# Notes for foundations of ggplot2

#### Eric Scott

```
library(palmerpenguins)
theme set(theme bw())
```

i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all

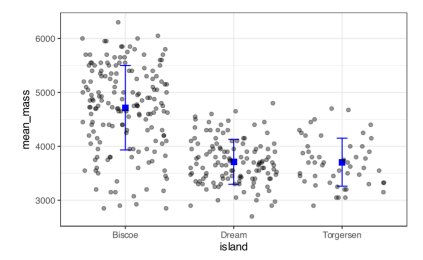
#### Data

This is the plot we're trying to re-create:

conflicts to become errors

\* dplyr::filter() masks stats::filter()
\* dplyr::lag() masks stats::lag()

```
Warning: Removed 2 rows containing missing values or values outside the scale range (`geom_point()`).
```



What geom can we use for error bars? Type geom\_ and hit tab to see a list of all possible geom\_ functions. geom\_errorbar() sounds about right! What data do we need to plot an error bar? View the help file with ?geom\_errorbar and check the "Aesthetics" section. Looks like we need x (island), ymin, and ymax.

Let's start by summarizing the data to calculate a mean and standard deviation for each island.

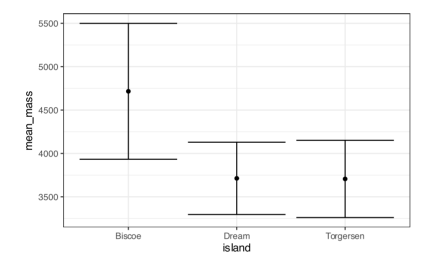
```
peng_summary <-
penguins |>
group_by(island) |> #for each island...
summarize(
   mean_mass = mean(body_mass_g, na.rm = TRUE),
   sd = sd(body_mass_g, na.rm = TRUE)
)
peng_summary
```

Then let's get values for the lower and upper parts of the error bars.

```
peng_summary <-
peng_summary |>
mutate(lower = mean_mass - sd, upper = mean_mass + sd)
peng_summary
```

Now we can plot the error bars and means.

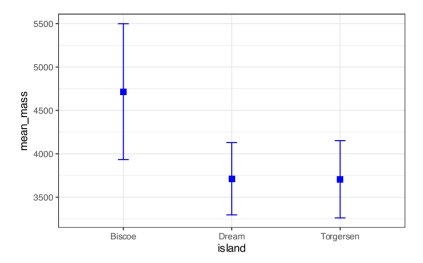
```
ggplot(peng_summary, aes(x = island)) +
    #mean:
    geom_point(aes(y = mean_mass)) +
    #sd:
    geom_errorbar(aes(ymin = lower, ymax = upper))
```



Let's do some tweaking to make this look more appealing

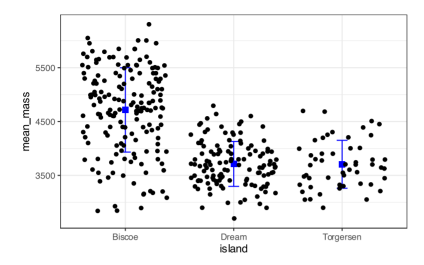
```
ggplot(peng_summary, aes(x = island)) +
    #mean:
    geom_point(
        aes(y = mean_mass),
        shape = "square",
        color = "blue",
        size = 2.5
)+
    #sd:
    geom_errorbar(
        data = peng_summary,
        aes(ymin = lower, ymax = upper),
        width = 0.1,
```

```
color = "blue"
)
```



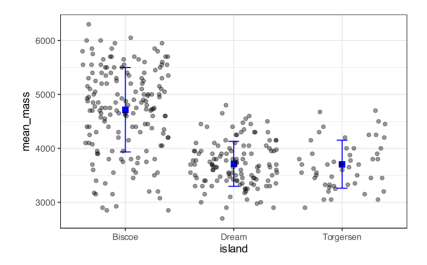
Now we can add the raw data using <code>geom\_jitter()</code> by overriding the data argument.

```
ggplot(peng_summary, aes(x = island)) +
 #mean:
 geom_point(
   aes(y = mean_mass),
   shape = "square",
   color = "blue",
   size = 2.5
 )+
 #sd:
 geom_errorbar(
   data = peng_summary,
   aes(ymin = lower, ymax = upper),
   width = 0.1,
   color = "blue"
 ) +
 #add raw data:
 geom jitter(
   data = penguins, #override data to use penguins instead of peng_summary
   aes(y = body_mass_g),
 )
```



