

ITS of Wheat

2023-08-23

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The relationship between fungi and wheat encompasses significant importance in agricultural ecosystems, transcending beyond the realm of infections. The coexistence of these two entities profoundly impacts agricultural productivity, with the specter of wheat root rot disease, wrought by soil-borne fungal pathogens, casting a formidable shadow over global crop yields, resulting in staggering annual economic losses. Beyond the purview of pathogenic interactions, the symbiotic and commensalistic associations of fungi with wheat rhizosphere foster a complex network of relationships that shape plant health and ecosystems. Investigating the multifaceted nexus between rhizosphere soil fungal diversity and wheat roots unveils a nuanced panorama, pivotal for comprehending the genesis and progression of wheat root rot disease.

Qiime2 to Phyloseq

Converting QIIME2 outcomes to the phyloseq object structure in R enables comprehensive analysis and visualisation of microbial community profiles. The phyloseq package provides functions for organising and manipulating data within the phyloseq object, allowing for diverse analyses such as diversity assessments, differential abundance testing, and taxonomic profile visualisation. By leveraging R's capabilities, researchers can perform advanced statistical analysis, integrate with other omics data, and gain deeper insights into microbiome datasets.

```
# Download qiime2R from Github
# if (!requireNamespace("devtools", quietly = TRUE)){install.packages("devtools")}
# devtools::install_github("jbisanz/qiime2R")
library("qiime2R")

# Download phyloseq from CRAN
# if (!require("BiocManager", quietly = TRUE))
#   install.packages("BiocManager")
# BiocManager::install("phyloseq")
library("phyloseq")

library("tidyverse")

# Convert qiime2 to phyloseq format
physeq.a <- qza_to_phyloseq(
  features = "ITS/table-its-fungi-with-phyla-no-mitochondria-no-chloroplast.qza", # table.qza
  tree = "ITS/phylogeny-align-to-tree-mafft-fasttree/rooted_tree.qza",
  taxonomy = "ITS/taxonomy-fungi.qza",
  metadata = "ITS/meta-table.txt"
)
physeq.a ## confirm the object

## phyloseq-class experiment-level object
```

```
## otu_table() OTU Table: [ 3904 taxa and 37 samples ]
## sample_data() Sample Data: [ 37 samples by 7 sample variables ]
## tax_table() Taxonomy Table: [ 3904 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 3904 tips and 3878 internal nodes ]
```

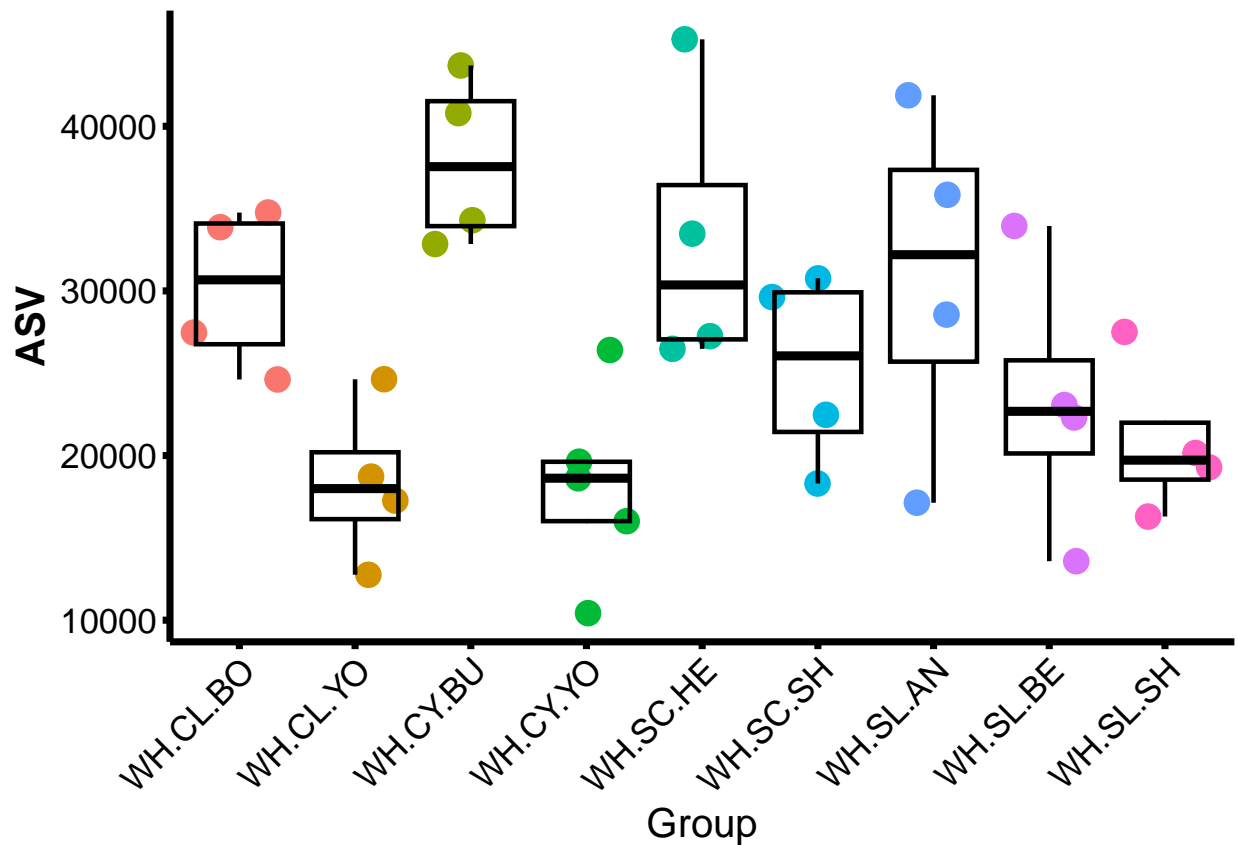
Assessing ASV Distribution (merged and filtered)

We evaluated the number of Amplicon Sequence Variants (ASVs) after merging and filtering, as well as visualised the raw read counts per sample within each group. This analysis provided insights into the distribution of ASVs and the sequencing depth of each sample, allowing for a comprehensive assessment of data quality and quantity.

```
# install.packages("ggplot2")
library("ggplot2")
```

```
ASV <- sample_sums(physeq.a)
ASV <- as.data.frame(ASV)
ASV$Group <- physeq.a@sam_data$Group

ggplot(ASV, aes(x = Group, y = ASV, colour = Group))+
  geom_point(alpha = 1, position = "jitter", size = 4) +
  geom_boxplot(alpha = 0, colour = "black", size = 0.8)+
  theme_classic() +
  theme(text = element_text(size=15, colour = "black"),
        axis.ticks = element_line(colour = "black", size = 1.25),
        axis.line = element_line(colour = 'black', size = 1.25),
        axis.text.x = element_text(angle=45, hjust=1, colour = "black", size = 13),
        axis.text.y = element_text(angle=0, hjust=0.5, colour = "black",size = 13),
        axis.title.y = element_text(color="black", size=15,face="bold"),
        legend.position = "none")
```



