Two methods for mapping and visualizing associated data on phylogeny using ggtree

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1. Examples of mapping and visualizing associated data on phylogenetic trees

ggtree (Yu et al. 2017) supports tree with data parsed by treeio package and thus supports evolutionary statistics inferred by commonly used software packages including ASTRAL (Mirarab and Warnow 2015), BEAST (Bouckaert et al. 2014), EPA (Berger, Krompass, and Stamatakis 2011), HyPhy (Pond, Frost, and Muse 2005), IQ-TREE (Nguyen et al. 2015), MrBayes (Huelsenbeck and Ronquist 2001), PAML (Yang 2007), PHYLODOG (Boussau et al. 2013), pplacer (Matsen, Kodner, and Armbrust 2010), r8s (Sanderson 2003), RAXML (Stamatakis 2014) and RevBayes (Höhna et al. 2014) to be used to annotate the tree. In addition, ggtree supports tree objects defined by other R packages, including obkData, phyloseq, phylo4 and phylo4d, so that associated data stored in these objects (e.g. outbreak data in obkData, microbiome data in phyloseq and matrix in phylo4d) can be used directly to annotate the tree (see Appendix of (Yu et al. 2017), Fig. S3 and S8A). The methods introduced in this paper emphasize on integrating external data. Here, we present several examples to demonstrate how data can be mapped and visualized to annotate phylogenetic trees. More examples can be found on published figures that collected on ggtree project website¹.

1.1. Integrating node/edge data

This example uses method 1, the %<+% operator, to integrate taxon (tip_data.csv) and internal node (inode_data.csv) information and map the data to different colors or shapes of symbolic points and labels. The tip data contains imageURL that links to online figures of the species, which can be parsed and used as tip labels in ggtree. ggtree supports labeling both internal or external nodes using local or online image files such as those deposited on phylopic database². This also provides a solution of using subplots to annotate the tree. Online tools such as iTOL (Letunic and Bork 2007) and EvolView (He et al. 2016) support displaying subplots on phylogenetic tree. However only bar and pie charts are supported by these tools. Users may want to visualize node-associated data with other visualization methods, such as violin plot (Grubaugh et al. 2017), venn diagram (Lott et al. 2015), sequence logo etc, and display them on the tree. In ggtree, all kinds of subplots are supported as we can export all subplots to image files and use them to label corresponding nodes on the tree.

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 $^{^{1}} https://guangchuangyu.github.io/software/ggtree/gallery/\\$

 $^{^2}$ http://phylopic.org/

```
geom_tiplab(geom = "label", offset = 1, hjust = .5) +
  geom_tippoint(aes(shape = trophic_habit, color = trophic_habit, size = mass_in_kg)) +
  theme(legend.position = "right") + scale_size_continuous(range = c(3, 10))

d2 <- read.csv(paste0(url, "inode_data.csv"))
p3 <- p2 %<+% d2 + geom_label(aes(label = vernacularName.y, fill = posterior)) +
    scale_fill_gradientn(colors = RColorBrewer::brewer.pal(3, "YlGnBu"))
print(p3)</pre>
```

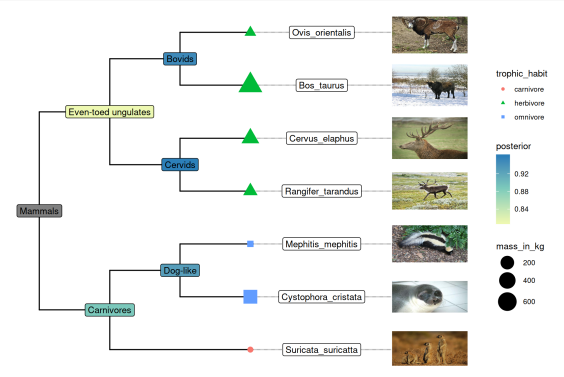


Fig. S1: Example of attaching multiple datasets and labelling taxa with image files

Although the data integrated by %<+% operator in ggtree is for tree visualization, the data attached to the ggtree graphic object can be converted to treedata object that contains the tree and the attached data. The data stored in the treedata object can be used directly to annotate the tree in ggtree.

```
## convert ggtree object to treedata object
y <- as.treedata(p3)
\textit{## all the data in 'tip\_data.csv' and 'inode\_data.csv' were incorporated}
## 'treedata' S4 object'.
##
## ...@ phylo:
## Phylogenetic tree with 7 tips and 6 internal nodes.
##
## Tip labels:
## Rangifer_tarandus, Cervus_elaphus, Bos_taurus, Ovis_orientalis, Suricata_suricatta, Cystophora_cris
## Node labels:
## [1] "Mammalia"
                       "Artiodactyla" "Cervidae"
                                                      "Bovidae"
## [5] "Carnivora"
                       "Caniformia"
```

```
##
## Rooted; includes branch lengths.
##
## with the following features available:
##
    'vernacularName.x', 'imageURL', 'imageLicense', 'imageAuthor',
    'infoURL.x',
                                   'trophic habit',
                                                         'ncbi_taxid',
##
                    'mass in kg',
    'vernacularName.y', 'infoURL.y',
                                                     'bootstrap'.
##
                                         'rank.y',
                                                                     'posterior'.
## Now all the features (from external data attahced to the tree), that stored in
## tree object can be used to annotate the tree.
ggtree(y) + geom_tiplab(aes(label = vernacularName.x, color = trophic_habit)) +
    geom_label(aes(x = branch, label = vernacularName.y, fill = bootstrap)) +
    scale_fill_gradientn(colors = RColorBrewer::brewer.pal(3, "YlGnBu")) +
    theme(legend.position = "right") + xlim(-.1, 3.5)
```