And finally we can do some tweaking of the jitter layer

```
ggplot(peng_summary, aes(x = island)) +
 #mean:
 geom_point(
   aes(y = mean_mass),
   shape = "square",
   color = "blue",
   size = 2.5
 )+
 #sd:
 geom_errorbar(
   data = peng_summary,
   aes(ymin = lower, ymax = upper),
   width = 0.1,
    color = "blue"
 ) +
 #add raw data:
 geom_jitter(
   data = penguins, #override data to use penguins instead of peng_summary
   aes(y = body_mass_g),
   alpha = 0.4, #add transparency
    height = 0 #don't jitter vertically, only horizontally
 )
```



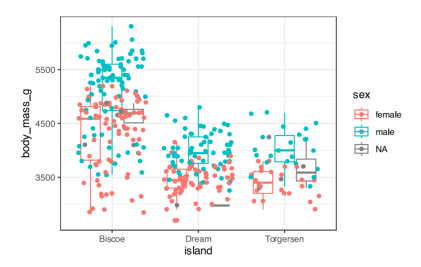
### **Aesthetics**

Let's use a boxplot on top of the jitter plot and have the points colored by sex but not the box plots

When color = sex is in the aes() call in ggplot(), this aesthetic mapping is inherited by all geoms.

```
ggplot(penguins, aes(x = island, y = body_mass_g, color = sex)) +
  geom_boxplot() +
  geom_jitter()
```

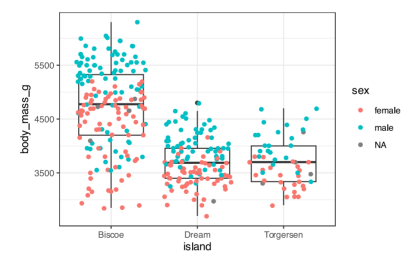
```
Warning: Removed 2 rows containing non-finite outside the scale range
(`stat_boxplot()`).
```



If we want to map sex to color only for the jitter layer, we can remove it from ggplot() and add it to geom\_jitter()

```
ggplot(penguins, aes(x = island, y = body_mass_g)) +
  geom_boxplot() +
  geom_jitter(aes(color = sex))
```

Warning: Removed 2 rows containing non-finite outside the scale range
 (`stat\_boxplot()`).

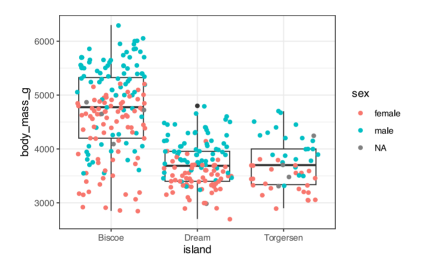


Or, we can use inherit.aes = FALSE and specify *all* the aesthetic mappings for the boxplot layer.

```
ggplot(penguins, aes(x = island, y = body_mass_g, color = sex)) +
geom_boxplot(aes(x = island, y = body_mass_g), inherit.aes = FALSE) +
geom_jitter()
```

Warning: Removed 2 rows containing non-finite outside the scale range
(`stat\_boxplot()`).

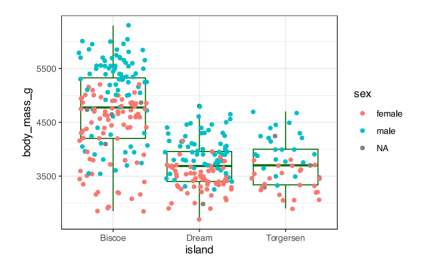
Warning: Removed 2 rows containing missing values or values outside the scale range (`geom point()`).



If you set aesthetic mappings to constants, it overrides the mappings to data.

```
ggplot(penguins, aes(x = island, y = body_mass_g, color = sex)) +
  geom_boxplot(color = "darkgreen") +
  geom_jitter()
```

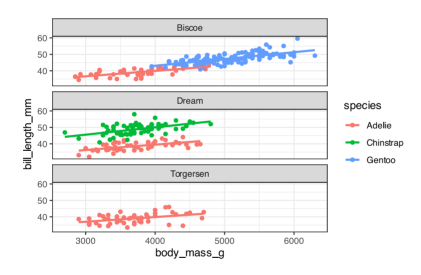
Warning: Removed 2 rows containing non-finite outside the scale range (`stat\_boxplot()`).



