Supplementary Text S1: Assessing the completeness of Chlorophyta genomes

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```
library(here)
library(cogeqc)
library(tidyverse)
library(Herper)

set.seed(123) # for reproducibility
options(timeout = 6000) # to load files from the web
```

1 Overview

Here, we will use *cogeqc* to assess the completeness of Chlorophyta genomes available on Pico-PLAZA 3.0 (Van Bel et al. 2018) using Best Universal Single-Copy Orthologs (BUSCOs).

2 Managing external dependencies with virtual environments

Here, for convenience, we will install BUSCO in a Conda environment for use with *cogeqc*. For that, we will use the Bioconductor package *Herper*

Below, you can find the code to install miniconda in a directory of your choice (here, " \sim /Documents") and create a virtual environment containing a BUSCO installation.

```
# Path to where BUSCO will be installed and env name
my_miniconda <- file.path("~/Documents", "miniconda")
env <- "cogeqc_env"

# Create env named `cogeqc_env` with BUSCO in it
install_CondaTools(
    tools = "busco==5.3.0",
    env = env,
    channels = c("conda-forge", "bioconda"),
    pathToMiniConda = my_miniconda
)</pre>
```

3 Data acquisition

Now, we will load all genomes directly from PLAZA as DNAStringSet objects and export them to a single directory of FASTA files, so we can run BUSCO in batch mode.

```
# Links to Chlorophyta genomes from Pico-PLAZA 3.0
base_url <- "ftp://ftp.psb.ugent.be/pub/plaza/plaza_pico_03/Genomes/"
links <- paste0(
   base_url,
   c("mpu.fasta.gz", "mrcc299.fasta.gz", "olu.fasta.gz", "ome.fasta.gz",
   "orcc809.fasta.gz", "ota.fasta.gz", "bprrcc1105.fasta.gz",
   "cre.fasta.gz", "vca.fasta.gz", "cvu.fasta.gz", "acg.fasta.gz",
   "pse3.fasta.gz", "prcc4223.fasta.gz", "cnc64a.fasta.gz",</pre>
```

```
"hsp.fasta.gz", "apr.fasta.gz")
)
# Load all genomes
genomes <- lapply(links, Biostrings::readDNAStringSet)</pre>
names(genomes) <- basename(links)</pre>
# Write all genomes to a subdirectory of tempdir
genomes_path <- file.path(tempdir(), "genomes")</pre>
if(!dir.exists(genomes_path)) { fs::dir_create(genomes_path) }
write <- lapply(seq_along(genomes), function(x) {</pre>
    Biostrings::writeXStringSet(
        x = genomes[[x]],
        filepath = file.path(genomes_path, names(genomes)[x])
    return(NULL)
})
dir(genomes_path)
## [1] "acg.fasta.gz"
                              "apr.fasta.gz"
                                                      "bprrcc1105.fasta.gz"
## [4] "cnc64a.fasta.gz"
                              "cre.fasta.gz"
                                                      "cvu.fasta.gz"
## [7] "hsp.fasta.gz"
                              "mpu.fasta.gz"
                                                      "mrcc299.fasta.gz"
## [10] "olu.fasta.gz"
                              "ome.fasta.gz"
                                                     "orcc809.fasta.gz"
                              "prcc4223.fasta.gz"
                                                     "pse3.fasta.gz"
## [13] "ota.fasta.gz"
## [16] "vca.fasta.gz"
```

4 Running BUSCO

Now that all genomes are stored as FASTA files in /tmp/RtmpcuLMqJ/genomes, we can assess their completeness with BUSCO.

```
# See all possible lineage datasets
with_CondaEnv(
    env, list_busco_datasets(), my_miniconda
)

# Run BUSCO using chlorophyta_odb10 as the lineage data set
busco <- with_CondaEnv(
    env,
    run_busco(
        sequence = genomes_path,
        outlabel = "chlorophyta_busco",
        mode = "genome",
        lineage = "chlorophyta_odb10",
        outpath = tempdir(),
        download_path = tempdir()
    ),
    my_miniconda
)</pre>
```

```
# Read and parse the output
outdir <- file.path(tempdir(), "chlorophyta_busco")
busco_summary <- read_busco(outdir)
save(
    busco_summary,
    file = here::here("products", "result_files", "busco_summary.rda"),
    compress = "xz"
)</pre>
```

The parsed BUSCO output (as returned by read_busco()) looks like this:

```
load(here("products", "result_files", "busco_summary.rda"))
head(busco_summary)
## Class Frequency Lineage File
## 1 Complete_SC 94.1 chlorophyta_odb10 pse3.fasta.gz
## 2 Complete_SC 95.1 chlorophyta_odb10 cre.fasta.gz
## 3 Complete_SC 96.8 chlorophyta_odb10 olu.fasta.gz
## 4 Complete_SC 98.7 chlorophyta_odb10 mrcc299.fasta.gz
## 5 Complete_SC 91.8 chlorophyta_odb10 apr.fasta.gz
## 6 Complete_SC 86.4 chlorophyta_odb10 acg.fasta.gz
```

5 Visualizing summary statistics

Finally, let's visualize summary BUSCO stats:

