# Notes for foundations of ggplot2

Eric Scott

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```
library(tidyverse)
```

```
— Attaching core tidyverse packages
—

/ dplyr 1.1.4 / readr 2.1.5

/ forcats 1.0.0 / stringr 1.5.1

/ ggplot2 3.5.1 / tibble 3.2.1

/ lubridate 1.9.3 / tidyr 1.3.1

/ purrr 1.0.2

— Conflicts
— tidyverse_conflicts()

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

/ dplyr::lag() masks stats::lag()

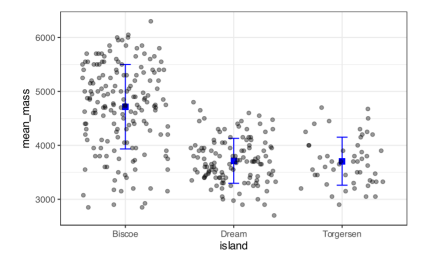
i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all conflicts to become errors
```

```
library(palmerpenguins)
theme_set(theme_bw())
```

#### Data

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

```
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.

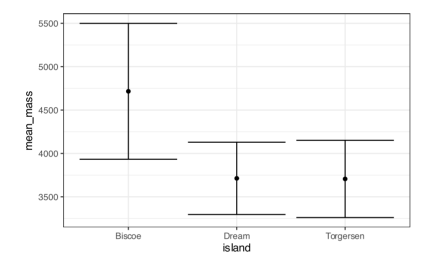
```
peng_summary <-
penguins |>
group_by(island) |> #for each island...
summarize(
   mean_mass = mean(body_mass_g, na.rm = TRUE),
   sd = sd(body_mass_g, na.rm = TRUE)
)
peng_summary
```

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.

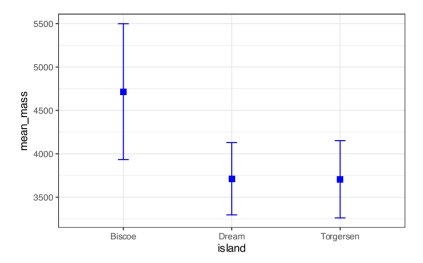
```
ggplot(peng_summary, aes(x = island)) +
    #mean:
    geom_point(aes(y = mean_mass)) +
    #sd:
    geom_errorbar(aes(ymin = lower, ymax = upper))
```



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

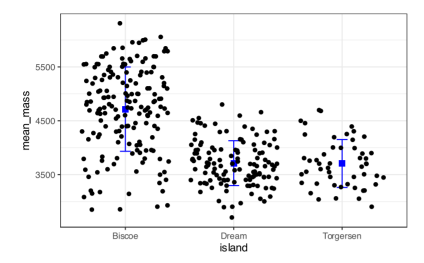
```
ggplot(peng_summary, aes(x = island)) +
    #mean:
    geom_point(
        aes(y = mean_mass),
        shape = "square",
        color = "blue",
        size = 2.5
)+
    #sd:
    geom_errorbar(
        data = peng_summary,
        aes(ymin = lower, ymax = upper),
        width = 0.1,
```

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



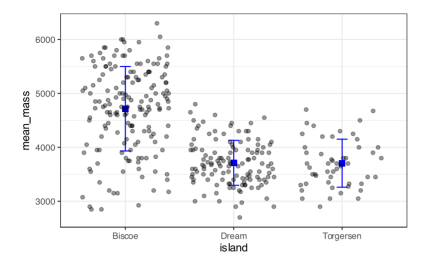
Now we can add the raw data using <code>geom\_jitter()</code> by overriding the data argument.

```
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
 )
```