# **Scales**

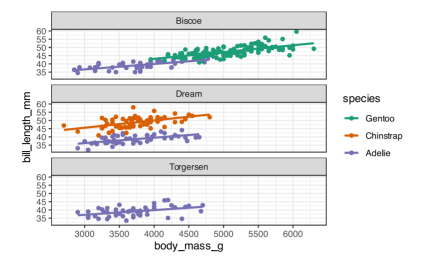
Here's the original plot, saved as p

 $geom_smooth()$  using formula = 'y ~ x'



And here's what it's going to end up like after modifying scales:

`geom\_smooth()` using formula = 'y  $\sim$  x'



First, let's address the color scale. Two changes need to happen: custom colors, and a re-ordering of the species in the legend. scale\_color\_manual() can take care of both.

We can supply whatever colors we want with a named vector where the names correspond to levels of the species variable that is mapped to color.

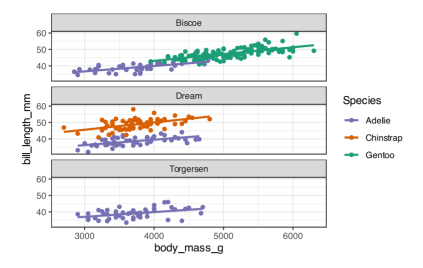
```
my_cols <- c(
   "Adelie" = "#7570b3",
   "Chinstrap" = "#d95f02",
   "Gentoo" = "#1b9e77"
)</pre>
```

These are hex-codes, but you can also used named colors in R.

Supply that named vector to the values argument.

```
p +
    scale_color_manual(
    name = "Species",
    values = my_cols
)
```

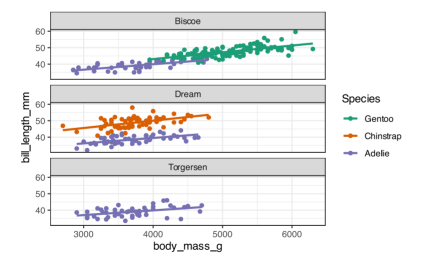
```
geom_smooth() using formula = 'y ~ x'
```



The re-ordering happens with the breaks argument like so:

```
p_new <- p +
    scale_color_manual(
    name = "Species",
    values = my_cols,
    breaks = c("Gentoo", "Chinstrap", "Adelie")
)
p_new</pre>
```

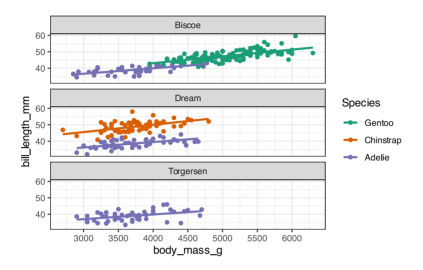




Now we can move on to the x and y axes. For the x-axis, let's increase the number of breaks to *about* 10.

```
p_new +
    scale_x_continuous(n.breaks = 10)
```

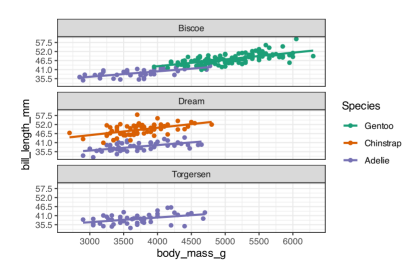
```
geom_smooth() using formula = 'y ~ x'
```



And we can supply *exact* breaks to the y-axis.

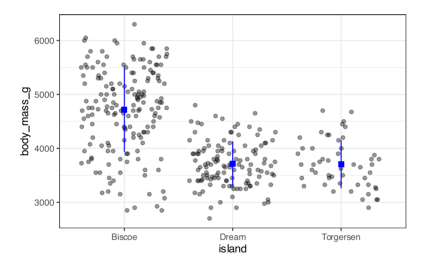
```
p_new +
    scale_x_continuous(n.breaks = 10) +
    scale_y_continuous(breaks = seq(from = 30, to = 65, by = 5.5))
```

```
geom_smooth() using formula = 'y ~ x'
```

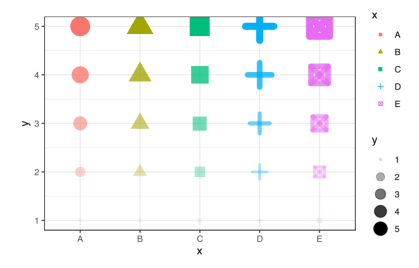


Warning: Removed 2 rows containing non-finite outside the scale range (`stat\_summary()`).

