Phylogenetic analysis of coronavirus sequences

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Contents

```
## Color palette per species
speciesPalette <- list(</pre>
 Human = "#880000",
 Bat = "#BBBBBB",
 Pangolin = "#00BB00",
 Camel = "#BB8800",
 Pig = "#FFBBBB",
 Civet = "#0000BB"
speciesPrefix <- c("Hu" = "Human",</pre>
              "Pn" = "Pangolin",
              "Pi" = "Pig",
              "Cv" = "Civet")
features <- c(
 "Recomb-reg-1",
 "Recomb-reg-2",
 "Recomb-reg-3",
 "CDS-ORF1ab",
 "CDS-ORF3a",
 "CDS-ORF6",
 "CDS-ORF7a",
 "CDS-ORF8",
 "CDS-N",
```

```
feature <- "S-gene"
## Define collections
collections <- c("around-CoV-2", "selected")</pre>
collection <- "around-CoV-2" # default for testing</pre>
## Outgroup per collection
outgroups <- list()</pre>
outgroups[["selected"]] <- c(</pre>
  "HuOC43",
  "PiPRCV",
  "HuTGEV",
  "PiSADS",
  "Hu229E",
  "HuNL63")
outgroups[["around-CoV-2"]] <- "BtBM48-31"</pre>
## Use GISAID data
useGISAID <- TRUE
if (useGISAID) {
  collections <- pasteO(collections, "-plus-GISAID")</pre>
  for (collection in names(outgroups)) {
    outgroups[[pasteO(collection, "-plus-GISAID")]] <- outgroups[[collection]]</pre>
  collection <- pasteO(collection, "-plus-GISAID")</pre>
dir <- vector()</pre>
dir["main"] <- ".."
dir["results"] <- file.path(dir["main"], "results")</pre>
```

```
dir <- vector()
dir["main"] <- ".."
dir["results"] <- file.path(dir["main"], "results")
dir["genomes"] <- file.path(dir["results"], "genome_phylogeny", "clustalw_alignments")

dir["R"] <- file.path(dir["main"], "scripts", "R")
# list.files(dir["R"])
source(file.path(dir["R"], "load_tree.R"))
source(file.path(dir["R"], "plot_my_tree.R"))</pre>
```

Phylogeny from full genomes

"CDS-ORF10")

We inferred a phylogeny of virus strains based ontheir full genomes.

A multiple alignment of genome sequences was performed with a progressive method (clustalw). The tree of virus strains was inferred with a maximum likelihood approach (phyml software).

```
#### Load and plot the genome tree ####
genomeTreeFile <- file.path(
    dir["genomes"],
    "coronavirus_selected-plus-GISAID_genomes_clustalw_gblocks.phy_phyml_tree.phb")

genomeTree <- loadTree(
    treeFile = genomeTreeFile,
    outgroup = outgroups[['selected']],
    rootNode = NULL,</pre>
```

Genome-based virus tree

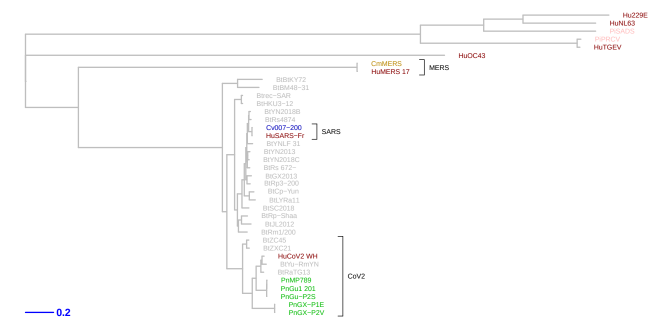


Figure 1: Genome tree of selected coronaviruses. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