Beta diversity

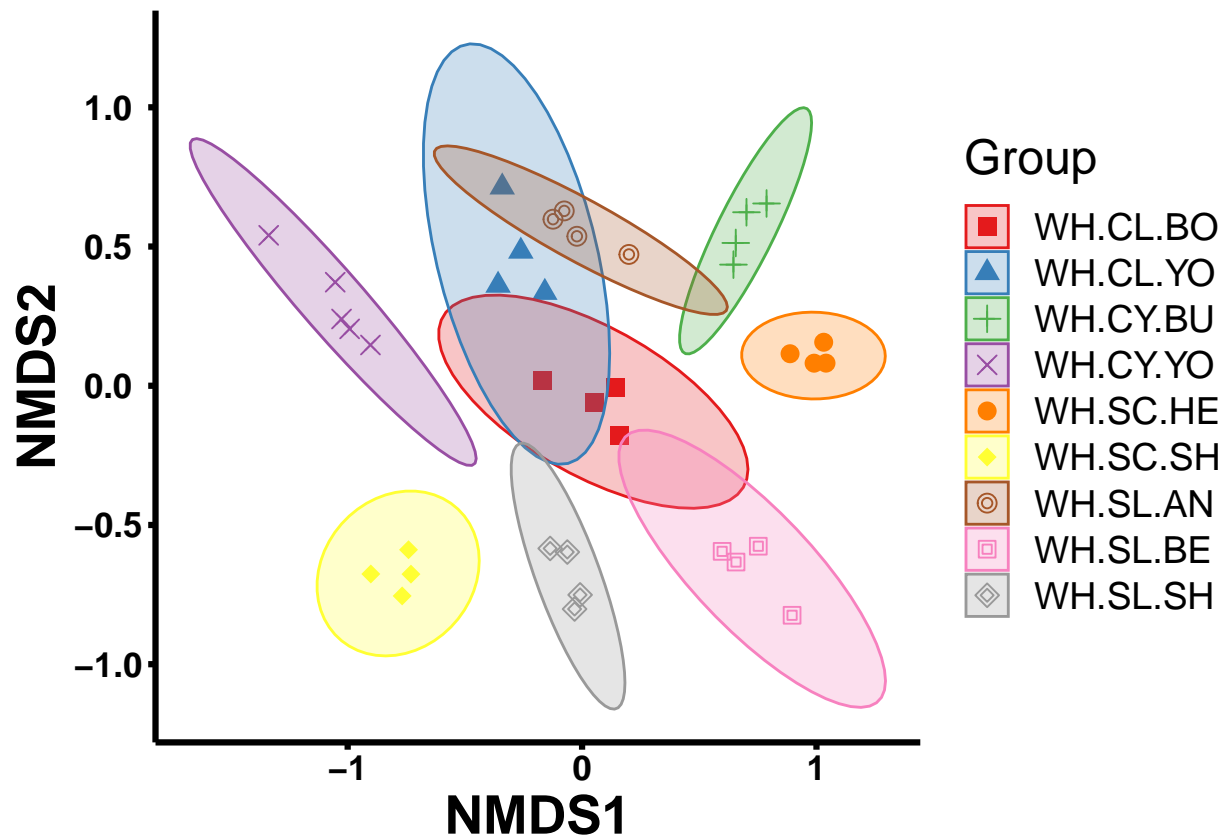
Beta diversity is a metric used in microbial community studies to assess species composition dissimilarity between samples, providing insights into microbial community dynamics and their ecological significance. By quantifying variation in community structure and employing visualisation techniques, beta diversity analysis allows comparisons across habitats or treatments, revealing key drivers of community variation and functional implications. It aids our understanding of microbial assemblages and their responses to environmental changes in ecology, environmental science, and human health research.

```
# method options: NMDS / PCoA
NMDS <- ordinate(physeq = physeq.a, method = "NMDS", distance = "bray")
```

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.196196
## Run 1 stress 0.1929835
## ... New best solution
## ... Procrustes: rmse 0.1085372 max resid 0.2861605
## Run 2 stress 0.23127
## Run 3 stress 0.2481474
## Run 4 stress 0.196196
## Run 5 stress 0.2110517
## Run 6 stress 0.2488192
## Run 7 stress 0.2362618
## Run 8 stress 0.2375049
```

```
## Run 9 stress 0.2110517
## Run 10 stress 0.2096686
## Run 11 stress 0.1929835
## ... New best solution
## ... Procrustes: rmse 1.120451e-06  max resid 2.738374e-06
## ... Similar to previous best
## Run 12 stress 0.2540123
## Run 13 stress 0.2188336
## Run 14 stress 0.2096244
## Run 15 stress 0.2110517
## Run 16 stress 0.196196
## Run 17 stress 0.2096178
## Run 18 stress 0.196196
## Run 19 stress 0.2357275
## Run 20 stress 0.196196
## *** Best solution repeated 1 times
```

```
# Plot ordination
plot_ordination(
  physeq = physeq.a,
  ordination = NMDS,
  color = "Group",
  shape = "Group"
) +
  theme_classic() +
  geom_point(aes(color = Group), alpha = 1, size = 3) +
  theme(
    text = element_text(size = 18, colour = "black"),
    axis.ticks = element_line(colour = "black", size = 1.1),
    axis.line = element_line(colour = "black", size = 1.1),
    axis.text.x = element_text(colour = "black", angle = 0, hjust = 0.5, size = 13, face = "bold"),
    axis.text.y = element_text(colour = "black", angle = 0, hjust = 0.5, size = 13, face = "bold"),
    axis.title.y = element_text(color = "black", size = 20, face = "bold"),
    axis.title.x = element_text(color = "black", size = 20, face = "bold")
  ) + stat_ellipse(geom = "polygon", type="norm", alpha=0.25, aes(fill = Group)) +
  scale_color_brewer(palette = "Set1") +
  scale_fill_brewer(palette = "Set1") +
  scale_shape_manual(values = c(15, 17, 3, 4, 16, 18, 21, 22, 23)) # Set custom shapes
```



