)))}) %>%
    `names<-`(c("Bitscore = 8", "Bitscore = 12"))}</pre>
#execute the functions
lcas <- getLca()</pre>
lcas <- lapply(lcas, function(x) {x <- x[!x$Query %in% toremove,]})</pre>
# add information on how many reads were captured by the bitscore threshold
merged_dfs <-
 lapply(1:length(lcas), function(x) lcas[[x]] %>%
    data.frame) %>%
  # create extra column with what bitscore was used and bind the dataframes
 Map(cbind, ., Bitscore_setting = names(lcas)) %% # info of bitscore for bth dfs
  do.call(rbind, .) %>% #combined them by row
  data.frame() %>%
  filter(!grepl("sp\\.", species)) # remove hits that contain sp. because theyre not informative.
# get factor levels in right order for nice looking plots
merged_dfs$lca.rank <- factor(merged_dfs$lca.rank, levels = c("no identification", colnames(merged_dfs)
```

plot number of taxa found per rank by bitscore setting

Bitscore 8 has a stronger bias towards genus level identifications because the it takes fewer reads into account for the lca determination so higher chance that theres only 1 read to do taxa determination with,

```
myPlot <- function(){</pre>
  merged_dfs %>%
    ggplot(aes(x= lca.rank, y = ..count.., group = Bitscore_setting)) +
    geom_bar(aes(fill=`Bitscore_setting`),
             position = "dodge", alpha = 0.8) +
    geom_text(stat = "count",
              aes(label = ..count.., colour = Bitscore setting),
              position = position_dodge(0.9),
              vjust = -0.5, size = 3) +
    scale_fill_jco() +
    #scale_fill_brewer(type = "qual", palette = "Pastel2") +
    #scale_fill_manual(values = alpha(c("#00AFBB", "#FC4E07"), 0.8)) +
    theme minimal() +
    scale_color_manual(values = c("gray30", "gray8"),
                       guide = F) +
    scale_y_log10() +
    labs(title = "Number of LCA identifications per rank",
         subtitle = "by bitscore setting",
         x = "Rank",
         y = "Count (log10)",
         fill = "Bitscore setting") +
    theme(plot.title = element_text(size = 15, vjust = 1),
          text = element_text(size = 20),
          axis.text.x = element text(angle = 45, hjust =1, size = 15))}
p <- myPlot()</pre>
```

Number of LCA identifications per rank by bitscore setting



Rank

```
ggsave(filename = "Bitscore plot", p, device = "png", width = 10, height = 7.5)
```

The plot shows that bitscore 12 has less identifications but also less of a bias toward genus level, at it takes more reads into the LCA step.

Combine OTU and sampling location data

Compare the number of centroids supplied to the blast file to the number of blast hits found that had at least one blast hit of 70% identity and 70% coverage. The LCA file with bitscore 12 is chosen. get LCA: numbers

```
## number of rows in OTU file (number of OTUs found): 17948
##
## number of rows in LCA file (OTUs that had at least one blast hit): 3848
##
## Percentage of dark taxa is: 81.87 %
```

Combine sampling location data with the OTU table

Functions

```
get_bin_tags <- function(df){</pre>
    mdf %>%
    filter(sample %in% colnames(df)) %>%
    mutate(habitat = as.character(habitat)) %>%
    mutate(habitat = replace_na(habitat, "Mixed")) %>%
    mutate(bin = paste(tag, habitat)) %>%
    t() %>%
    as.data.frame() %>%
    row_to_names(1) %>%
    rownames_to_column(var = "Query") %>%
    filter(Query == "bin")
}
get_tags <- function(df){</pre>
  mdf %>%
    filter(sample %in% colnames(df)) %>%
    t() %>%
    as.data.frame() %>%
    row_to_names(1) %>%
    rownames_to_column(var = "Query") %>%
    filter(Query == "tag")}
replace_colnames <- function(df){</pre>
  df <- df %>% rownames_to_column(var = "Query")
  with_bins <- rbind.fill(tags, df) %>%
  row to names(row number = 1)
  colnames(with_bins)[1] <- "Query"</pre>
  return(with_bins)
}
# make a new df with one column calles Query (for merging), bind by col and remove rownames
#otu <- rownames_to_column(otu, var = "Query")
```

Execution:

```
tags <- get_bin_tags(otu)
for_network <- replace_colnames(otu)</pre>
```

Warning in row_to_names(., row_number = 1): Row 1 does not provide unique names.

