# Introduction to using gcplyr

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### Getting started

gcplyr is a package that implements a number of functions to make it easier to import, manipulate, and analyze microbial growth from data collected in multiwell plate readers ("growth curves"). Without gcplyr, importing and analyzing plate reader data can be a complicated process that has to be tailored for each experiment, requiring many lines of code. With gcplyr many of those steps are now just a single line of code.

This document gives an introduction of how to use gcplyr for each step of a growth curve analysis.

To get started, you need your growth curve data file saved to your computer (.csv, .xls, .xlsx, or any other format that can be read by read.table).

Users often want to combine their data with some information on the experimental design of their plate(s). You can save this information into a tabular file as well, or you can just keep it handy to enter directly in R (see vignette("gc03\_incorporate\_designs")).

Let's get started by loading gcplyr. We're also going to load a couple other packages we'll need.

```
library(gcplyr)
library(dplyr)
library(ggplot2)
```

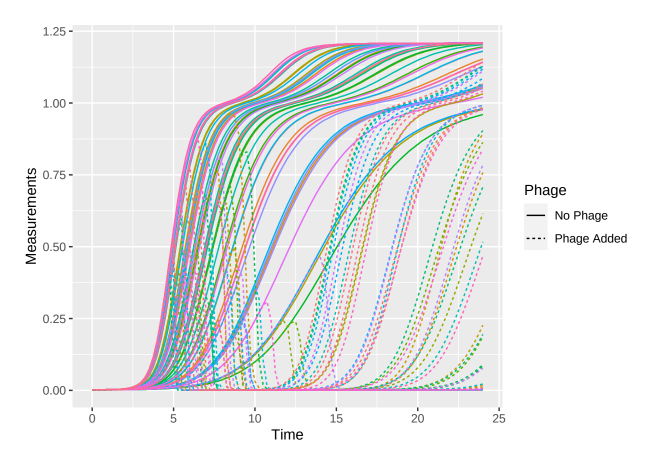
## A quick demo of gcplyr

Before digging into the details, here's a simple demonstration of what a final gcplyr script can look like. This script:

- 2. combines it with design files created by the user
- 3. calculates the lag time, maximum growth rate, maximum density, and area-under-the-curve

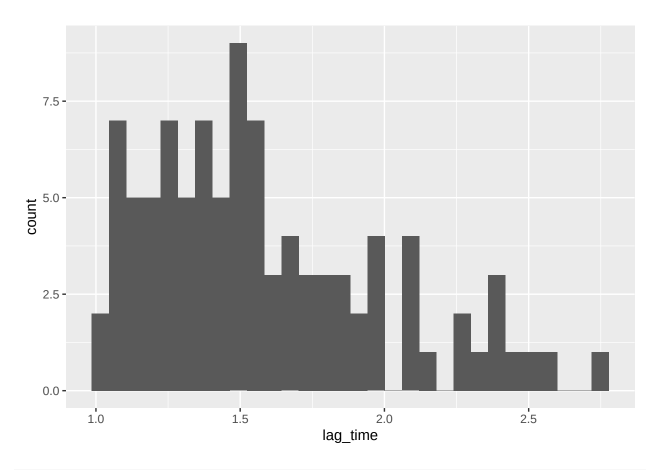
Don't worry about understanding all the details of how the code works right now. Each of these steps is explained in depth in later articles.

```
#For the purposes of this demo, we have to create our example data and
# design files. Normally, the data would be created by a plate reader, and
# the design would be created by you, the user
\#Generate\ our\ example\ data\ file,\ widedata.csv
make_example(vignette = 1, example = 1)
#> Files have been written
#> [1] "./widedata.csv"
#Generate our example design files, Bacteria strain.csv and Phage.csv
make_example(vignette = 1, example = 2)
#> Files have been written
#> [1] "./Bacteria_strain.csv" "./Phage.csv"
# Read in our data
data wide <- read wides(files = "widedata.csv")</pre>
# Transform our data to be tidy-shaped
data_tidy <-
  trans_wide_to_tidy(wides = data_wide, id_cols = c("file", "Time"))
# Convert our time into hours
data_tidy$Time <- as.numeric(data_tidy$Time)/3600</pre>
# Import our designs
designs <- import_blockdesigns(files = c("Bacteria_strain.csv", "Phage.csv"))</pre>
# Merge our designs and data
data_merged <- merge_dfs(data_tidy, designs)</pre>
#> Joining, by = "Well"
# Plot the data
ggplot(data = data merged,
       aes(x = Time, y = Measurements, color = Well)) +
  geom_line(aes(lty = Phage)) +
 guides(color = "none")
```