```
# Manually create tree based on Pico-PLAZA's tree
c_branches <- function(b1, b2) {</pre>
    x <- paste0("(", b1, ",", b2, ")")
ostreococcus_root <- "((((Ostreococcus_lucimarinus, Ostreococcus_sp_RCC809), Ostreococcus_tauri), Ostreococcus_</pre>
micromonas <- "(Micromonas_pusilla_strain_CCMP1545, Micromonas_sp_RCC299)"</pre>
chlamydomonadales <- "(Volvox_carteri, Chlamydomonas_reinhardtii)"</pre>
picochlorum <- "(Picochlorum_sp_SENEW3, Picochlorum_RCC4223)"</pre>
chlorellales <- "((Helicosporidium_sp, Auxenochlorella_protothecoides), Chlorella_sp_NC64A)"</pre>
trebouxiophyceae <- c_branches(</pre>
    "(Coccomyxa_subellipsoidea_C-169, Asterochloris_sp_Cgr/DA1pho_v2)",
    c_branches(picochlorum, chlorellales)
)
chlo_tree <- c_branches(</pre>
    c_branches(
        ostreococcus_root, micromonas
    ),
    c_branches(
        chlamydomonadales, trebouxiophyceae
chlo_tree <- paste0(chlo_tree, ";")</pre>
```

```
# Read tree as a phylo object and clean species names
chlo_tree <- treeio::read.tree(text = chlo_tree)</pre>
chlo_tree$tip.label <- gsub("_", " ", chlo_tree$tip.label)</pre>
# Plot species tree and get species order from tree topology
p_{tree} < - plot_species_tree(chlo_tree, xlim = c(0, 12))
taxa_order <- rev(ggtree::get_taxa_name(p_tree))</pre>
# Plot BUSCO summary stats
p_busco <- busco_summary %>%
    mutate(File = str_replace_all(File, "\\.fasta.*", "")) %>%
    mutate(File = str_replace_all(
        File,
        c (
            "pse3" = "Picochlorum sp SENEW3",
            "cre" = "Chlamydomonas reinhardtii",
            "olu" = "Ostreococcus lucimarinus",
            "mrcc299" = "Micromonas sp RCC299",
            "apr" = "Auxenochlorella protothecoides",
            "acg" = "Asterochloris sp Cgr/DA1pho v2",
            "cvu" = "Coccomyxa subellipsoidea C-169",
            "bprrcc1105" = "Bathycoccus prasinos",
            "orcc809" = "Ostreococcus sp RCC809",
            "prcc4223" = "Picochlorum RCC4223",
            "ota" = "Ostreococcus tauri",
            "hsp" = "Helicosporidium sp",
            "mpu" = "Micromonas pusilla strain CCMP1545",
            "vca" = "Volvox carteri",
            "ome" = "Ostreococcus mediterraneus",
            "cnc64a" = "Chlorella sp NC64A"
        )
    )) %>%
    mutate(File = factor(File, taxa_order)) %>%
    plot_busco() +
    theme(axis.text.y = element_blank()) +
    labs(y = "")
# Combining phylogeny with BUSCO plot
combined <- patchwork::wrap_plots(p_tree, p_busco)</pre>
combined
```

Except for *Helicosporidium sp.*, Chlorophyta genomes on Pico-PLAZA 3.0 have a high quality, as denoted by their high completeness.

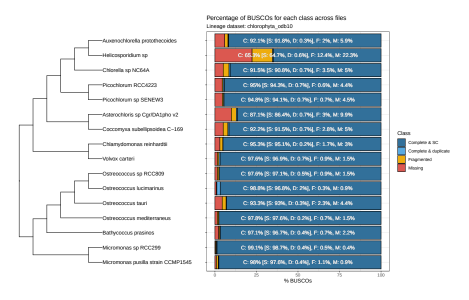


Figure 1: BUSCO scores for Chlorophyta genomes on Pico-PLAZA 3.0.

Session info

This document was created under the following conditions:

```
sessioninfo::session_info()
## - Session info -----
## setting value
## version R version 4.2.2 Patched (2022-11-10 r83330)
## os
         Ubuntu 20.04.5 LTS
## system x86_64, linux-gnu
         X11
## ui
## language (EN)
## collate en_US.UTF-8
## ctype en_US.UTF-8
## tz
         Europe/Brussels
         2023-02-02
## date
## pandoc 2.19.2 @ /usr/lib/rstudio/resources/app/bin/quarto/bin/tools/ (via rmarkdown)
##
## package
                * version date (UTC) lib source
                  5.6-2 2022-03-02 [1] CRAN (R 4.2.0)
0.1.8 2022-10-09 [1] CRAN (R 4.2.1)
## ape
## aplot
                  0.2.1 2019-03-21 [1] CRAN (R 4.2.0)
## assertthat
                  1.4.1 2021-12-13 [1] CRAN (R 4.2.0)
0.4.0 2021-06-01 [1] CRAN (R 4.2.2)
## backports
## beeswarm
## BiocGenerics
                  0.42.0 2022-04-26 [1] Bioconductor
## BiocManager
                  1.30.18 2022-05-18 [1] CRAN (R 4.2.0)
## BiocStyle
                * 2.25.0 2022-06-15 [1] Github (Bioconductor/BiocStyle@7150c28)
                  2.64.1
## Biostrings
                            2022-08-18 [1] Bioconductor
                  1.0-7 2021-04-24 [1] CRAN (R 4.2.0)
## bitops
```

```
0.29
## bookdown
                                  2022-09-12 [1] CRAN (R 4.2.1)
                       1.0.1 2022-08-29 [1] CRAN (R 4.2.1)
## broom
                                  2016-07-27 [1] CRAN (R 4.2.0)
## cellranger
                       1.1.0
## cli
                       3.4.1 2022-09-23 [1] CRAN (R 4.2.1)
## cogegc
                    * 1.3.1 2023-01-24 [1] Bioconductor
                    2.0-3 2022-02-21 [1] CRAN (R 4.2.0)
1.5.2 2022-09-29 [1] CRAN (R 4.2.1)
## colorspace
## crayon
## DBI
                      1.1.3 2022-06-18 [1] CRAN (R 4.2.0)
## dbplyr
                      2.2.1 2022-06-27 [1] CRAN (R 4.2.1)
                      0.6.29 2021-12-01 [1] CRAN (R 4.2.0)
## digest
                   * 1.0.10 2022-09-01 [1] CRAN (R 4.2.1)
## dplyr
                    0.3.2 2021-04-29 [1] CRAN (R 4.2.0)
## ellipsis
                     0.17 2022-10-07 [1] CRAN (R 4.2.1)
1.0.3 2022-03-24 [1] CRAN (R 4.2.0)
2.1.1 2022-07-06 [1] CRAN (R 4.2.1)
## evaluate
## fansi
## farver
## fastmap
                      1.1.0 2021-01-25 [1] CRAN (R 4.2.0)
                   * 0.5.2 2022-08-19 [1] CRAN (R 4.2.1)
1.5.2 2021-12-08 [1] CRAN (R 4.2.0)
1.2.1 2022-09-08 [1] CRAN (R 4.2.1)
## forcats
## fs
## gargle
## generics
                      0.1.3 2022-07-05 [1] CRAN (R 4.2.1)
## GenomeInfoDb 1.32.4 2022-09-06 [1] Bioconductor
## GenomeInfoDbData 1.2.8 2022-05-06 [1] Bioconductor
## ggbeeswarm 0.7.1 2022-12-16 [1] CRAN (R 4.2.2)
## ggfun
                      0.0.8 2022-11-07 [1] CRAN (R 4.2.1)
                   * 3.4.0 2022-11-04 [1] CRAN (R 4.2.1)
0.1.0 2021-09-02 [1] CRAN (R 4.2.0)
## ggplot2
## ggplotify
                     3.7.1.001 2022-11-10 [1] Github (YuLab-SMU/ggtree@b7ef83e)
## ggtree
## glue
                      1.6.2 2022-02-24 [1] CRAN (R 4.2.0)
## googledrive 2.0.0 2021-07-08 [1] CRAN (R 4.2.0)

## googlesheets4 1.0.1 2022-08-13 [1] CRAN (R 4.2.1)

## gridGraphics 0.5-1 2020-12-13 [1] CRAN (R 4.2.0)
                   0.3.1 2022-09-01 [1] CRAN (R 4.2.1)
2.5.1 2022-08-22 [1] CRAN (R 4.2.1)
* 1.0.1 2020-12-13 [1] CRAN (R 4.2.0)
## gtable
## haven
## here
                  * 1.1.2 2022-05-18 [1] Github (RockefellerUniversity/Herper@5bceeb4)
## Herper
                  1.1.2 2022-08-19 [1] CRAN (R 4.2.1)
0.5.3 2022-07-18 [1] CRAN (R 4.2.1)
## hms
## htmltools
                      1.4.4 2022-08-17 [1] CRAN (R 4.2.1)
## httr
## igraph
                      1.3.5 2022-09-22 [1] CRAN (R 4.2.1)
                      2.30.1 2022-08-18 [1] Bioconductor
## IRanges
                     1.8.3
## jsonlite
                                  2022-10-21 [1] CRAN (R 4.2.1)
## knitr
                      1.40 2022-08-24 [1] CRAN (R 4.2.1)
                      0.4.2 2020-10-20 [1] CRAN (R 4.2.0)
## labeling
                      0.20-45 2021-09-22 [1] CRAN (R 4.2.0)
## lattice
                      0.2.2
                                  2019-03-15 [1] CRAN (R 4.2.0)
##
   lazyeval
## lifecycle
                      1.0.3 2022-10-07 [1] CRAN (R 4.2.1)
                      1.8.0 2021-10-07 [1] CRAN (R 4.2.0)
## lubridate
                      2.0.3 2022-03-30 [1] CRAN (R 4.2.0)
1.5-1 2022-09-13 [1] CRAN (R 4.2.1)
## magrittr
## Matrix
                      0.1.9 2022-08-19 [1] CRAN (R 4.2.1)
## modelr
               0.5.0 2018-06-12 [1] CRAN (R 4.2.0)
## munsell
```