```
## Consider running clean_names() after row_to_names().
```

```
tags <- get_tags(otu)
for_shared <- replace_colnames(otu)

## Warning in row_to_names(., row_number = 1): Row 1 does not provide unique names.
## Consider running clean_names() after row_to_names().</pre>
```

Prep data for sumarising per region

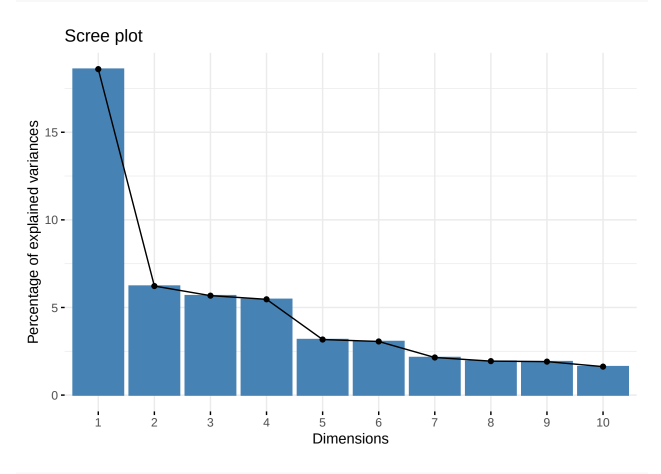
```
mixedmdf <- mdf %>%
  mutate(habitat = as.character(habitat)) %>%
  mutate(habitat = replace na(habitat, "Mixed")) %>%
  mutate(habitat = gsub(" layer", "", habitat)) %>%
  filter(!habitat == "100 m above bottom") %>%
  filter(!is.na(lat))
try <- otu %>%
  data.matrix() %>%
  t() %>%
  as.data.frame %>%
  rownames_to_column(var = "sample") %>%
  merge(mixedmdf, ., by = "sample", all.y = TRUE)
with row <- column to rownames(try, var = "sample")
with_row[,6:ncol(with_row)][with_row[,6:ncol(with_row)] > 0 ] <- 1</pre>
with_row <- with_row %>% filter(!is.na(lat))
with_row <- with_row[!rowSums(with_row[,6:ncol(with_row)])==0,]</pre>
res.pca <- prcomp(with_row[,6:ncol(with_row)],</pre>
                   center = TRUE, scale. = FALSE)
get_eig(res.pca)
```