Warning: Removed 2 rows containing missing values or values outside the scale range (`geom\_point()`).



#### Geoms



## **Stats**

## stat\_summary()

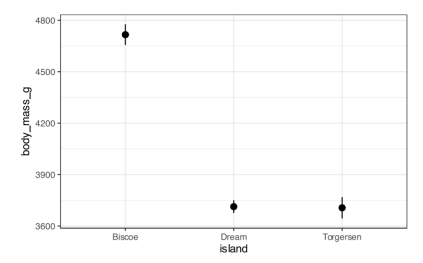
stat\_summary() calculates some summary statistics as y, ymin, and ymax (and possibly other aesthetic mappings) and supplies them to a geom (default = "pointrange"). This is a shortcut for doing the sort of plot we did in the "Data" section without having to create a separate dataset.

First, let's see what the default looks like:

```
ggplot(penguins, aes(x = island, y = body_mass_g)) +
   stat_summary()
```

Warning: Removed 2 rows containing non-finite outside the scale range (`stat\_summary()`).

No summary function supplied, defaulting to `mean\_se()`



As you can see in the warning, by default it is plotting mean  $\pm$  SE (standard error) with the mean\_se() function. To instead plot mean  $\pm$  SD we can either create our own function or use mean\_sdl() and change it's mult argument from the default 2 which doubles the SD.

Let's see what mean sdl() does on its own:

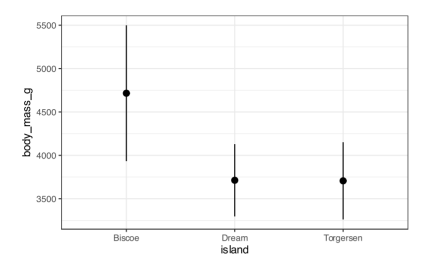
```
mean_sdl(rnorm(100), mult = 1)
```

```
y ymin ymax
1 -0.08363667 -1.061473 0.8941994
```

It creates a tibble with the columns y, ymin, and ymax. Any function that does this will work with stat\_summary() by supplying it to the fun.data argument. To pass along the mult argument, we have to use the fun.args argument.

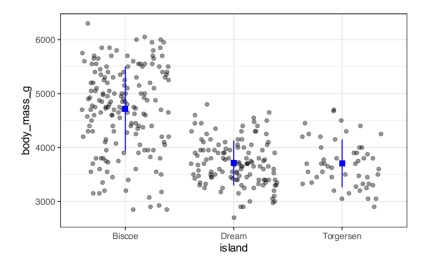
```
ggplot(penguins, aes(x = island, y = body_mass_g)) +
   stat_summary(fun.data = mean_sdl, fun.args = list(mult = 1))
```

Warning: Removed 2 rows containing non-finite outside the scale range
(`stat\_summary()`).



Now we can add our data!

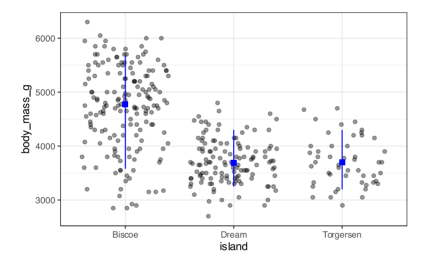
Warning: Removed 2 rows containing non-finite outside the scale range
(`stat\_summary()`).



We could instead use our own custom function that plots the median and the middle 80% of data points, for example.

```
Warning: Removed 2 rows containing non-finite outside the scale range (`stat_summary()`).
```

```
Warning: Removed 2 rows containing missing values or values outside the scale range (`geom_point()`).
```



#### Binned density plot with geom\_histogram() and after\_stat()

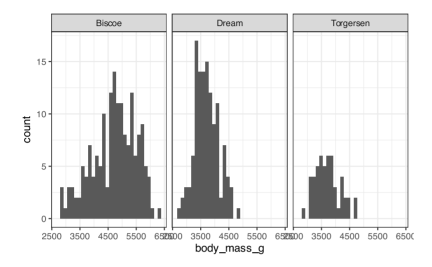
Some "stats" calculate multiple values available with after\_stat(). For example, geom\_histogram() uses the count variable calculated by stat\_bin() to plot the number of data points in each bit on the y-axis.

