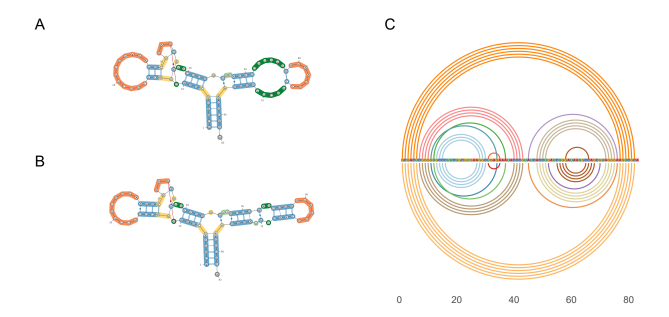
# ggmsa: a visual exploration tool for multiple sequence alignment and associated data

```
library(ggmsa)
library(ggplot2)
library(magick)
library(ggplotify)
library(aplot)
library(RColorBrewer)
library(Biostrings)
library(treeio)
library(ggtree)
```

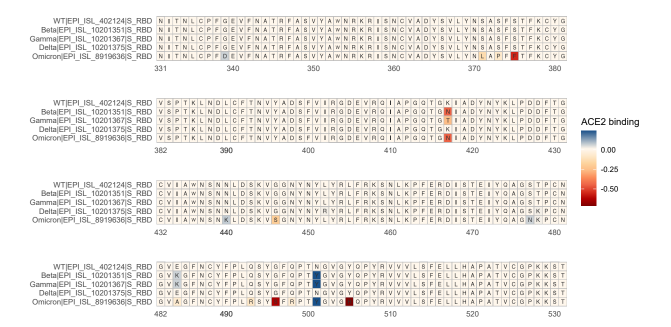
#### Reviewer1: Require2

```
tpp_seq <- "data/tpp_riboswitch.fasta"</pre>
arc_4NYG <- "data/riboswitch_thiamine.txt"</pre>
arc_4NYD <- "data/riboswitch_hypoxanthine.txt"</pre>
thiamine <- readSSfile(arc 4NYG, type = "Vienna" )</pre>
hypoxanthine <- readSSfile(arc_4NYD, type = "Vienna")</pre>
p <- ggmsa(tpp_seq,</pre>
           color = "Chemistry_NT",
           seq_name = F,
           show.legend = F,
           border = NA) +
    geom_helix(helix_data = list(known = hypoxanthine,
                                   predicted = thiamine)) +
    theme(axis.text.y = element_blank())
p1 <- image_read_pdf("data/bpRNA_PDB_590_ColorCodedStructures_4NYG.pdf")
p2 <- image_read_pdf("data/bpRNA_PDB_589_ColorCodedStructures_4NYD.pdf")</pre>
q1 <- as.ggplot(p1)
q2 <- as.ggplot(p2)
p_loop <- plot_list(gglist = list(q1, q2), ncol = 1)</pre>
pp <- plot_list(gglist = list(p_loop, p), ncol = 2, tag_levels = "A")</pre>
```