```
##
           eigenvalue variance.percent cumulative.variance.percent
## Dim.1 1.235636e+02
                         1.860083e+01
                                                        18.60083
## Dim.2 4.133834e+01
                         6.222927e+00
                                                        24.82376
## Dim.3 3.769930e+01
                         5.675119e+00
                                                        30.49888
## Dim.4 3.631158e+01
                        5.466217e+00
                                                        35.96510
## Dim.5 2.106870e+01
                         3.171608e+00
                                                        39.13670
## Dim.6 2.033141e+01
                                                        42.19732
                         3.060618e+00
## Dim.7 1.426110e+01
                        2.146815e+00
                                                        44.34414
## Dim.8 1.287627e+01
                        1.938348e+00
                                                        46.28249
## Dim.9 1.269507e+01
                         1.911070e+00
                                                        48.19356
## Dim.10 1.079537e+01
                         1.625097e+00
                                                        49.81865
## Dim.11 1.029378e+01
                        1.549589e+00
                                                        51.36824
## Dim.12 9.961104e+00
                        1.499509e+00
                                                        52.86775
## Dim.13 9.367197e+00
                                                        54.27786
                        1.410105e+00
## Dim.14 9.019676e+00
                         1.357790e+00
                                                        55.63565
## Dim.15 8.318713e+00
                        1.252270e+00
                                                        56.88792
## Dim.16 8.105968e+00
                        1.220244e+00
                                                        58.10816
## Dim.17 8.056506e+00
                         1.212798e+00
                                                        59.32096
```

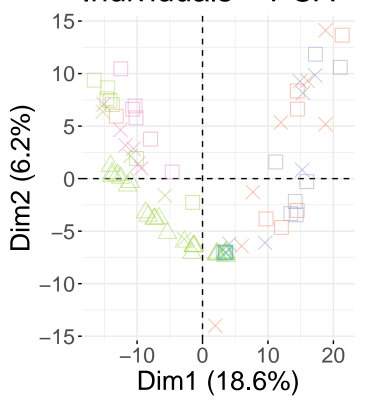
##	Dim 10	7.999300e+00	1.204186e+00	60.52514
		7.833405e+00	1.179213e+00	61.70436
			1.157416e+00	
		7.688611e+00		62.86177
		7.593018e+00	1.143026e+00	64.00480
		7.459436e+00	1.122917e+00	65.12772
		7.397463e+00	1.113588e+00	66.24130
		7.249856e+00	1.091368e+00	67.33267
		7.128307e+00	1.073070e+00	68.40574
		7.104494e+00	1.069485e+00	69.47523
		6.996115e+00	1.053170e+00	70.52840
##	Dim.28	6.936785e+00	1.044239e+00	71.57264
##	Dim.29	6.902948e+00	1.039145e+00	72.61178
##	Dim.30	6.660621e+00	1.002666e+00	73.61445
##	Dim.31	6.501130e+00	9.786571e-01	74.59310
##	Dim.32	6.407848e+00	9.646147e-01	75.55772
##	Dim.33	6.306231e+00	9.493177e-01	76.50704
##	Dim.34	6.256552e+00	9.418392e-01	77.44888
##	Dim.35	6.210241e+00	9.348676e-01	78.38374
##	Dim.36	6.112079e+00	9.200906e-01	79.30383
##	Dim.37	5.929396e+00	8.925902e-01	80.19642
##	Dim.38	5.763768e+00	8.676572e-01	81.06408
##	Dim.39	5.631673e+00	8.477721e-01	81.91185
##	Dim.40	5.450972e+00	8.205700e-01	82.73242
##	Dim.41	5.389881e+00	8.113736e-01	83.54380
##	Dim.42	5.323411e+00	8.013674e-01	84.34516
##	Dim.43	5.267317e+00	7.929232e-01	85.13809
##	Dim.44	5.056815e+00	7.612351e-01	85.89932
##	Dim.45	4.881622e+00	7.348621e-01	86.63418
##	Dim.46	4.844067e+00	7.292086e-01	87.36339
##	Dim.47	4.763941e+00	7.171468e-01	88.08054
##	Dim.48	4.637051e+00	6.980453e-01	88.77859
##	Dim.49	4.560209e+00	6.864777e-01	89.46506
##	Dim.50	4.440668e+00	6.684825e-01	90.13355
##	Dim.51	4.281621e+00	6.445401e-01	90.77809
		4.097152e+00	6.167707e-01	91.39486
			5.962724e-01	91.99113
		3.905791e+00	5.879640e-01	92.57909
		3.783217e+00	5.695120e-01	93.14860
		3.588145e+00	5.401466e-01	93.68875
		3.485210e+00	5.246511e-01	94.21340
		3.460860e+00	5.209855e-01	94.73439
		3.422107e+00	5.151519e-01	95.24954
		3.350430e+00	5.043619e-01	95.75390
		3.237250e+00	4.873241e-01	96.24123
		3.000120e+00	4.516274e-01	96.69285
		2.251840e+00	3.389840e-01	97.03184
		2.221142e+00	3.343628e-01	97.36620
		2.221142e+00 2.051549e+00	3.088329e-01	97.67503
		1.950424e+00	2.936099e-01	
				97.96864
		1.563194e+00	2.353176e-01	98.20396
		1.406802e+00	2.117750e-01	98.41574
		1.358941e+00	2.045701e-01	98.62031
		1.274400e+00	1.918436e-01	98.81215
##	1./1 חוע	1.101545e+00	1.658226e-01	98.97797