Tree per feature

```
#### Load and plot feature-specific trees ####

## Define vectors to hold the results and enable tree comparisons
treeFiles <- vector()
treeData <- list()

## Define the path to the tre file
feature <- "S1"

for (feature in features) {
   message("\n\tReading tree for feature ", feature)
   prefix <- pasteO(feature, "_", collection)</pre>
```

```
treeFile <- file.path(</pre>
  dir["results"],
  prefix,
  pasteO(prefix, "_clustalw_gblocks.phy_phyml_tree_GTR.phb"))
treeFiles[feature] <- treeFile</pre>
treeData[[feature]] <- loadTree(</pre>
  treeFile = treeFiles[feature],
  outgroup = outgroups[[collection]],
  rootNode = NULL,
  nodesToRotate = NULL)
plotMyTree(treeData[[feature]],
           main = pasteO(feature, " tree"),
           scaleLength = 0.02,
           show.node.label = FALSE)
# cladelabels(genomeTree$tree, "CoV2", 46, cex = 0.7, orientation = "horizontal", offset = 5)
# cladelabels(genomeTree$tree, "MERS", 43, cex = 0.7, orientation = "horizontal", offset = 5)
# cladelabels(genomeTree$tree, "SARS", 69, cex = 0.7, orientation = "horizontal", offset = 5)
```