```
# Clean up by removing objects that are no longer needed
rm(pcoa, physeq.W.B)
```

Alpha diversity

Alpha diversity is a fundamental concept in ecology and refers to the diversity or richness of species within a specific community or habitat. In the context of microbial ecology, alpha diversity represents the diversity of microorganisms within a given sample or microbiome. It provides insights into the variety and evenness of microbial species present in a particular environment. Common measures of alpha diversity include species richness, which counts the number of unique species, and evenness, which assesses the distribution of species abundances. Alpha diversity is crucial for understanding the stability, resilience, and functional potential of microbial communities. It can be influenced by various factors, including environmental conditions, host factors, and perturbations. By comparing alpha diversity across different samples or experimental groups, researchers can gain insights into the impact of factors such as disease, habitat changes, or interventions on microbial community structure.

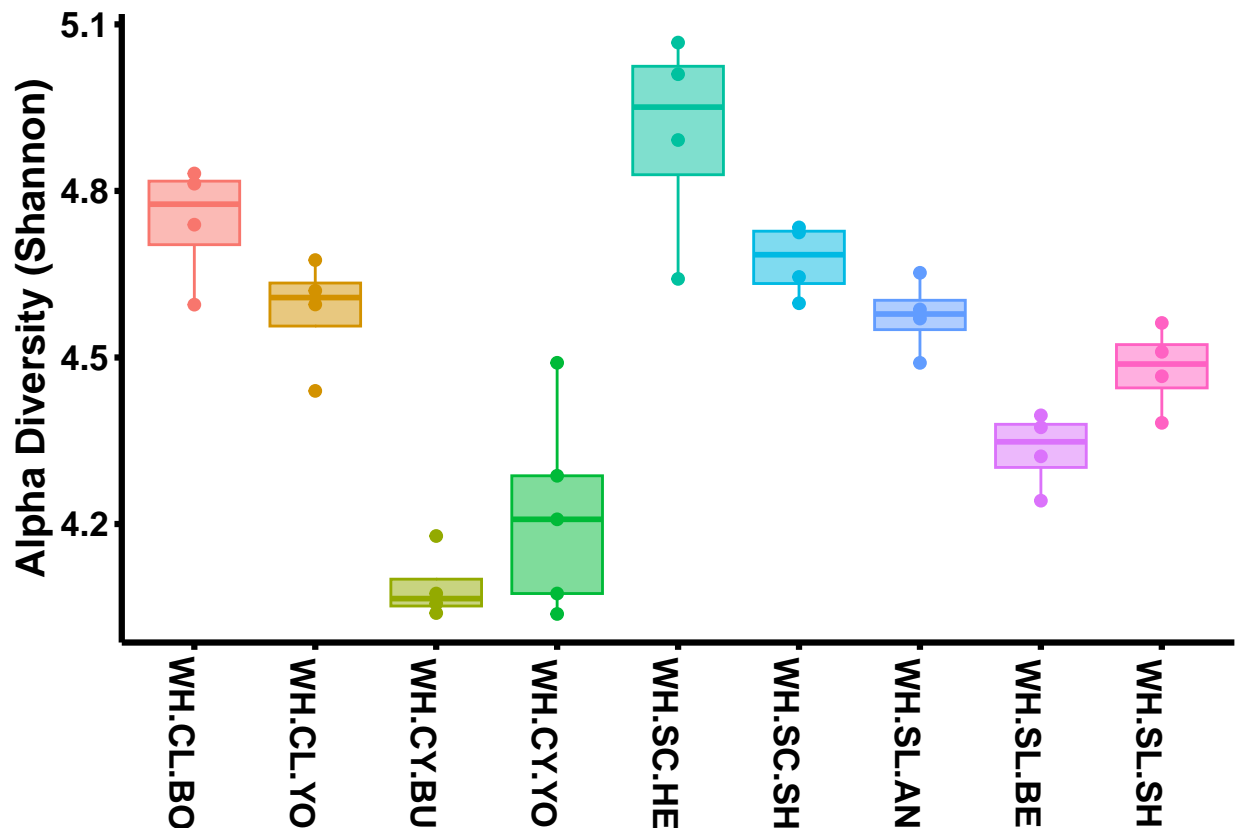
```
# available measurements [c("Observed", "Chao1", "ACE", "Shannon", "Simpson", "InvSimpson", "Fisher")]
alpha.object <- cbind(
  x = sample_data(physeq.a),
  y = estimate_richness(physeq.a, measures = 'Shannon')
)

# Create a plot for alpha diversity
ggplot(data = alpha.object, aes(x = x.Group, y = Shannon, color = x.Group, fill = x.Group)) +
  theme_classic() +
```

```

labs(
  x = element_blank(),           # No x-axis label
  y = "Alpha Diversity (Shannon)" # y-axis label
) +
geom_point(size = 1.75) +        # Add points
geom_boxplot(alpha = 0.5) +      # Add boxplot with transparency
theme(
  text = element_text(size = 18, colour = "black"),
  axis.ticks = element_line(colour = "black", size = 1.1),
  axis.line = element_line(colour = "black", size = 1.1),
  axis.text.x = element_text(colour = "black", angle = 270, size = 13, face = "bold"),
  axis.text.y = element_text(angle = 0, hjust = 0, colour = "black", size = 13, face = "bold"),
  axis.title.y = element_text(color = "black", size = 15, face = "bold"),
  legend.position = "none"       # Hide legend
)

```



```

# Clean up by removing the alpha.object
rm(alpha.object)

```

Pairwise comparison using PERMANOVA

Pairwise PERMANOVA is a statistical method used to compare multiple groups or treatments in ecological and microbial community studies. It assesses dissimilarity between samples and provides a p-value to determine the significance of observed differences. This approach is valuable for targeted group comparisons,

allowing researchers to investigate the effects of specific factors on microbial communities and uncover significant variations in community composition. By considering within- and between-group variation, pairwise PERMANOVA provides robust statistical analysis and insights into microbial community dynamics and functioning.