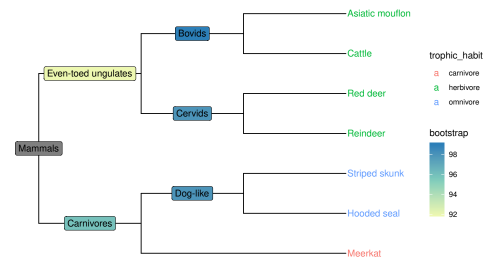


Fig. S2: Example of visualizing treedata object that contains tree with associated data

The treedata object can be exported to BEAST compatible NEXUS file³, which can be parsed by treeio or FigTree⁴. This creates the possibility to integrate tree-associated data with the tree into a single file and visualize the data not only with ggtree, but also FigTree, etc.

1.2. Visualizing species abundance distribution with phylogenetic tree

Species abundance is continuous numerical data and usually summarized as boxplot, violinplot or density curve, which are all suported by facet_plot (Table S1). This example uses microbome data that provided in phyloseq package and density ridgeline is employed to visualize the abundance data. facet_plot automatically re-arranges the abundance data according to the tree structure, visualizes the data using the specify geom function, *i.e.* geom_density_ridges, and align the density curves with the tree as demonstrated in Fig. S3. Note that data stored in the phyloseq object is visible to ggtree and can be used directly in tree visualization (Phylum was used to color tips and density ridgelines in this example).

```
library(phyloseq)
library(ggridges)
library(dplyr)
library(ggtree)
```

 $^{^3}$ https://bioconductor.org/packages/release/bioc/vignettes/treeio/inst/doc/Exporter.html

⁴http://tree.bio.ed.ac.uk/software/figtree/

```
data("GlobalPatterns")
GP <- GlobalPatterns
GP <- prune_taxa(taxa_sums(GP) > 1000, GP)
sample_data(GP)$human <- get_variable(GP, "SampleType") %in% c("Feces", "Skin")</pre>
mergedGP <- merge_samples(GP, "SampleType")</pre>
mergedGP <- rarefy_even_depth(mergedGP,rngseed=394582)</pre>
mergedGP <- tax_glom(mergedGP,"Order")</pre>
melt_simple <- psmelt(mergedGP) %>%
  filter(Abundance < 120) %>%
  select(OTU, val = Abundance)
p <- ggtree(mergedGP) +</pre>
  geom_tippoint(aes(color = Phylum), size = 1.5)
facet_plot(p, panel="Abundance", data=melt_simple,
           geom_density_ridges, mapping = aes(x = val, group = label,
                                          fill = Phylum),
           color = 'grey80', lwd = .3)
```

