# Chlorophyll analysis

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#### Import data

```
# Define function to read in chl data
read_chl <- function(file) {</pre>
  chl_data <- read_csv(file, skip = 24, n_max = 24) %>%
    select(-1) %>%
    magrittr::set_colnames(c("row", 1:12, "wavelength")) %>%
    fill(row) %>%
    gather("col", "absorbance", -wavelength, -row) %>%
    unite("well", c(row, col), sep = "")
}
# List chlorophyll data files
chl_path <- "baseline_TPC_physio/data/TPC_chlorophyll/"</pre>
                                                                                                   # Path t
all_chl_files <- list.files(path = chl_path, pattern = "*.csv")</pre>
                                                                           # List all files in directory
chl_platemaps <- list.files(path = chl_path, pattern = "platemap")</pre>
                                                                          # List platemap files
chl_data_files <- setdiff(all_chl_files, chl_platemaps)</pre>
                                                                           # List absorbance data files
# Read in all files into tibble
df <- tibble(file = chl_data_files) %>%
  mutate(platemap = map(file, ~ read_csv(paste0(chl_path, tools::file_path_sans_ext(.), "_platemap.csv"
         chl_data = map(file, ~ read_chl(pasteO(chl_path, .))))
# Merge platemap and data for each plate
df <- df %>%
  mutate(merged = map2(platemap, chl_data, ~ right_join(.x, .y)))
```

## Calculate chlorophyll concentrations

```
df <- df %>%
  group_by(file) %>%
  mutate(blank750 = `750`[fragment_ID == "BK"]) %>%
  ungroup() %>%
  mutate(adj630 = `630` - blank750,
         adj663 = `663` - blank750)
# calculate chla and chlc2 values based on equations from Jeffrey and Humphrey 1975
# units µq/ml
#path length adjustment = 0.6
df <- df %>%
  mutate(chla.ug.ml = (11.43 * adj663)/0.6 - (0.64 * adj630)/0.6,
        chlc2.ug.ml = (27.09 * adj630)/0.6 - (3.63 * adj663)/0.6)
#previous, with no pathlength adjustment
#df <- df %>%
  \#mutate(chla.ug.ml = (11.43 * adj663) - (0.64 * adj630),
        \#chlc2.uq.ml = (27.09 * adj630) - (3.63 * adj663))
```

#### Normalize to surface area

```
# Load homogenate volume
homog.vol <- read_csv("baseline_TPC_physio/data/TPC_homogenate_vols/homogenate_vols.csv") %>%
  select(fragment_ID, homog_vol_ml)
chl <- full_join(df, homog.vol)</pre>
# Load surface area
sa <- read_csv("baseline_TPC_physio/output/surface.area.calc.csv")</pre>
chl <- full_join(chl, sa)</pre>
# Multiply chlorophyll by the homogenate volume and divide by surface area
chl <- chl %>%
 mutate(chla.ug.cm2 = chla.ug.ml * homog_vol_ml / surface.area.cm2,
         chlc2.ug.cm2 = chlc2.ug.ml * homog_vol_ml / surface.area.cm2)
# remove blanks and NAs
chl <- filter(chl, !fragment_ID %in% c("NA", "BK"))</pre>
# write chlorophyll data to file
chl %>%
 select(fragment_ID, chla.ug.cm2, chlc2.ug.cm2) %>%
 mutate(timepoint="May2022")%>%
 filter(!is.na(chla.ug.cm2))%>%
 filter(!is.na(chlc2.ug.cm2))%>%
  write_csv(path = "baseline_TPC_physio/output/chlorophyll.csv")
```

### Plot results by species and site

```
# Join with sample metadata
meta <- read_csv("baseline_TPC_physio/metadata_POC_TPC.csv")</pre>
chl <- right_join(chl, meta, by = "fragment_ID")</pre>
# Plot chlorophyll a
chl %>%
 ggplot(aes(x = site , y = chla.ug.cm2, color = species)) +
  #facet_wrap(~species) +
  labs(x = "", y = "chlorophyll a (\mug/cm2)") +
  geom_jitter(width = 0.1) +
                                                                            # Plot all points
  stat_summary(fun.data = mean_cl_normal, fun.args = list(mult = 1),
                                                                            # Plot standard error
               geom = "errorbar", color = "black", width = 0.5) +
  stat_summary(fun.y = mean, geom = "point", color = "black")
                                                                            # Plot mean
   6 -
   5 -
chlorophyll a (µg/cm2)
                                                                               species
                                                                                   Pocillopora
   2 -
                                     Manava
# Plot chlorophyll c2
chl %>%
  ggplot(aes(x = site, y = chlc2.ug.cm2, color = species)) +
  #facet_wrap(~species) +
  labs(x = "", y = "chlorophyll c2 (\mug/cm2)") +
  geom_jitter(width = 0.1) +
                                                                            # Plot all points
  stat_summary(fun.data = mean_cl_normal, fun.args = list(mult = 1),
                                                                            # Plot standard error
               geom = "errorbar", color = "black", width = 0.5) +
  stat_summary(fun.y = mean, geom = "point", color = "black")
                                                                            # Plot mean
```