```
# devtools::install_github("pmartinezarbizu/pairwiseAdonis/pairwiseAdonis")
library("pairwiseAdonis")
```

```
metdat = as.data.frame(as.matrix(physeq.a@sam_data))
dat = as.data.frame(t(as.data.frame(physeq.a@otu_table)))
pairwise.adonis(dat, metdat$Group, sim.function = "vegdist",
  sim.method = "bray", p.adjust.m = "bonferroni",
  reduce = NULL, perm = 100000)
```

##		pairs	Df	SumsOfSqs	F.Model	R2	p.value	p.adjusted
## 1	WH.CL.BO vs WH.CL.YO	1	0.6836316	7.141412	0.5434281	0.028571429	1.0000000	
## 2	WH.CL.BO vs WH.CY.BU	1	0.8424259	11.626774	0.6596087	0.028571429	1.0000000	
## 3	WH.CL.BO vs WH.CY.YO	1	0.9402290	8.830922	0.5578274	0.008089919	0.2912371	
## 4	WH.CL.BO vs WH.SC.HE	1	0.7165156	7.172596	0.5445089	0.028571429	1.0000000	
## 5	WH.CL.BO vs WH.SC.SH	1	0.7061539	5.851479	0.4937341	0.028571429	1.0000000	
## 6	WH.CL.BO vs WH.SL.AN	1	0.5239285	4.907670	0.4499284	0.028571429	1.0000000	
## 7	WH.CL.BO vs WH.SL.BE	1	0.6106859	4.633587	0.4357501	0.028571429	1.0000000	
## 8	WH.CL.BO vs WH.SL.SH	1	0.6328513	6.156368	0.5064315	0.028571429	1.0000000	
## 9	WH.CL.YO vs WH.CY.BU	1	1.0094973	16.080161	0.7282629	0.028571429	1.0000000	
## 10	WH.CL.YO vs WH.CY.YO	1	0.7288589	7.424016	0.5146983	0.008119919	0.2923171	
## 11	WH.CL.YO vs WH.SC.HE	1	0.8085318	8.961816	0.5989792	0.028571429	1.0000000	
## 12	WH.CL.YO vs WH.SC.SH	1	0.7401712	6.668033	0.5263669	0.028571429	1.0000000	
## 13	WH.CL.YO vs WH.SL.AN	1	0.5871301	6.047873	0.5019868	0.028571429	1.0000000	
## 14	WH.CL.YO vs WH.SL.BE	1	0.7148304	5.853562	0.4938230	0.028571429	1.0000000	
## 15	WH.CL.YO vs WH.SL.SH	1	0.7150048	7.678351	0.5613506	0.028571429	1.0000000	
## 16	WH.CY.BU vs WH.CY.YO	1	1.2795885	16.357105	0.7003053	0.008139919	0.2930371	
## 17	WH.CY.BU vs WH.SC.HE	1	0.6135266	9.164290	0.6043336	0.028571429	1.0000000	
## 18	WH.CY.BU vs WH.SC.SH	1	0.9171570	10.454216	0.6353518	0.028571429	1.0000000	
## 19	WH.CY.BU vs WH.SL.AN	1	0.7116171	9.641420	0.6164031	0.028571429	1.0000000	
## 20	WH.CY.BU vs WH.SL.BE	1	0.8752828	8.854946	0.5960941	0.028571429	1.0000000	
## 21	WH.CY.BU vs WH.SL.SH	1	0.9093068	13.018466	0.6845171	0.028571429	1.0000000	
## 22	WH.CY.YO vs WH.SC.HE	1	1.2661487	12.443868	0.6399893	0.007559924	0.2721573	
## 23	WH.CY.YO vs WH.SC.SH	1	0.8514764	7.121568	0.5043043	0.007979920	0.2872771	
## 24	WH.CY.YO vs WH.SL.AN	1	0.8690449	8.074413	0.5356370	0.008039920	0.2894371	
## 25	WH.CY.YO vs WH.SL.BE	1	0.9576115	7.418112	0.5144995	0.007559924	0.2721573	
## 26	WH.CY.YO vs WH.SL.SH	1	0.9647977	9.256035	0.5693907	0.008349917	0.3005970	
## 27	WH.SC.HE vs WH.SC.SH	1	0.8233765	7.149139	0.5436964	0.028571429	1.0000000	
## 28	WH.SC.HE vs WH.SL.AN	1	0.6725153	6.642198	0.5253990	0.028571429	1.0000000	
## 29	WH.SC.HE vs WH.SL.BE	1	0.5757410	4.558976	0.4317631	0.028571429	1.0000000	
## 30	WH.SC.HE vs WH.SL.SH	1	0.7799919	8.017346	0.5719589	0.028571429	1.0000000	
## 31	WH.SC.SH vs WH.SL.AN	1	0.6679211	5.473318	0.4770475	0.028571429	1.0000000	
## 32	WH.SC.SH vs WH.SL.BE	1	0.6649246	4.521124	0.4297187	0.028571429	1.0000000	
## 33	WH.SC.SH vs WH.SL.SH	1	0.6153668	5.211822	0.4648506	0.028571429	1.0000000	
## 34	WH.SL.AN vs WH.SL.BE	1	0.6718066	5.045557	0.4567952	0.028571429	1.0000000	
## 35	WH.SL.AN vs WH.SL.SH	1	0.8360346	8.027306	0.5722628	0.028571429	1.0000000	
## 36	WH.SL.BE vs WH.SL.SH	1	0.6155645	4.764900	0.4426330	0.028571429	1.0000000	
##	sig							
## 1								

```
## 2
## 3
## 4
## 5
## 6
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## 9
## 10
## 11
## 12
## 13
## 14
## 15
## 16
## 17
## 18
## 19
## 20
## 21
## 22
## 23
## 24
## 25
## 26
## 27
## 28
## 29
## 30
## 31
## 32
## 33
## 34
## 35
## 36
```

Bar plot - relative abundance

Bar plots illustrating relative abundance in microbiome enable to visually represent the proportions of different microbial taxa within a given sample. Each bar corresponds to a specific taxonomic group, with its height indicating the relative abundance of that group in the microbiome.