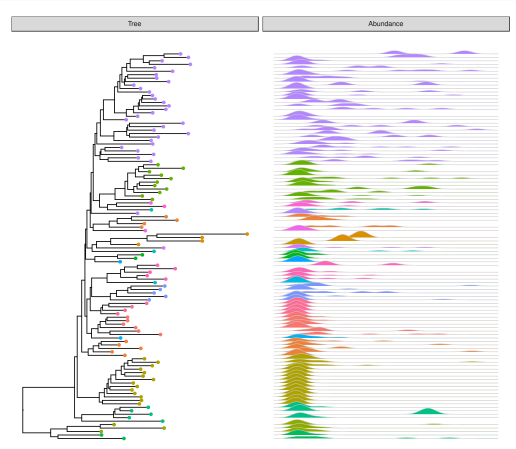


Fig. S3: Phylogenetic tree with OTU abundance densities

1.3 Visualizing pairwise nucleotide sequence distance with phylogenetic tree

This example reproduces Fig. 1 of (Chen et al. 2017). It extracts accession numbers from tip labels of the HPV58 tree and calculates pairwise nucleotide sequence distances. The distance matrix is visualized as dot and line plots. This example demonstrating the abilities of adding multiple layers to a specific panel. As illustrated on Fig. S4, the facet_plot function displays sequence distances as a dot plot and then adds a layer of line plot to the same panel, *i.e.* Sequence Distance. In addition, the tree in facet_plot can be fully annotated with multiple layers (clade labels, bootstrap support values, *etc*).

```
library(tibble)
library(tidyr)
library(Biostrings)
library(treeio)
library(ggplot2)
library(ggtree)
hpvtree <- paste0("https://raw.githubusercontent.com/GuangchuangYu/",
                  "plotting_tree_with_data/master/HPV58.tree")
tree <- read.tree(hpvtree)</pre>
clade \leftarrow c(A3=92, A1=94, A2=108, B1=156, B2=159, C=163, D1=173, D2=176)
tree <- groupClade(tree, clade)</pre>
cols <- c(A1="#EC762F", A2="#CA6629", A3="#894418", B1="#0923FA",
         B2="#020D87", C="#000000", D1="#9ACD32", D2="#08630A")
## visualize the tree with tip labels and tree scale
p <- ggtree(tree, aes(color = group), ladderize = FALSE) %% rotate(rootnode(tree)) +
    geom_tiplab(aes(label=paste0("italic('", label, "')")), parse = TRUE, size = 2.5) +
    geom\_treescale(x = 0, y = 1, width = 0.002) +
    scale_color_manual(values = c(cols, "black"), na.value = "black", name = "Lineage",
                      breaks = c("A1", "A2", "A3", "B1", "B2", "C", "D1", "D2")) +
    guides(color = guide_legend(override.aes = list(size = 5, shape = 15))) +
    theme_tree2(legend.position = c(.1, .88))
## Optional
## add labels for monophyletic (A, C and D) and paraphyletic (B) groups
p <- p + geom_cladelabel(94, "italic(A1)", color = cols[["A1"]], offset = .003, align = TRUE,
                    offset.text = -.001, barsize = 1.2, extend = c(0, 0.5), parse = TRUE) +
    geom_cladelabel(108, "italic(A2)", color = cols[["A2"]], offset = .003, align = TRUE,
                   offset.text = -.001, barsize = 1.2, extend = 0.5, parse = TRUE) +
    geom_cladelabel(131, "italic(A3)", color = cols[["A3"]], offset = .003, align = TRUE,
                   offset.text = -.001, barsize = 1.2, extend = c(0.5, 0), parse = TRUE) +
    geom_cladelabel(92, "italic(A)", color = "darkgrey", offset = .00315, align = TRUE,
                   offset.text = 0.0002, barsize = 2, fontsize = 5, parse = TRUE) +
    geom_cladelabel(156, "italic(B1)", color = cols[["B1"]], offset = .003, align = TRUE,
                   offset.text = -.001, barsize = 1.2, extend = c(0, 0.5), parse = TRUE) +
    geom_cladelabel(159, "italic(B2)", color = cols[["B2"]], offset = .003, align = TRUE,
                   offset.text = -.001, barsize = 1.2, extend = c(0.5, 0), parse = TRUE) +
    geom_strip(65, 71, "italic(B)", color = "darkgrey", offset = 0.00315, align = TRUE,
                   offset.text = 0.0002, barsize = 2, fontsize = 5, parse = TRUE) +
    geom_cladelabel(163, "italic(C)", color = "darkgrey", offset = .0031, align = TRUE,
                   offset.text = 0.0002, barsize = 3.2, fontsize = 5, parse = TRUE) +
    geom_cladelabel(173, "italic(D1)", color = cols[["D1"]], offset = .003, align = TRUE,
                   offset.text = -.001, barsize = 1.2, extend = c(0, 0.5), parse = TRUE) +
    geom_cladelabel(176, "italic(D2)", color = cols[["D2"]], offset = .003, align = TRUE,
```

```
offset.text = -.001, barsize = 1.2, extend = c(0.5, 0), parse = TRUE) +
    geom_cladelabel(172, "italic(D)", color = "darkgrey", offset = .00315, align = TRUE,
                   offset.text = 0.0002, barsize = 2, fontsize = 5, parse = TRUE)
## Optional
## display support values
p <- p + geom_nodelab(aes(subset = (node == 92), label = "*"),
                    color = "black", nudge_x = -.001, nudge_y = 1) +
    geom nodelab(aes(subset = (node == 155), label = "*"),
                    color = "black", nudge_x = -.0003, nudge_y = -1) +
    geom_nodelab(aes(subset = (node == 158), label = "95/92/1.00"),
                    color = "black", nudge_x = -0.0001, nudge_y = -1, hjust = 1) +
    geom_nodelab(aes(subset = (node == 162), label = "98/97/1.00"),
                    color = "black", nudge_x = -0.0001, nudge_y = -1, hjust = 1) +
    geom_nodelab(aes(subset = (node == 172), label = "*"),
                    color = "black", nudge_x = -.0003, nudge_y = -1)
## extract accession numbers from tip labels
tl <- tree$tip.label
acc <- sub("\\w+\\|", "", tl)
names(tl) <- acc</pre>
## read sequences from GenBank directly into R
## and convert the object to DNAStringSet
tipseq <- ape::read.GenBank(acc) %>% as.character %>%
    lapply(.,paste0,collapse = "") %>% unlist %>%
    DNAStringSet
## align the sequences using muscle
tipseq_aln <- muscle::muscle(tipseq)</pre>
tipseq aln <- DNAStringSet(tipseq aln)</pre>
## calculate pairwise hamming distances among sequences
tipseq_dist <- stringDist(tipseq_aln, method = "hamming")</pre>
## calculate percentage of differences
tipseq_d <- as.matrix(tipseq_dist)/width(tipseq_aln[1]) * 100</pre>
## convert the matrix to tidy data frame for facet_plot
dd <- as data frame(tipseq d)</pre>
dd$seq1 <- rownames(tipseq d)</pre>
td <- gather(dd,seq2, dist, -seq1)
td$seq1 <- tl[td$seq1]
td$seq2 <- t1[td$seq2]
g <- p$data$group
names(g) <- p$data$label</pre>
td$clade <- g[td$seq2]
## visualize the sequence differences using dot plot and line plot
## and align the sequence difference plot to the tree using facet_plot
p2 <- facet_plot(p, panel = "Sequence Distance", data = td, geom_point,
            mapping = aes(x = dist, color = clade, shape = clade), alpha = .6) %%
    facet_plot(panel = "Sequence Distance", data = td, geom = geom_path,
            mapping=aes(x = dist, group = seq2, color = clade), alpha = .6) +
    scale_shape_manual(values = 1:8, guide = FALSE)
```