```
## Dim.72 1.001609e+00
                            1.507787e-01
                                                             99.12875
## Dim.73 8.549059e-01
                            1.286945e-01
                                                             99.25745
## Dim.74 8.429984e-01
                            1.269020e-01
                                                             99.38435
## Dim.75 7.747584e-01
                            1.166294e-01
                                                             99.50098
## Dim.76 7.136975e-01
                            1.074375e-01
                                                             99.60841
## Dim.77 6.076474e-01
                            9.147308e-02
                                                             99.69989
## Dim.78 5.186211e-01
                            7.807138e-02
                                                             99.77796
## Dim.79 4.197801e-01
                            6.319220e-02
                                                             99.84115
## Dim.80 3.859904e-01
                            5.810562e-02
                                                             99.89926
## Dim.81 3.474255e-01
                            5.230020e-02
                                                             99.95156
## Dim.82 3.218061e-01
                            4.844355e-02
                                                            100.00000
## Dim.83 2.292289e-25
                            3.450731e-26
                                                            100.00000
```

fviz_eig(res.pca)



Individuals - PCA



Region

- × Saba North
- Saba South
- △ Statia

Habitat

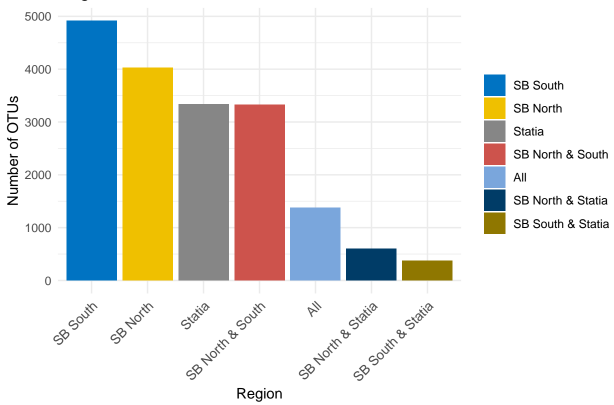
- 0,05 m above bottom
- 50 m above bottom
- 5 m above bottom
- Chlorofil max
- Mixed

```
ggsave(p, filename = "pcatest", device = "png", height = 7, width = 12)
```

```
# sum the values per row
summed <-
  data.matrix(for_shared[,-1]) %>% t(.) %>%
  rowsum(., group = sub("\\.\\d+$", "", rownames(.))) %>%
  t() %>%
  data.frame %>% `rownames<-`(for_shared[,1]) %>%
  rownames_to_column(var = "Query")
plotdf <- melt(summed, id.vars = "Query", value.name = "total_present")</pre>
# make dummy: O=not found & 1 = found
plotdf$dummy <- ifelse(plotdf$total_present > 0, yes=1, no=0)
plotdfcomb <- plotdf %>%
  group_by(variable) %>%
  mutate(presence = if else(dummy == 1,
                            ifelse(variable == 'Saba.North', 'SB North',
                                   ifelse(variable == 'Saba.South', 'SB South', 'Statia')),
                            NULL)) %>%
  na.omit() %>%
  group_by(Query) %>%
  summarise(presence = paste(presence, collapse = ' & '), .groups = 'drop') %>%
  mutate(shortpresence = ifelse(presence == 'SB North & SB South & Statia', 'All',
                                ifelse(presence == 'SB North & SB South', 'SB North & South',
```