```
# install.packages("plyr")
library("plyr")
```

```
# https://github.com/joey711/phyloseq/issues/901
```

```
## (3A) Merge the replicate samples for each Group
physeq.a.group = merge_samples(physeq.a, "Group") # Sum between replicate samples
```

```
# (3B) repair factors in the sample metadata
```

```
sample_data(physeq.a.group)$Group <- levels(sample_data(physeq.a)$Group)[get_variable(physeq.a.group, "Group")]
sample_data(physeq.a.group)$Soil <- levels(sample_data(physeq.a)$Soil)[get_variable(physeq.a.group, "Soil")]
```



```
# Filter taxa with mean abundance > 0.1
physeq2 = filter_taxa(physeq.a.group, function(x) mean(x) > 0.1, TRUE)

# Transform sample counts to relative abundances
physeq3 = transform_sample_counts(physeq2, function(x) x / sum(x))

# Aggregate taxa at the Phylum level
glom <- tax_glom(physeq3, taxrank = 'Phylum')

# Convert the aggregated phyloseq object to a dataframe
data <- psmelt(glom)
# Convert Phylum column to character
data$Phylum <- as.character(data$Phylum)

# Rename phyla with abundance < 1% to "< 1% abund."
data$Phylum[data$Abundance < 0.01] <- "< 1% abund."

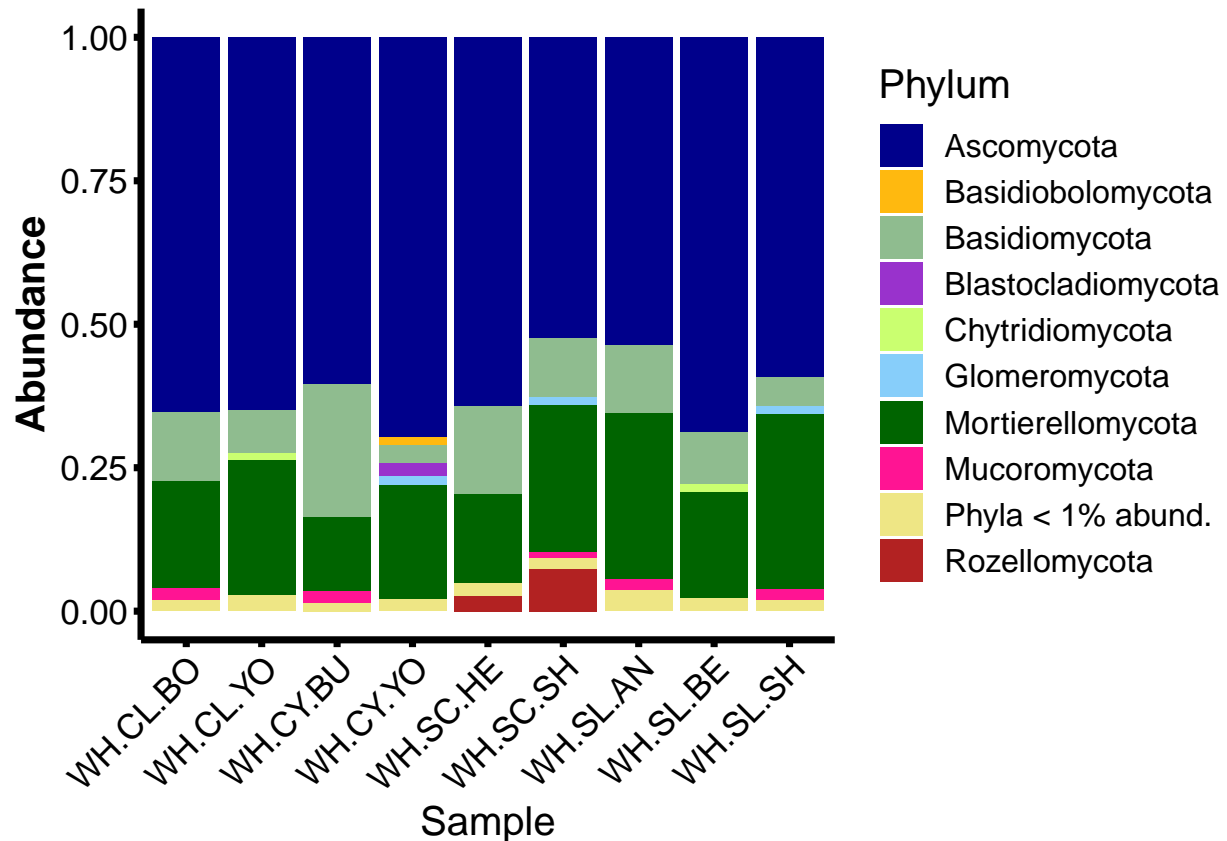
# Calculate medians of abundance for each Phylum
medians <- ddply(data, ~Phylum, function(x) c(median = median(x$Abundance)))

# Extract phyla with median abundance <= 0.01
remainder <- medians[medians$median <= 0.01, ]$Phylum

# Rename selected phyla to "Phyla < 1% abund."
data[data$Phylum %in% remainder, ]$Phylum <- "Phyla < 1% abund."

# Rename phyla with abundance < 1% to "Phyla < 1% abund."
data$Phylum[data$Abundance < 0.01] <- "Phyla < 1% abund."

# Plot the data with condensed phyla categories
p <- ggplot(data = data, aes(x = Sample, y = Abundance, fill = Phylum))
p + geom_bar(aes(), stat = "identity", position = "stack") +
  scale_fill_manual(values = c("darkblue", "darkgoldenrod1", "darkseagreen", "darkorchid", "darkolivegreen3"),
    theme(text = element_text(size=15, colour = "black"),
      axis.ticks = element_line(colour = "black", size = 1.25),
      axis.line = element_line(colour = 'black', size = 1.25),
      axis.text.x = element_text(angle=45, hjust=1, colour = "black", size = 13),
      axis.text.y = element_text(angle=0, hjust=0.5, colour = "black",size = 13),
      axis.title.y = element_text(color="black", size=15,face="bold"), legend.position = "right") # +
```



Number of taxa in groups Examining the cumulative count of taxa within a particular taxonomic group for each distinct subgroup involves the summation of the total number of taxa within that group while eliminating any overlapping taxa based on their names within individual subgroups. This process aids in comprehending the dynamics of microbiome populations.

```
library("dplyr")
```

```
physeq <- merge_samples(physeq.a, "Group", fun = sum)
gentab_levels <- list()
observationThreshold <- 1

# Define the taxonomic levels
genus_levels <- c("Phylum", "Class", "Order", "Family", "Genus", "Species")

# Something wrong with Kingdom Level, removed from the analysis
# loop through all the taxonomic levels
for (level in genus_levels) {

  # create a factor variable for each level
  genfac <- factor(tax_table(physeq)[, level])

  # calculate the abundance of each genus within each sample
  gentab <- apply(otu_table(physeq), MARGIN = 1, function(x) {
    tapply(x, INDEX = genfac, FUN = sum, na.rm = TRUE, simplify = TRUE)
  })
}
```

```

# calculate the number of samples in which each genus is observed above the threshold
level_counts <- apply(gentab > observationThreshold, 2, sum)

# create a data frame of level counts with genus names as row names
BB <- as.data.frame(level_counts)
BB$name <- row.names(BB)