print(p2)

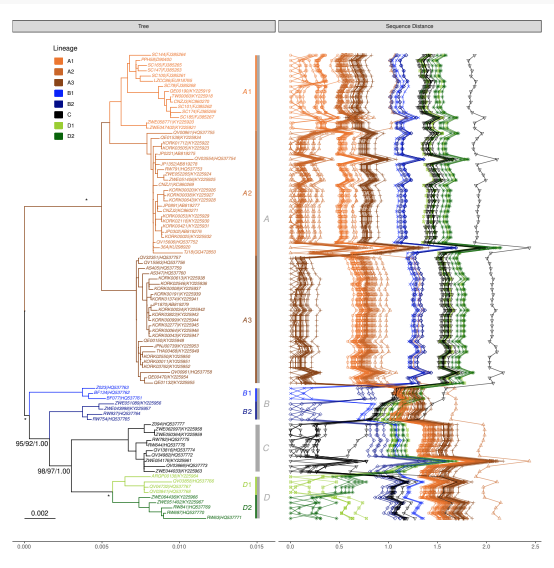


Fig. S4: Phylogeny of HPV58 complete genomes with dot and line plots of pairwise nucleotide sequence distances

2. Geometric layers that supported by facet_plot

facet_plot is a general solution for linking graphic layer to a tree. The function internally re-order the input data based on the tree structure and visualize the data at the specific panel by the geometric function. Users are freely to visualize several panels to plot different types of data as demonstrated in Fig. S6 and use different geometric functions to plot the same dataset (Fig. S4) or different datasets (Fig. S8) on the same panel.

facet_plot is designed to work with most of the geom functions defined in ggplot2 (Wickham 2016) and other ggplot2-based packages. Here is the list of the geometric functions that works seamlessly with facet_plot. As the ggplot2 community keeps expanding and more geom functions will be implemented in either ggplot2 or other extensions, facet_plot will gains more power to present data in future. Note that different geom functions can be combined to present data on the same panel (Fig. S4 and S8) and the

Table S1: Geometric layers that supported by 'facet_plot()'

Package	Geom Layer	Description
ggalt	$geom_dumbbell$	create dumbbell charts
ggbio	geom_alignment	show interval data as alignmen
ggfittext	geom_fit_text	shrinks, grows or wraps text to fit inside a defined rectangular area
gggenes	geom_gene_arrow	draws genes as arrows
ggimage	geom_image geom_phylopic	visualizes image files queries image files from phylopic database and visualizes them
	geom_hline geom_jitter geom_label geom_point	adds horizontal lines adds a small amount of random variation to the location of each point draws a rectangle behind the text creats scatterplots
ggplot2	geom_raster geom_rect geom_segment geom_spoke geom_text geom_tile geom_vline	a high performance special case for all the tiles are the same size draw rectangle by using the locations of the four coners draws a straight line between points a polar parameterisation of 'geom_segment()' adds text to the plot draw rectangle by using the center of the tile and its size adds vertical lines
ggrepel	geom_text_repel geom label repel	adds text to the plot. The text labels repel away from each other and away from the data points draws a rectangle underneath the text. The text labels repel away from each other and away from the data points
ggridges	geom_density_ridges geom_density_ridges_gradient geom_ridgeline geom_ridgeline_gradient	arranges multiple density plots in a staggered fashion works just like 'geom_density_ridges' except that the 'fill' aesthetic can vary along the x axis plots the sum of the 'y' and 'height' aesthetics versus 'x', filling the area between 'y' and 'y + height' with a color works just like 'geom_ridgeline' except that the 'fill' aesthetic can vary along the x axis
ggstance	geom_barh geom_boxploth geom_crossbarh geom_errorbarh geom_histogramh geom_linerangeh geom_pointrangeh geom_violinh	horizontal version of 'geom_bax()' horizontal version of 'geom_boxplot()' horizontal version of 'geom_erossbar()' horizontal version of 'geom_errorbarh()' horizontal version of 'geom_histogram()' horizontal version of 'geom_linerange()' horizontal version of 'geom_pointrange()' horizontal version of 'geom_violin()'
ggtree	geom motif	draws aligned motifs