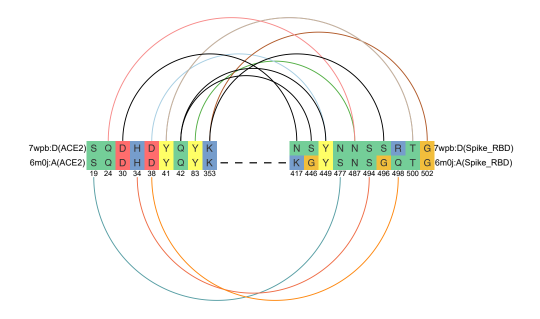
#### Reviewer1: Require2

```
colRD <- colorRampPalette(rev(brewer.pal(n = 9, name = "OrRd")))</pre>
colBU <- colorRampPalette(colors = rev(c("#185089","#FFF7EC")))</pre>
data <- "data/s_RBD.fasta"</pre>
mds <- read.csv("data/MDS.csv")</pre>
del <- c("expr_lib1", "expr_lib2",</pre>
          "expr_avg","bind_lib1",
          "bind_lib2")
mds <- mds[,!colnames(mds) %in% del]</pre>
tidymsa <- tidy_msa(data)</pre>
tidymsa <- assign_mds(tidymsa, mds)</pre>
tidymsa$position <- tidymsa$position + 330</pre>
p <- ggplot() +</pre>
    geom_msa(data = tidymsa,
              char_width = 0.5,
              mds = TRUE,
              seq_name = TRUE,
              show.legend = TRUE) +
    ggmsa:::theme_msa() +
    scale_fill_gradientn(name = "ACE2 binding",
                           colors = c(colRD(75), colBU(25))) +
    facet_msa(50)
p
```



```
#read sequences
x <- readAAStringSet("data/ACE2.fasta")</pre>
y <- readAAStringSet("data/Spike_RBD.fasta")</pre>
#read protein-protein position
inter1 <- read.csv("data/6m0j_inter.csv")</pre>
inter2 <- read.csv("data/7wpb_inter.csv")</pre>
t1 \leftarrow tidy_msa(x, start = 19)
t2 <- tidy_msa(y, start = 331)</pre>
t_merge \leftarrow merge_seq(t1, gap = 100, t2)
h1 <- tidy hdata(100, inter1, t1, t2)
h2 <- tidy_hdata(100, inter2, t1, t2)
ACE2 <- t_merge[t_merge$position %in% c(h1$i,h2$i),]
spike <- t_merge[t_merge$position %in% c(h1$j,h2$j),]</pre>
t1_reset <- reset_pos(ACE2)</pre>
t2_reset <- reset_pos(spike)</pre>
sim_merge <- merge_seq(previous_seq = t1_reset,</pre>
                          subsequent_seq = t2_reset,
                          gap = 5,
                         adjust_name = FALSE)
sim_h1 <- simplify_hdata(h1, sim_merge)</pre>
sim_h2 <- simplify_hdata(h2, sim_merge)</pre>
#break and label
b <- sim_merge[sim_merge$character != "-", "position"] %>% unique()
1 <- c(inter1$Res.no.1,inter1$Res.no..2,</pre>
         inter2$Res.no.1,inter2$Res.no..2) %>% unique
p_sim <- ggplot() +</pre>
```

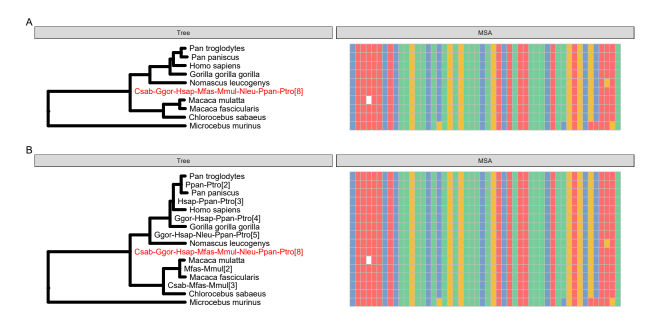
```
geom_msa(data = sim_merge,border = NA, char_width = 0.5, seq_name = F) +
    ggmsa:::theme_msa() +
    geom_helix(helix_data = list(known = sim_h1,
                             predicted = sim_h2),
               overlap = T) +
     geom_text(mapping = aes(x = b,
                             y = 0.25,
                             label = l[order(1)]),
               size = 3) +
     theme(axis.text.x = element_blank(),
           axis.text.y = element_blank()) +
     scale_y_discrete(labels = c("6m0j:A(ACE2)", "7wpb:D(ACE2)")) +
     geom_text(aes(x = c(27.2, 27.2),
                   y = c(1,2),
                   label = c("6m0j:A(Spike_RBD)",
                             "7wpb:D(Spike_RBD)")),
               size = 3.5) +
    geom_text(aes(x = c(-1.5, -1.5),
                  y = c(1,2),
                  label = c("6m0j:A(ACE2)",
                            "7wpb:D(ACE2)")),
              size = 3.5) + xlim(-2,29)
p_sim
```



### Reviewer2: Require1

```
x <- read.phyloxml("data/msa_phyloxml.xml")
p <- ggtree(x, size = 2)

p1 <- adjust_ally(p, node = 11, sub = FALSE)
tidy1 <- extract_seq(p1)</pre>
```



## Reviewer2: Require2