# add the data frame to the gentab_levels list
gentab_levels[[level]] <- BB
}

# Combine all level counts data frames into one data frame
B2 <- gentab_levels %>% reduce(full_join, by = "name")

# Set row names and column names
row.names(B2) <- B2$name
B2$name <- NULL
colnames(B2)[1:6] <- genus_levels

print(B2)

```

```

##           Phylum Class Order Family Genus Species
## WH.CL.BO      14     38    80    162   233    263
## WH.CL.YO      14     35    70    129   189    218
## WH.CY.BU      14     37    70    137   191    210
## WH.CY.YO      13     36    66    128   180    206
## WH.SC.HE      13     41    87    165   245    282
## WH.SC.SH      14     43    85    174   259    305
## WH.SL.AN      14     36    81    156   230    268
## WH.SL.BE      14     41    77    144   202    227
## WH.SL.SH      14     36    78    154   241    276

```

```

# write.csv(B2, file = "ITS/level_counts_by_group.csv", row.names = TRUE)

```

Plotting the top 10 taxa at family level

Generating a visual representation through the plotting of the top 10 taxa at the family level offers a concise overview of the predominant microbial groups within a dataset. This approach is instrumental in identifying and emphasizing significant contributors to the structure and function of microbial communities across diverse groups or conditions.

```

## Transform normalised ASVs to proportions
proportions = transform_sample_counts(physeq.a.group, function(x) 100 * x/sum(x))

##
Top30ASVs = names(sort(taxa_sums(proportions), TRUE)[1:13])
Taxtab30 = cbind(tax_table(proportions), Family30 = NA)
Taxtab30[Top30ASVs, "Family30"] <- as(tax_table(proportions)[Top30ASVs, "Family"], "character")
tax_table(proportions) <- tax_table(Taxtab30)

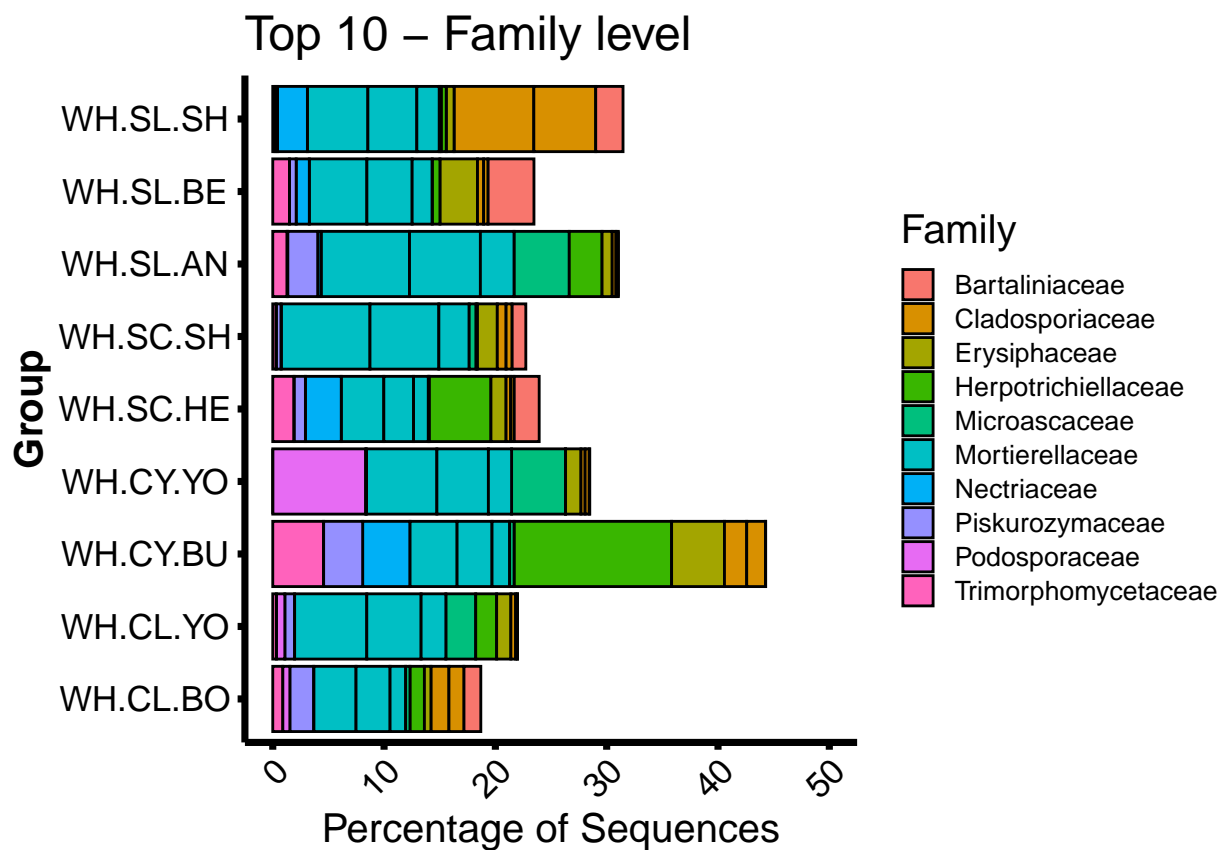
Rgsm30 = prune_taxa(Top30ASVs, proportions)

```

```

# plotting
title = "Top 10 - Family level"
plot_bar(Rgsm30, "Group", fill = "Family", title = title) +
  coord_flip() +
  ylab("Percentage of Sequences") + ylim(0, 50) + theme_classic() +
  theme(text = element_text(size=15, colour = "black"),
        axis.ticks = element_line(colour = "black", size = 1.25),
        axis.line = element_line(colour = 'black', size = 1.25),
        axis.text.x = element_text(angle=45, hjust=1, colour = "black", size = 13),
        axis.text.y = element_text(angle=0, hjust=0.5, colour = "black", size = 13),
        axis.title.y = element_text(color="black", size=15, face="bold"),
        legend.position = "right",
        legend.text = element_text(size = 9.5),
        legend.key.height= unit(0.45, 'cm'),
        legend.key.width= unit(0.45, 'cm')
  )

```



```

# Clean up by removing unnecessary objects
rm(proportions, Top30ASVs, Taxtab30, Rgsm30, title)

```

Upset plot using UpsetR

When it comes to representing sets visually, the go-to option is usually a Venn diagram. These diagrams work well when dealing with up to five sets, providing a clear visualisation. However, as the dataset expands,

such as when dealing with five sets, deriving the desired insights from the diagram becomes more complex. As a result, considering an UpSet graph for data visualisation becomes an appealing choice. UpSet graphs offer a more streamlined way to display intersections and complements, particularly when dealing with larger datasets or multiple sets. This option ensures a more intuitive and informative representation of the data.