```
presence)))
pres <-
  data.frame(table(plotdfcomb$shortpresence)) %>%
  setNames(c('Region', 'Freq')) %>%
  arrange(desc(Freq))
pres$Region<- factor(pres$Region, levels = reorder(pres$Region, -pres$Freq))</pre>
pres %>%
  ggplot() +
  geom_bar(aes(x = Region,
               y = Freq,
               fill = Region),
           stat = 'identity') +
  theme_minimal() +
  theme(axis.text.x = element_text(angle = 45, hjust = 1, size = '10'),
        legend.title = element_blank()) +
  labs(title = 'Regions where OTUs was detected',
       x = 'Region',
       y = 'Number of OTUs') +
  scale_fill_jco()
```

Regions where OTUs was detected



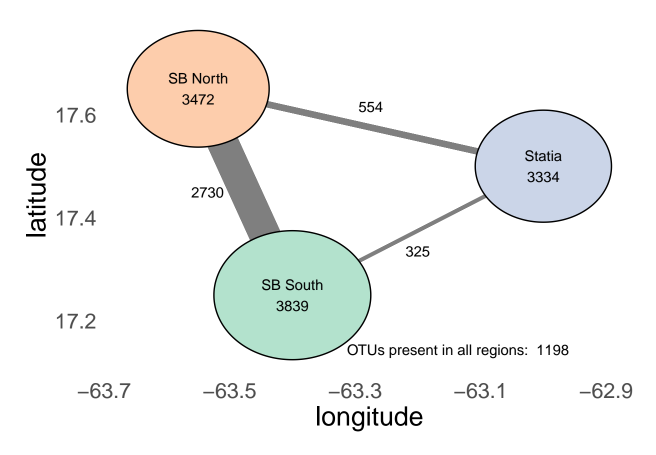
Region plot

```
lat = c(17.25, 17.65, 17.5, NA, NA, NA, NA)
long = c(-63.4, -63.55, -63.0, NA, NA, NA, NA)
lat_text <- c(NA, NA, NA, mean(c(17.25, 17.65)), mean(c(17.25, 17.65, 17.5)), mean(c(17.65, 17.5))+0.04
long_text <- c(NA, NA, NA, mean(c(-63.4, -63.55))-0.06, mean(c(-63.4, -63.55, -63.0)), mean(c(-63.55, -63.55))
coordinates <- cbind(lat, long, lat_text, long_text)</pre>
test <- cbind(pres, coordinates)</pre>
northlength <- length(grep("Saba North", colnames(for_shared)))</pre>
southlength <- length(grep("Saba South", colnames(for_shared)))</pre>
statialength <- length(grep("Statia", colnames(for_shared)))</pre>
test$Freq <- round((test$Freq/c(southlength, northlength, statialength,
                         mean(c(southlength, northlength)),
                         mean(c(southlength, northlength, statialength)),
                         mean(c(northlength, statialength)),
                         mean(c(southlength, statialength)))) * 25, 0)
allregions <- paste("OTUs present in all regions: ", test[5,2])
test <- test %>% filter(!Region == "All")
p <- ggplot(data=test, aes(x0=long, y0=lat, r=Freq/(8*max(Freq)), fill = Region)) +
  geom_segment(aes(x=long[1],
                   y=lat[1],
                   xend=long[2],
                   yend=lat[2]), size = test$Freq[4]/250, color = "gray50") +
  geom_segment(aes(x=long[2],
                   y=lat[2],
                   xend=long[3],
                   yend=lat[3]), size = test$Freq[5]/250, color = "gray50") +
  geom_segment(aes(x=long[3],
                   y=lat[3],
                   xend=long[1],
                   yend=lat[1]), size = test$Freq[6]/250, color = "gray50") +
  geom_circle() +
  geom_text(data = test, aes(x=long, y=lat, label = paste(Region, "\n", as.character(Freq), sep = "")))
  geom_text(data = test, aes(x =long_text, y=lat_text, label = Freq)) +
  theme_minimal() +
  labs(
    #title = "OTUs that are shared between regions",
    \#subtitle = "Circle correspond to number of OTUs found per region\setminusnthickness of line corresponds to
       x = "longitude",
       y = "latitude") +
  theme(text = element text(size = 20), panel.grid = element blank(),
        panel.grid.minor = element_blank(),
        legend.position = "none") +
  scale_fill_brewer(type = "qual", palette = "Pastel2") +
  annotate(geom = "text", label = allregions, x = Inf, y = -Inf, hjust = 1.3, vjust = -2)
```

Warning: Removed 3 rows containing non-finite values (stat_circle).

```
## Warning: Removed 3 rows containing missing values (geom_text).
```

Warning: Removed 3 rows containing missing values (geom_text).