combinations of different geom functions create the possibility to present more complex data with phylogeny.

3. Comparing ggtree with other R packages

We have presented detail comparison of ggtree with ape (Paradis, Claude, and Strimmer 2004), phytools (Revell 2012), phyloseq (McMurdie and Holmes 2013) and OutbreakTools (Jombart et al. 2014) on Appendix S1 of (Yu et al. 2017). Here we extend the comparison with plotTree scripts⁵, metacoder (Foster, Sharpton, and Grünwald 2017) and phylobase⁶, which have the ability to integrate external data.

3.1 plotTree scripts

Although plotTree is not an R package, we include the scripts for comparision as the scripts has the abilities of plotting tree with basic information, heatmap and bar plots. Phylogenetic tree is visualized by plot.phylo provided by ape (Paradis, Claude, and Strimmer 2004). The tip information used in plotTree is not really integrated, as it cannot be mapped to visual characteristics, such as color, shape and size that are supported in ggtree. The information is hard-coded to only color tip with circle symbols and print the text next to the tips. In addition, there is no solution for mapping data to internal nodes. For plotting tree with data, plotTree only supports heatmap for matrix, barplot for numerical data, dotplot for allele data and line segments for genome blocks. The capabilities is restricted to specific needs and only applied to specific data. An example of plotting tree with data using plotTree is demonstrated in Fig. S5 and a corresponding ggtree version is illustrated in Fig. S6.

⁵https://github.com/katholt/plotTree

⁶https://CRAN.R-project.org/package=phylobase

```
## need to clone the repo before running the scrtip
## access date: 2018-07-23
## qit clone https://qithub.com/katholt/plotTree.qit
setwd("plotTree/tree_example_april2015")
source("../plotTree.R")
## the 'location' that used to color the nodes, cannot be used to color bar and snp data.
##
## issues:
## 1. bar data can not be displayed properly.
## 2. legend not shown properly,
        there is no taxa sampled from locations label as 'other' or 'VN'.
## 3. the positions of data panels are hard-coded.
plotTree(tree = "tree.nwk",
         infoFile = "info.csv", infoCols = NA,
         colourNodesBy = "location", legend.pos = "topleft",
         barData = "bar.csv", snpFile = "alleles.csv")
```

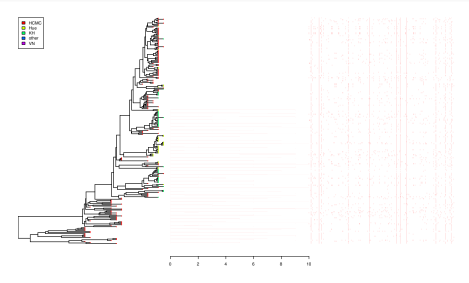


Fig. S5: Example of plotTree

ggtree provide general solutions for integrating data. Method 1, the %<+% operator, can integrate external and internal node data and map the data as visual characteristics to visualize the tree (Fig. S1) and other datasets used in facet_plot (in Fig. S6, SNP and Trait data were colored by isolation location that was attached by %<+%). Method 2, the facet_plot function, has no restriction of input data as long as there is a geom function available to plot the data (e.g. species abundance displayed by geom_density_ridges that demonstrated in Fig. S3). Users are free to combine different panels (Fig. S6) and combine different geom layers in the same panel (Fig. S4 and S8). The plotTree scritps can be easily reproduced using ggtree⁷, while ggtree offers more flexibility and can do much more. As ggtree is more powerful, the authors of plotTree also use ggtree to present tree with associated data including genotypes, isolation locations, patient status, mutations etc, in their recent paper (Britto et al. 2018).