```
library("UpSetR")
library("reshape2")
library("plyr")
library("dplyr")
# BiocManager::install("microbiome")
library("microbiome")

# Aggregate taxa at the genus level
B <- aggregate_taxa(physeq.a.group, "Genus", verbose = TRUE)

## [1] "Remove taxonomic information below the target level"
## [1] "Mark the potentially ambiguous taxa"
## [1] "-- split"
## [1] "-- sum"
## [1] "Create phyloseq object"
## [1] "Remove ambiguous levels"
## [1] "-- unique"
## [1] "-- Rename the lowest level"
## [1] "-- rownames"
## [1] "-- taxa"
## [1] "Convert to taxonomy table"
## [1] "Combine OTU and Taxon matrix into Phyloseq object"
## [1] "Add the metadata as is"

# Remove undesired genera
# B2 <- subset_taxa(B, !get("Genus") %in% c("uncultured", "Unknown"))

# Remove unwanted taxon names
taxa_to_remove <- c("uncultured", "Unknown")
B2 <- subset_taxa(B, !get("Genus") %in% taxa_to_remove)

# Extract relevant data from the phyloseq object
sample_data <- sample_data(B2)
otu_table <- otu_table(B2)
abundance <- as.vector(otu_table)

# Create a tibble with the extracted data
D <- tibble(
  Sample = rep(sample_data$Group, each = nrow(otu_table)),
  ASV = rep(rownames(otu_table), times = ncol(otu_table)),
  Abundance = abundance
) %>%
  group_by(Sample) %>%
  mutate(rank = rank(desc(Abundance))) %>%
  filter(Abundance > 0) %>%
  ungroup() %>%
  select(Sample, Abundance, ASV)
```

```

# Remove the Abundance column
D$Abundance <- NULL

# Rename the second column to "ASV"
names(D)[2] <- "ASV"

# Convert data from long to wide format
E <- dcast(D, ASV ~ Sample)

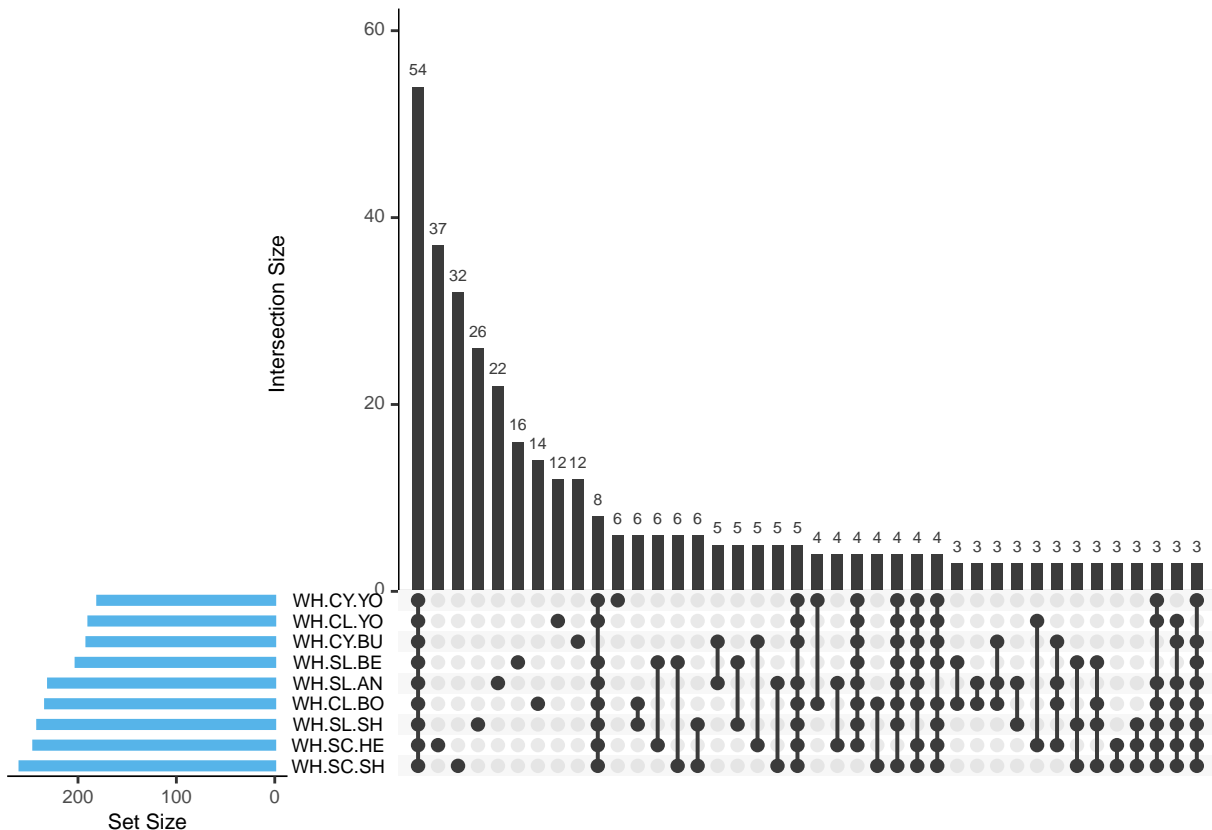
# Define a binary function
binary_fun <- function(x) {
  x[is.na(x)] <- 0
  ifelse(x > 0, 1, 0)
}

# Apply the binary function to columns 2 to 10
temp_df <- apply(E[2:10], 2, binary_fun)
temp_df <- as.data.frame(temp_df)

# Create an UpSet plot
upset_plot <- upset(temp_df, sets = colnames(temp_df), sets.bar.color = "#56B4E9",
                    order.by = "freq", empty.intersections = "on")