```
ggsave("regions", p, device = "png")
```

```
## Saving 6.5 \times 4.5 in image
```

Warning: Removed 3 rows containing non-finite values (stat_circle).

Warning: Removed 3 rows containing missing values (geom_text).

Warning: Removed 3 rows containing missing values (geom_text).

prepare data for summarising per habitat

```
prep <- function(df){
   df %>%
   `rownames<-`(NULL) %>%
   column_to_rownames(var = "Query") %>%
   data.matrix() %>%
   t() %>%
```

```
as.data.frame() %>%
  rownames_to_column(var = "habitat") %>%
  filter(!habitat == "Saba.North.100.m.above.bottom") %>%
 mutate(habitat = sub("\\.\\d+$", "", habitat)) %>%
 filter(!habitat == "Saba.North.100.m.above.bottom")
aggr <- function(df){</pre>
  aggregate(df[,-1], list(habitat = df$habitat), mean) %>%
   mutate(habitat = habitat %>%
             sub("Saba.South", "SS", .) %>%
             sub("Saba.North", "SN", .) %>%
             sub(".above.bottom", "ab", .) %>%
             sub(".layer", "", .) %>%
             gsub("\\.", " ", .) %>%
             sub("0 05", "0.05", ., fixed = TRUE)) %>%
    column_to_rownames(var = "habitat") %>%
   t() %>%
   as.data.frame %>%
   rownames_to_column(var = "habitat")
}
```

Execution:

```
try <- prep(for_network)

try[,-1][try[,-1]>0] <- 1

agr <- try %>% aggr()
```

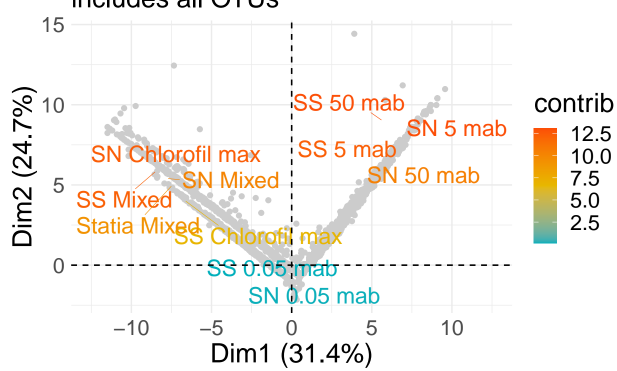
```
myPca <- function(df){</pre>
  res.pca <- prcomp(column_to_rownames(df, var = "habitat"),</pre>
                     center = TRUE, scale. = TRUE)
  print(get_eig(res.pca))
  fviz_eig(res.pca)
  fviz_pca_biplot(res.pca,
                  label = "var",
                   col.var = "contrib",
                   gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
                   col.ind = "gray80",
                   repel = TRUE,
                   geom.ind = "point",
                   geom.var = "text",
                   labelsize = 6) + theme_minimal() + theme(text = element_text(size = 20))
}
p_agr <- myPca(agr) +</pre>
  labs(title = "Presence-absence per bin",
       subtitle = "includes all OTUs") + xlim(-12.5, 12.5)
```

```
## Dim.1 3.4579667 31.436061 31.43606
## Dim.2 2.7160396 24.691269 56.12733
```

```
## Dim.3
           0.9972303
                              9.065730
                                                           65.19306
## Dim.4
           0.9955104
                              9.050094
                                                           74.24315
## Dim.5
           0.7590921
                              6.900837
                                                           81.14399
           0.5116063
                                                           85.79496
## Dim.6
                              4.650966
## Dim.7
           0.4595809
                              4.178008
                                                           89.97297
## Dim.8
           0.3532729
                              3.211572
                                                           93.18454
## Dim.9
           0.2794858
                              2.540780
                                                           95.72532
           0.2540913
                              2.309921
                                                           98.03524
## Dim.10
## Dim.11
           0.2161238
                              1.964762
                                                           100.00000
```

p_agr

Presence—absence per bin includes all OTUs



ggsave("finalpcapresence", p_agr, device = "png")

Saving 6.5×4.5 in image