```
## nlme
                                 3.1-160 2022-10-10 [1] CRAN (R 4.2.1)
                         3.1-160 2022-10-10 [1] CRAN (R 4.2.1)
1.1.2 2022-08-19 [1] CRAN (R 4.2.1)
1.8.1 2022-08-19 [1] CRAN (R 4.2.1)
2.0.3 2019-09-22 [1] CRAN (R 4.2.0)
1.8.7 2022-03-24 [1] CRAN (R 4.2.0)
0.1-7 2013-12-03 [1] CRAN (R 4.2.0)
* 0.3.5 2022-10-06 [1] CRAN (R 4.2.1)
## patchwork
 ## pillar
 ## pkgconfig
 ## plyr
 ## png
 ## purrr
## R6
                              2.5.1 2021-08-19 [1] CRAN (R 4.2.0)
                                 1.0.9 2022-07-08 [1] CRAN (R 4.2.1)
 ## Rcpp
                               1.98-1.9 2022-10-03 [1] CRAN (R 4.2.1)
## RCurl 1.98-1.9 2022-10-03 [1] CRAN (R 4.2.1)
## readr * 2.1.3 2022-10-01 [1] CRAN (R 4.2.1)
## readxl 1.4.1 2022-08-17 [1] CRAN (R 4.2.1)
## reprex 2.0.2 2022-08-17 [1] CRAN (R 4.2.1)
## reshape2 1.4.4 2020-04-09 [1] CRAN (R 4.2.0)
## reticulate * 1.26 2022-08-31 [1] CRAN (R 4.2.1)
 ## RCurl
## rlang 1.0.6 2022-09-24 [1] CRAN (R 4.2.1)
## rmarkdown 2.17 2022-10-07 [1] CRAN (R 4.2.1)
## rprojroot 2.0.3 2022-04-02 [1] CRAN (R 4.2.0)
## rstudioapi 0.14 2022-08-22 [1] CRAN (R 4.2.1)
## rvest 1.0.3 2022-08-19 [1] CRAN (R 4.2.1)
## S4Vectors 0.34.0 2022-04-26 [1] Bioconductor
## scales 1.2.1 2022-08-20 [1] CRAN (R 4.2.1)
 ## rjson
                              0.2.21 2022-01-09 [1] CRAN (R 4.2.0)
                                 1.2.1 2022-08-20 [1] CRAN (R 4.2.1)
 ## sessioninfo 1.2.2 2021-12-06 [1] CRAN (R 4.2.0)
1.23.0 2022-11-10 [1] Github (Guangcl 0.3.0 2022-03-28 [1] CRAN (R 4.2.0) 1.2.2 2021-07-24 [1] CRAN (R 4.2.0) 0.5.0 2022-10-22 [1] CRAN (R 4.2.1) 0.4.5 2017-03-22 [1] CRAN (R 4.2.1) 2.5.0 2022-03-03 [1] CRAN (R 4.2.0) 0.33 2022-09-12 [1] CRAN (R 4.2.0)
                              1.23.0 2022-11-10 [1] Github (GuangchuangYu/treeio@db85803)
 ## treeio
 ## tzdb
 ## utf8
 ## vctrs
 ## vipor
 ## withr
##
 ## [1] /home/faalm/R/x86_64-pc-linux-gnu-library/4.2
 ## [2] /usr/local/lib/R/site-library
 ## [3] /usr/lib/R/site-library
 ## [4] /usr/lib/R/library
 ##
 ## -----
```

References

Van Bel, Michiel, Tim Diels, Emmelien Vancaester, Lukasz Kreft, Alexander Botzki, Yves Van de Peer, Frederik Coppens, and Klaas Vandepoele. 2018. "PLAZA 4.0: An Integrative Resource for Functional, Evolutionary and Comparative Plant Genomics." *Nucleic Acids Research* 46 (D1): D1190–96.