# Print the UpSet plot
print(upset_plot)

```



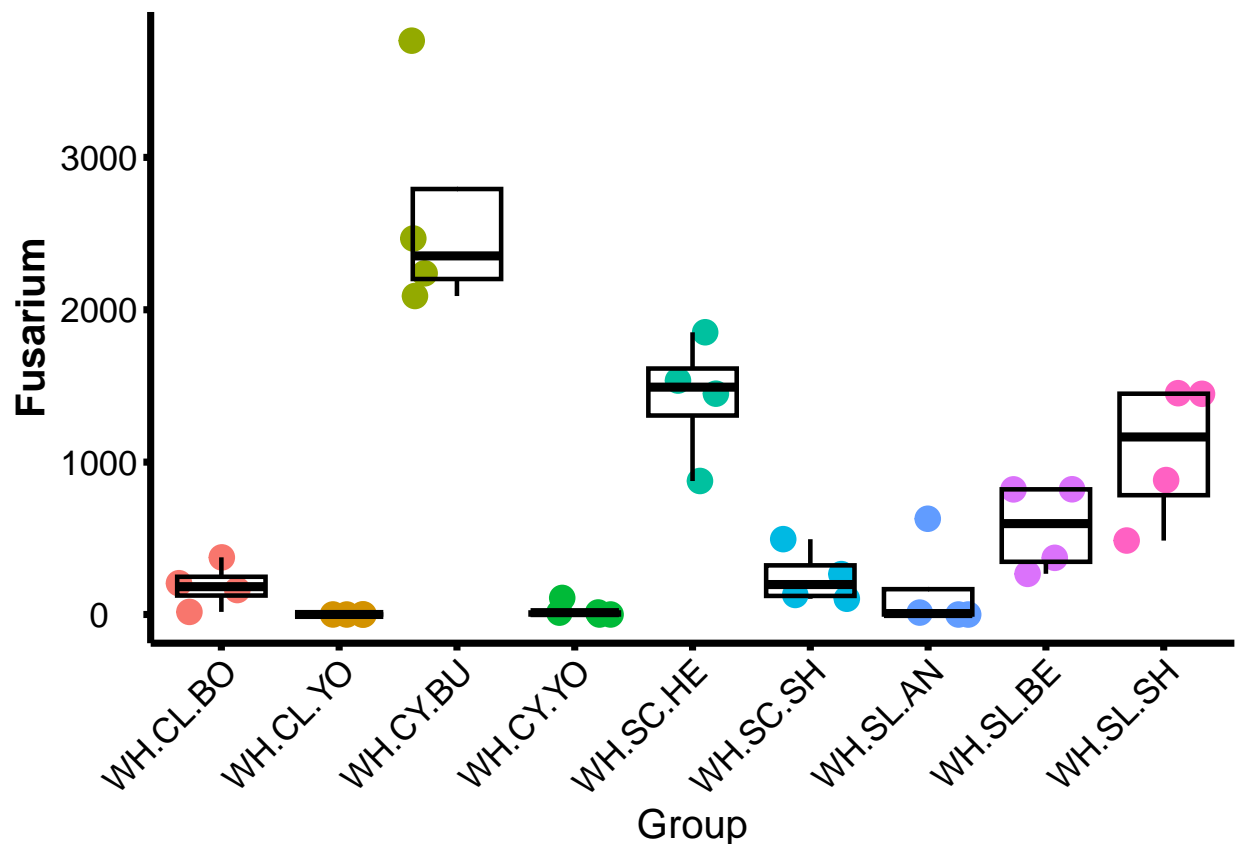
```

# Plotting for the abundance of one specific fungi
# Subset the taxa to Genus from physeq.wheat
physeq.a.genus <- subset_taxa(physeq.a, Genus == "Fusarium")

# Calculate the total abundance of Fusarium for each sample
meta = physeq.a.genus@sam_data
otudf = as.data.frame(t(as.data.frame(physeq.a.genus@otu_table)))
meta$Fusarium = rowSums(otudf)

# Plot a graph of the abundance of Fusarium for each sample grouped by Group:
ggplot(subset(meta, Group %in% c("WH.CL.BO", "WH.CL.YO",
                                "WH.CY.BU", "WH.CY.YO",
                                "WH.SC.HE", "WH.SC.SH",
                                "WH.SL.AN", "WH.SL.BE",
                                "WH.SL.SH")),
  aes(x = Group, y = Fusarium, colour = interaction(Group))) +
  geom_point(alpha = 1, position = "jitter", size = 4) +
  geom_boxplot(alpha = 0, colour = "black", size = 0.8) +
  theme_classic() +
  theme(text = element_text(size=15, colour = "black"),
    axis.ticks = element_line(colour = "black", size = 1.25),
    axis.line = element_line(colour = 'black', size = 1.25),
    axis.text.x = element_text(angle=45, hjust=1, colour = "black", size = 13),
    axis.text.y = element_text(angle=0, hjust=0.5, colour = "black", size = 13),
    axis.title.y = element_text(color="black", size=15, face="bold"), legend.position = "none")

```



###Phylogenetic tree Phylogenetic trees are branching diagrams that show the evolutionary relationships between different species or other entities. They are constructed by comparing the organisms' DNA, proteins, or other features. The more similar the features, the more closely related the organisms are thought to be. It can be used to answer a variety of questions about evolution, such as how different species are related to each other, when different species diverged from each other, and what are the common ancestors of different species.

```
# Load required libraries
library(phyloseq)  # For handling phylogenetic sequencing data
library(ggtree)    # For tree visualisation
library(scales)    # For scaling transformations
```

There could be an error message and it could be ignored (<https://github.com/YuLab-SMU/tidytrees/issues/10>)

```
# Load the GlobalPatterns dataset and prune taxa
GP <- prune_taxa(taxa_sums(physeq.a) > 0, physeq.a) # Remove taxa with zero sums
GP.chl <- subset_taxa(GP, Genus == "Fusarium") # Subset data based on Phylum value

# Create a ggtree plot
p <- ggtree(GP.chl, ladderize = FALSE) +
  geom_text2(aes(subset = !isTip, label = label), hjust = -0.3, size = 3.5) +
  geom_tiplab(aes(label = Species), as_ylab=TRUE) +
  geom_point(aes(x = x + hjust, color = Group,
                 shape = Group, size = Abundance), na.rm = TRUE) +
  scale_size_continuous(trans = log_trans(2)) +
  scale_shape_manual(values = c(15, 17, 3, 4, 16, 18, 21, 22, 23)) + # Set custom
  # theme_classic() +
  theme(text = element_text(size=15, colour = "black"),
        axis.ticks = element_line(colour = "black", size = 1.25),
        axis.line = element_line(colour = "black", size = 1.25),
        # axis.text.x = element_text(angle=45, hjust=1, colour = "black", size = 13),
        axis.text.y = element_text(angle=0, hjust=0.5, colour = "black",size = 2.5),
        axis.title.y = element_text(color="black", size=2.5,face="bold"), legend.position = "top")

# Print the ggtree plot
print(p)
```


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- ◆ WH.SL.AN