S1 tree

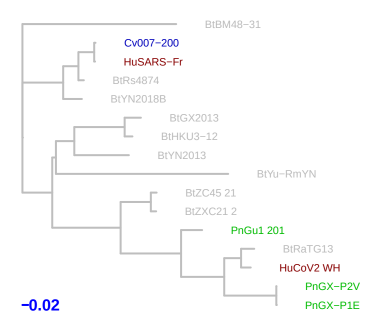


Figure 2: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

S2 tree

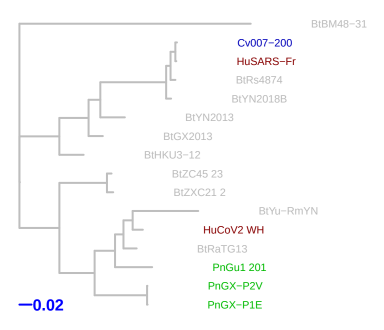


Figure 3: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

RBD tree

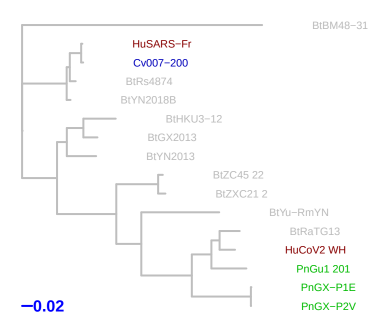


Figure 4: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

Recomb-reg-1 tree

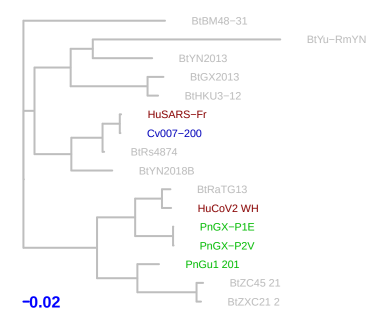


Figure 5: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

Recomb-reg-2 tree

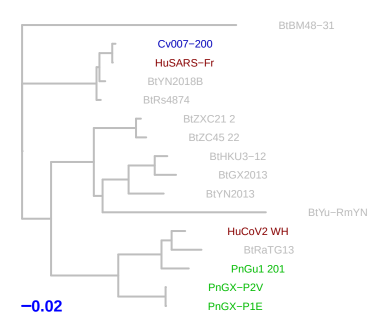


Figure 6: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

Recomb-reg-3 tree

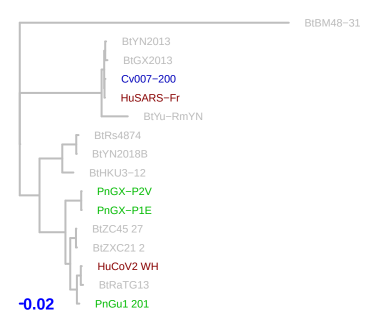


Figure 7: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF1ab tree

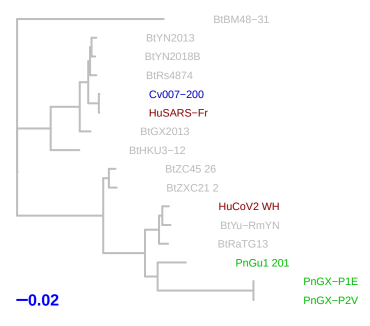


Figure 8: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-S tree

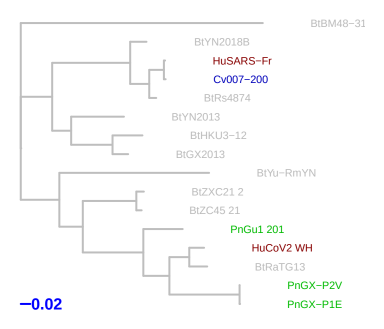


Figure 9: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF3a tree

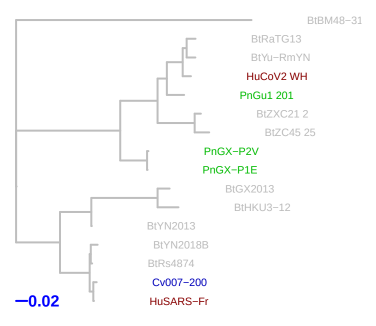


Figure 10: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-E tree

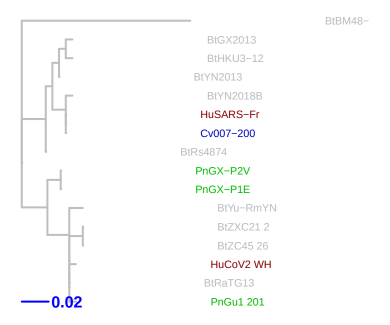


Figure 11: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-M tree

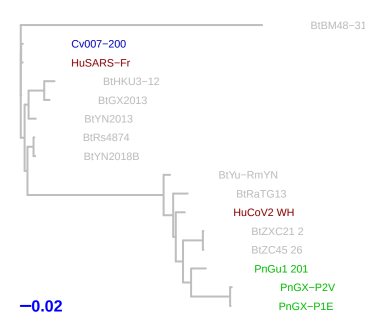


Figure 12: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF6 tree

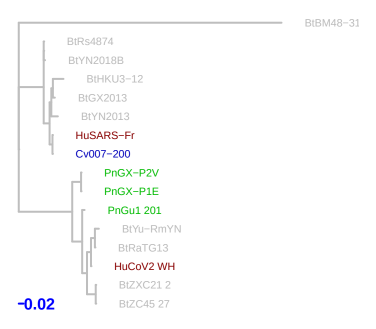


Figure 13: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF7a tree

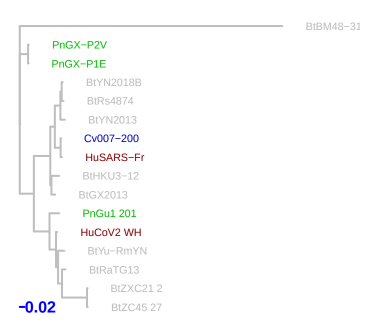


Figure 14: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF8 tree

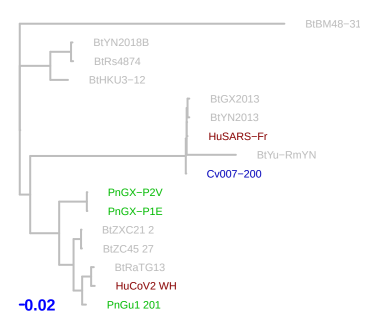


Figure 15: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-N tree

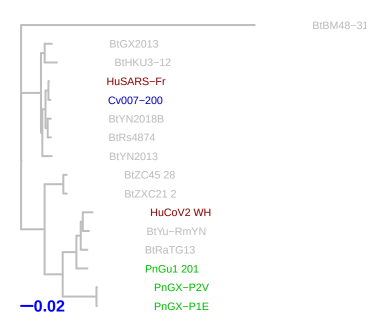


Figure 16: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).

CDS-ORF10 tree



Figure 17: Feature-specific tree. The tree was inferred by maximum likelihood apprroach (PhyML) based on a progressive multiple alignment (clustalw).