Here's a histogram of body mass faceted by island:

```
ggplot(penguins) +
  geom_histogram(aes(x = body_mass_g)) +
  facet_wrap(vars(island))
```

```
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```

Warning: Removed 2 rows containing non-finite outside the scale range (`stat bin()`).



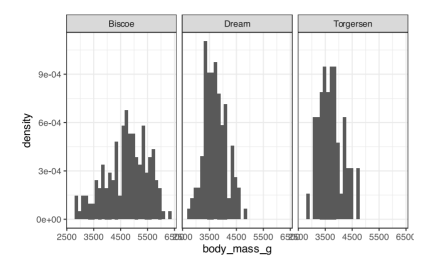
Torgersen island clearly just has fewer penguins, making it somewhat difficult to compare the relative *distribution* of body mass among the islands.

In the "Computed variables" section of ?stat\_bin() you see that it also calculates density, ncount, ndensity, and width. We can plot density of points rather than count using after\_stat(density).

```
ggplot(penguins) +
  geom_histogram(aes(x = body_mass_g, y = after_stat(density))) +
  facet_wrap(vars(island))
```

```
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```

Warning: Removed 2 rows containing non-finite outside the scale range  $(\dot stat_bin()\dot )$ .

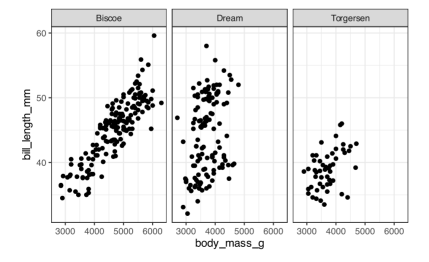


Now the bars for each island add up to 1, taking sample size out of the equation.

#### **Facets**

facet\_wrap() facets by a single variable

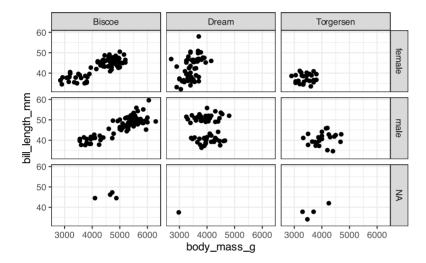
```
ggplot(penguins, aes(x = body_mass_g, y = bill_length_mm)) +
  facet_wrap(vars(island)) +
  geom_point()
```



facet\_grid() facets by two variables

```
ggplot(penguins, aes(x = body_mass_g, y = bill_length_mm)) +
facet_grid(vars(sex), vars(island)) +
geom_point()
```

Warning: Removed 2 rows containing missing values or values outside the scale range (`geom\_point()`).

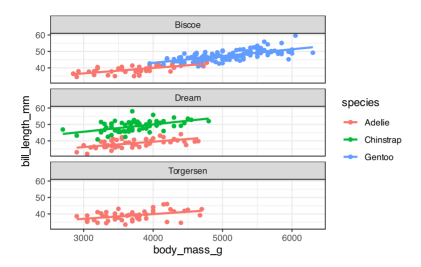


### Coords

There are multiple ways to change the axis limits in ggplot2. First, you can change them with scale\_x\_continuous() or scale\_y\_continuous()

```
p
```

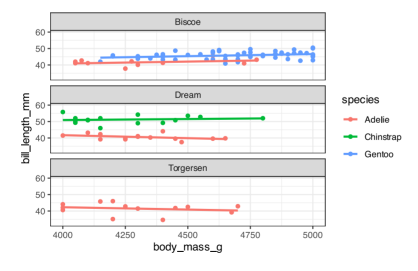
```
geom_smooth() using formula = 'y ~ x'
```



```
p + scale_x_continuous(limits = c(4000, 5000))
```

```
geom_smooth() using formula = 'y ~ x'
```

Warning: Removed 222 rows containing non-finite outside the scale range ( $`stat\_smooth()`)$ .

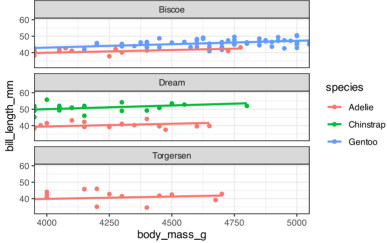


You can see in the warning message printed that 222 rows have been **removed** before drawing this plot. You can tell because the trend lines produced by <code>geom\_smooth()</code> now have different slopes because they are fit to only a subset of data!

You can also change the limits with coord\_cartesian()

```
p + coord_cartesian(xlim = c(4000, 5000))

`geom_smooth()` using formula = 'y ~ x'
Biscoe
```



This has the effect of *zooming in* on the x-axis. The lines and points just outside of the limits are cut off and the slopes of the trend lines are unaffected because no data has been removed.