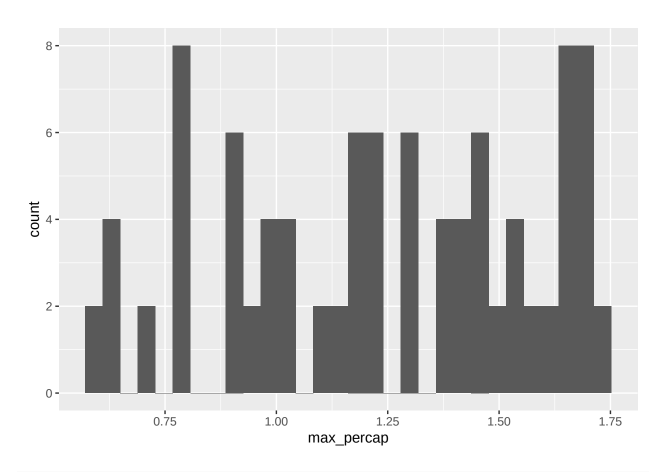


```
# Voila! 8 lines of code and all your data is imported & plotted!
# Calculate the per-capita growth rate over time in each well
data_merged <- mutate(</pre>
  group_by(data_merged, Well),
 percap_deriv = calc_deriv(y = Measurements, x = Time, percapita = TRUE,
                            blank = 0, window_width_n = 5))
# Calculate four common metrics of bacterial growth:
# the lag time, saving it to a column named lag_time
# the maximum growth rate, saving it to a column named max_percap
# the maximum density, saving it to a column named max_dens
# the area-under-the-curve, saving it to a column named 'auc'
data_sum <- summarize(</pre>
  group_by(data_merged, Well, Bacteria_strain, Phage),
 lag_time = lag_time(x = Time, y = Measurements, deriv = percap_deriv),
 max_percap = max(percap_deriv, na.rm = TRUE),
 max_dens = max(Measurements),
 auc = auc(y = Measurements, x = as.numeric(Time)))
#> `summarise()` has grouped output by 'Well', 'Bacteria_strain'. You can override
#> using the `.groups` argument.
# Print some of the values
head(data sum)
#> # A tibble: 6 x 7
```

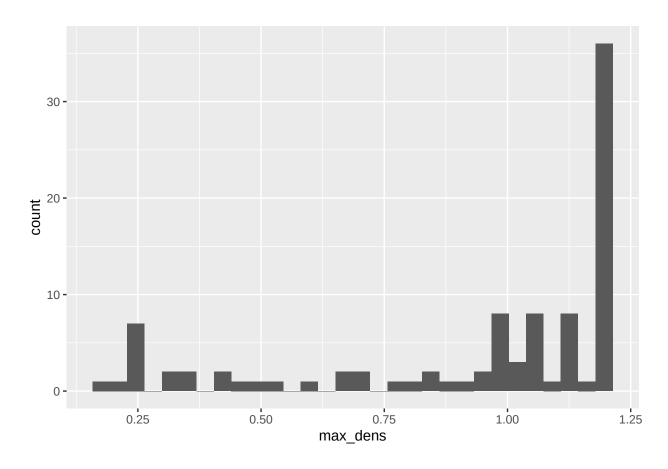
```
#> # Groups: Well, Bacteria_strain [6]
#> Well Bacteria_strain Phage lag_time max_percap max_dens
#> <chr> <chr>
                         <chr>
                                       <dbl>
                                                 <dbl>
                                                          <dbl> <dbl>
                        No Phage
                                        2.11
                                                 1.00
#> 1 A1
          Strain 1
                                                          1.18 15.9
#> 2 A10 Strain 4
                       Phage Added
                                        1.19
                                                1.43
                                                          0.984 5.57
#> 3 A11 Strain 5
                        Phage Added
                                        1.20
                                                1.47
                                                          0.984 5.99
#> 4 A12 Strain 6
                        Phage Added
                                        1.54
                                                 0.789
                                                          0.19 0.395
          Strain 2
#> 5 A2
                         No Phage
                                        1.74
                                                 1.31
                                                          1.21 19.3
                                        2.14
#> 6 A3 Strain 3
                         No Phage
                                                 0.915
                                                          1.15 15.1
# Plot the results for each of the metrics
ggplot(data = data_sum, aes(x = lag_time)) +
 geom_histogram()
#> `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



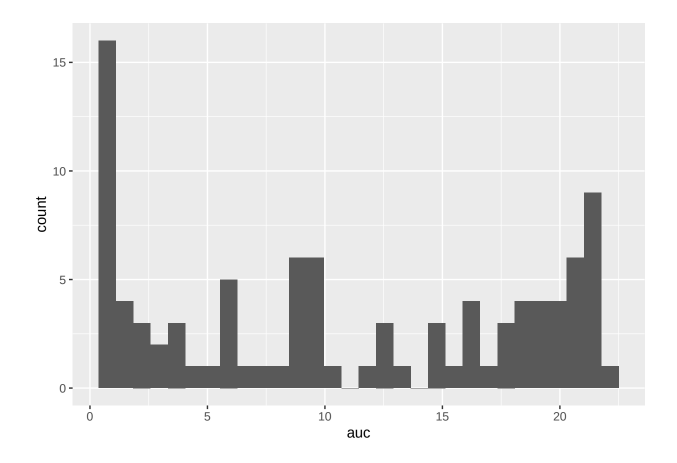
```
ggplot(data = data_sum, aes(x = max_percap)) +
  geom_histogram()
#> `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



```
ggplot(data = data_sum, aes(x = max_dens)) +
  geom_histogram()
#> `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



```
ggplot(data = data_sum, aes(x = auc)) +
  geom_histogram()
#> `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



### What's next?

Generally, working with gcplyr will follow a number of steps, each of which is likely to be only one or a few lines of code in your final script. We've explained each of these steps in a page linked below. To start, we'll learn how to import our data into R and transform it into a convenient format.

- 1. Introduction: vignette("gc01\_gcplyr")
- 2. Importing and transforming data: vignette("gc02\_import\_reshape")
- 3. Incorporating experimental designs: vignette("gc03 incorporate designs")
- 4. Pre-processing and plotting your data: vignette("gc04\_preprocess\_plot")
- 5. Processing your data: vignette("gc05\_process")
- 6. Analyzing your data: vignette("gc06\_analyze")
- 7. Dealing with noise: vignette("gc07\_noise")
- 8. Statistics, merging other data, and other resources: vignette("gc08\_conclusion")
- 9. Working with multiple plates: vignette("gc09\_multiple\_plates")