⁷https://github.com/GuangchuangYu/plotTree-ggtree

```
tree <- read.tree(paste0(remote_folder, "tree.nwk"))</pre>
## read the sampling information data set
info <- read.csv(paste0(remote_folder, "info.csv"))</pre>
## read and process the allele table
snps<-read.csv(paste0(remote_folder, "alleles.csv"), header = F,</pre>
                 row.names = 1, stringsAsFactor = F)
snps strainCols <- snps[1,]</pre>
snps<-snps[-1,] # drop strain names</pre>
colnames(snps) <- snps_strainCols</pre>
gapChar <- "?"</pre>
snp <- t(snps)</pre>
lsnp <- apply(snp, 1, function(x) {</pre>
        x != snp[1,] & x != gapChar & snp[1,] != gapChar
lsnp <- as.data.frame(lsnp)</pre>
lsnp$pos <- as.numeric(rownames(lsnp))</pre>
lsnp <- tidyr::gather(lsnp, name, value, -pos)</pre>
snp_data <- lsnp[lsnp$value, c("name", "pos")]</pre>
## read the trait data
bar_data <- read.csv(paste0(remote_folder, "bar.csv"))</pre>
## visualize the tree
p <- ggtree(tree)</pre>
## attach the sampling information data set
## and add symbols colored by location
p <- p %<+% info + geom_tippoint(aes(color=location))</pre>
## visualize SNP and Trait data using dot and bar charts,
## and align them based on tree structure
p2 <- facet_plot(p, panel = "SNP", data = snp_data, geom = geom_point,
                 mapping=aes(x = pos, color = location), shape = '|') %>%
    facet_plot("Trait", bar_data, ggstance::geom_barh,
                 aes(x = dummy_bar_value, color = location, fill = location),
                 stat = "identity", width = .6) +
    theme_tree2(legend.position=c(.05, .85))
print(p2)
```

3.2 metacoder

metacoder (Foster, Sharpton, and Grünwald 2017) is designed for visualizing herachical data, mainly for community taxonomic diversity data. It produces heat tree that uses e.g. taxa abundance to scale color and size of nodes and edges. This is similar to method 1 presented in our manuscript. ggtree (Yu et al. 2017) is more flexible and can do more, such as coloring text labels and plotting variety of node shapes using categorical data.

The heat tree implemented in metacoder is designed for herachical data, but not for phylogeny. metacoder internally use graph layout algorithms implemented in igraph (Csardi and Nepusz 2006) and doesn't support widely used phylogenetic tree layouts (e.g. rectangular and slanted layouts). Edge length for phylogenetic

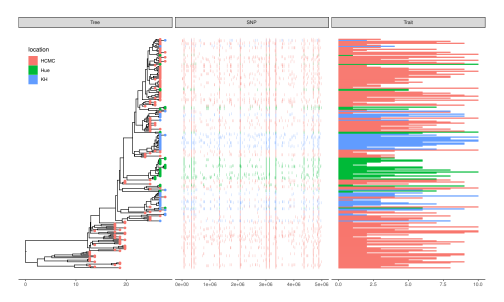


Fig. S6: Example of plotting SNP and trait data using ggtree

tree is also not supported.

ggtree is designed for general purpose and can be easily extended for specific research applications. For example, the R package, microbiomeViz⁸ for visualizing microbiome data, is developed based on ggtree. microbiomeViz is similar to the python library, GraPhlAn, which supports phylogeny and has better annotation abilities than metacoder (according to (Foster, Sharpton, and Grünwald 2017)).

In addition, ggtree supports visualizing microbiome data stored in the phyloseq object (McMurdie and Holmes 2013). Examples can be found on Appendix S1 of (Yu et al. 2017) and Fig. S3.

3.3 phylobase

The phylobase⁹ package defines phylo4d class that combines a tree with a data frame, and provides plot method, which internally call the treePlot function, to display the tree with the data. However there are some restrictions of the plot method, it can only plot numeric values for tree-associated data as bubbles and cannot generate figure legend. Phylobase doesn't implement visualization method to display categorical values. Using associated data as visual characteristics such as color, size and shape, is also not supported. Although it is possible to color the tree using associated data, it requires users to extract the data and map them to color vector manually follow by passing the color vector to the plot method. This is tedious and error-prone since the order of the color vector needs to be consistent with the edge list stored in the object.

Here is the example of plotting associated data with phylobase:

```
library(phylobase)
data(geospiza_raw)
g1 <- as(geospiza_raw$tree, "phylo4")
g2 <- phylo4d(g1, geospiza_raw$data, missing.data="warn")

## Warning in formatData(phy = x, dt = tip.data, type = "tip", ...): The
## following nodes are not found in the dataset: olivacea
plot(g2)</pre>
```

⁸https://github.com/lch14forever/microbiomeViz

⁹https://CRAN.R-project.org/package=phylobase

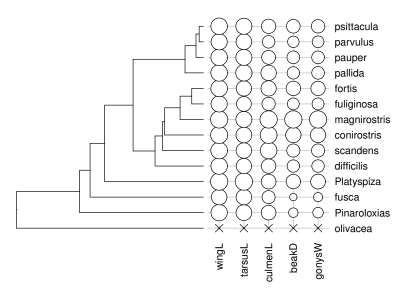


Fig. S7: Example of phylobase

The phylo4d object is supported directly by ggtree, so that we can use ggtree(g2) to directly visualize the tree. All the associated data stored in the phylo4d object can be used directly to annotate the tree (Fig. S8A). In addition, users can use facet_plot to visualize the associated data (Fig. S8B).

```
d1 \leftarrow data.frame(x = seq(0.93, 1.15, length.out = 5),
                lab = names(geospiza_raw$data))
## plot bubbles directly using data stored in 'q2'
p1 <- ggtree(g2) + geom_tippoint(aes(size = wingL), x = d1*x[1], shape = 1) +
    geom_tippoint(aes(size = tarsusL), x = d1$x[2], shape = 1) +
    geom_tippoint(aes(size = culmenL), x = d1$x[3], shape = 1) +
    geom_tippoint(aes(size = beakD), x = d1$x[4], shape = 1) +
    geom_tippoint(aes(size = gonysW), x = d1$x[5], shape = 1) +
    scale size continuous(range = c(3,12), name="") +
    geom_text(aes(x = x, y = 0, label = lab), data = d1, angle = 90) +
    geom tiplab(offset = .3) + xlim(0, 1.3) +
    theme(legend.position = c(.1, .75)) + labs(tag = "A")
library(dplyr)
library(tidyr)
## extract tip data from 'g2' and use 'facet_plot' to visualze the data
d <- tipData(g2)</pre>
d$tip <- rownames(d)</pre>
dd <- gather(d, feature, value, -tip)</pre>
cat <- seq(ncol(d))
names(cat) <- names(d)</pre>
dd$cat <- cat[dd$feature]</pre>
d2 <- select(dd, -value) %>% filter(tip == 'fuliginosa')
p <- ggtree(g2) + geom_tiplab() + xlim_tree(c(0, 1.2))</pre>
p2 <- facet_plot(p, "Morphometric data", dd, geom_point, aes(x = cat, size = value), shape = 1) %>%
    facet_plot("Morphometric data", d2, geom_text, aes(x = cat, y = 0, label = feature), angle = 90) +
    scale_size_continuous(range = c(3, 12)) +
    theme(legend.position = "right") +
```

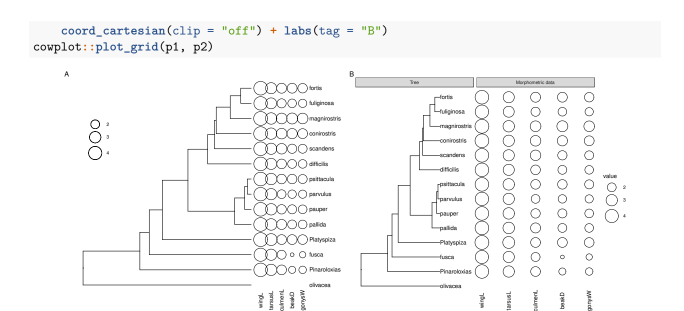
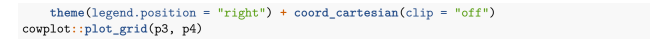


Fig. S8: Visualizing phylo4d data using ggtree

phylobase only provides plot method to visualize numerical values, and will ignore categorical values if there are any. In ggtree, we are able to attach additional information, either numerical or categorical data, and are able to display these information in our favorite way. Here as an example, the diet information was attached to the tree and used to color symbolic points, tip labels (Fig. S9A) and phenotypic data (Fig. S9B). The numerical values can be visualized not only as bubble plot (Fig. S8) but also heatmap (Fig. S9A) or other types (e.g. stacked bars). Heatmap is commonly used in phylogeny for comparative study. Before developing facet_plot, ggtree implemented a gheatmap function (Yu et al. 2017) to visualize phylogenetic tree with heatmap of numerical or categorical values. facet_plot is a general solution for plotting data with the tree, including heatmap (Fig. S9A). gheatmap is specifically designed for plotting heatmap with tree and provides shortcut for handling column labels and color palette. Another difference is that facet_plot only supports rectangular and slanted tree layouts while gheatmap supports rectangular, slanted and circular layouts. gheatmap works seemlessly with facet_plot as illustrated in Fig. S9B.

```
diet = data.frame(species=tipLabels(g2),
    Diet = c("Seeds", "Seeds", "Cacti", "Cacti", "Seeds", "Insects",
         "Insects", "Insects", "Insects", "Fruits", "Insects", "Insects", "Insects"))
p <- ggtree(g2) %<+% diet + geom_tiplab(aes(color = Diet), offset = .05) +
    geom_tippoint(aes(color = Diet), size = 5, alpha = .5) +
    xlim_tree(c(0, 1.2))
p2 <- gheatmap(ggtree(g2), tipData(g2), colnames_angle = 90) %<+% diet
p3 <- facet_plot(p, "Morphometric data", dd, geom_tile, aes(x = cat, fill = value)) %>%
    facet_plot("Morphometric data", d2, geom_text,
               aes(x = cat, y = 0, label = feature), angle = 90) +
    scale_fill_viridis_c(na.value = "white") + labs(tag = "A") + theme(legend.position="right")
p4 <- facet_plot(p2, "Morphometric data", dd, geom_point,
                 aes(x = cat, size = value, color = Diet)) %>%
    facet_plot("Morphometric data", d2, geom_text,
               aes(x = cat, y = 0, label = feature), angle = 90) +
    scale_size_continuous(range = c(3, 12)) + labs(tag = "B") +
```



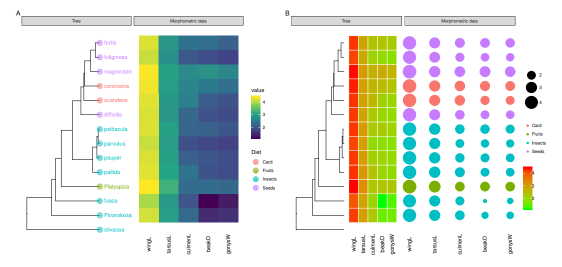


Fig. S9: Visualizing phylo4d data as heatmap using 'facet_plot()' with additional data to color taxa (A), 'facet_plot()' works with 'gheatmap' (B)

3.4 Summary

Although there are many software packages support visualizing phylogenetic tree, plotting tree with data is often missing or with only limited supports. Some of the packages defines \$4 classes to store phylogenetic tree with domain specific data, such OutbreakTools (Jombart et al. 2014) defined obkData for storing tree with epidemiology data and phyloseq (McMurdie and Holmes 2013) defines phyloseq for storing tree with microbiome data. These packages are capable to present some of the data stored in the object on the tree. However, not all the associated data are supported. For example, species abundance stored in phyloseq object is not supported to be visualized using phyloseq package. These packages did not provide any utilities to integrate external data for tree visualization. metacoder is able to integrate external data to be used as visual characteristics (limited to size and color) to produce heat tree. However, phylogenetic tree layouts are not supported and branch lengthes will be ignored in metacoder. None of these packages support visualizing external data and align the plot to tree based on the tree structure.

The phylobase package support integrating external data (especially numerical matrix) to phylogenetic tree and visualize the data as bubble plot with the tree side by side. The plotTree scripts supports visualizing predefined datasets as heatmap or bar plots and aligns the plot to the tree. They only provide limited supports to visualize specific data types. The funtions provided by these packages or scripts are hard-coded to plot simple tree with specific panel (bubble plot or heatmap, etc) and there is no utilities provided to add further layer of annotations.

In contrast, ggtree has many unique features that cannot be found in all these implementations:

- 1. Integrating node/edge data to the tree can be mapped to visual characteristics of the tree or other datasets (Fig. S1).
- 2. Capable of parsing expression (math symbols or text formatting), emoji and image files (Fig. S1).
- 3. No predefined of input data types or how the data should be plotted in facet plot (Table S1).
- 4. Combining different geom functions to visualize associated data is supported (Fig. S4).
- 5. Visualizing different datasets on the same panel is supported (Fig. S8B and S9).
- 6. Data integrated by %<+% can be used in facet_plot (Fig. S6 and S9B).
- 7. Able to add further annotation to specific layers (Fig. S8B and S9).

8. Modular design by separating tree visualization, data integration (method 1) and graph alignment (method 2).

Modular design is a unique feature for ggtree to stand out from other packages. The tree can be visualized with data stored in tree object or external data linked by %<+% operator, and fully annotated with multiple layers of annotations (Fig. S1 and S4), before passing it to facet_plot. facet_plot can be called progresively to add multiple panels (Fig. S6) or multiple layers on the same panels (Fig. S4). This creates the possibility of plotting fully annotated tree with complex data panels that contains multiple graphic layers.

ggtree fits the R ecosystem and extends the abilities of integrating and presenting data with trees to existing phylogenetic packages. As demonstrated in this paper, we are now able to plot species abundance distribution with phyloseq object (Fig. S3), visualize numerical matrix stored in phylo4d object using other methods instead of just bubble plot (Fig. S9A) and integrate categorical values to color other data (Fig. S9), etc. All these cannot be easily done without ggtree. With ggtree, we are able to attach additional data to these tree objects using %<+% and align graph to tree using facet_plot. Integrating ggtree to existing workflows will definitely extends the abilities and broadens the applications to present phylogeny-associated data, especially for comparative studies.

NOTE: source code to produce this file can be obtained online¹⁰.

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¹⁰https://github.com/GuangchuangYu/plotting_tree_with_data

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