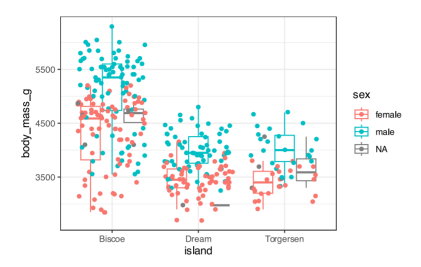
### **Aesthetics**

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()
```

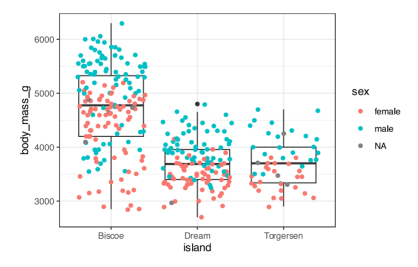
```
Warning: Removed 2 rows containing non-finite outside the scale range
(`stat_boxplot()`).
```



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()`).

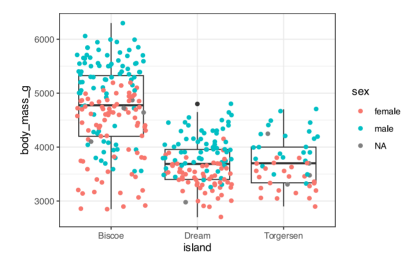


Or, we can use inherit.aes = FALSE and specify *all* the aesthetic mappings for the boxplot layer.

```
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()`).

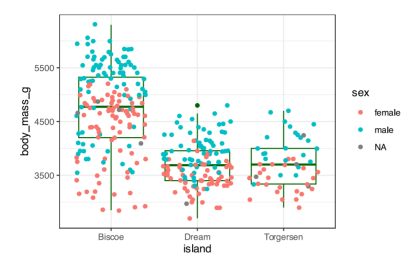
Warning: Removed 2 rows containing missing values or values outside the scale range (`geom point()`).



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()`).

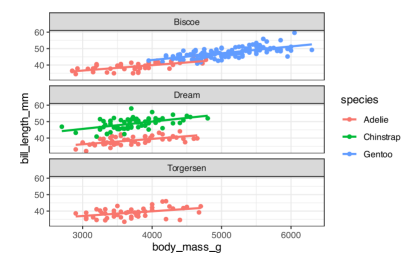


# **Scales**

Here's the original plot, saved as p

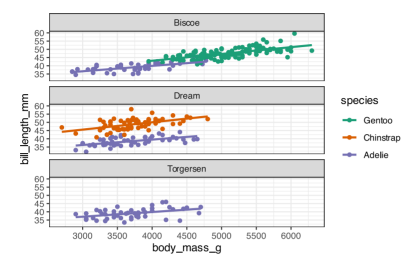
```
p <-
    ggplot(penguins |> drop_na(), aes(x = body_mass_g, y = bill_length_mm, color
= species)) +
    geom_point() +
    geom_smooth(method = "lm", se = FALSE) +
    facet_wrap(vars(island), ncol = 1)
p
```

```
`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 ~ 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.

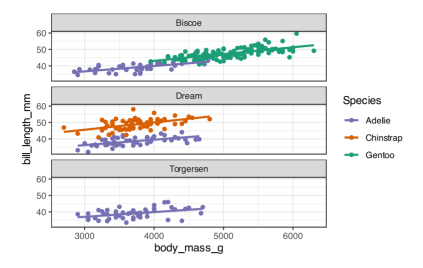
```
my_cols <- c(
   "Adelie" = "#7570b3",
   "Chinstrap" = "#d95f02",
   "Gentoo" = "#1b9e77"
)</pre>
```

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
)
```

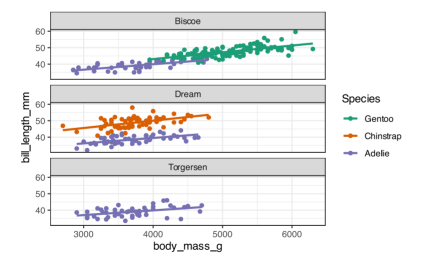
```
`geom_smooth()` using formula = 'y ~ x'
```



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>
```

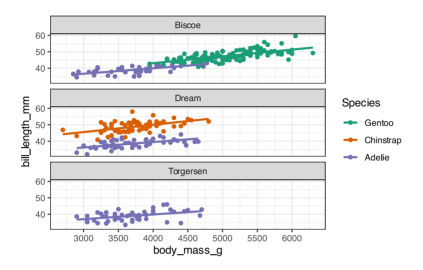




Now we can move on to the x and y axes. For the x-axis, let's increase the number of breaks to *about* 10.

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

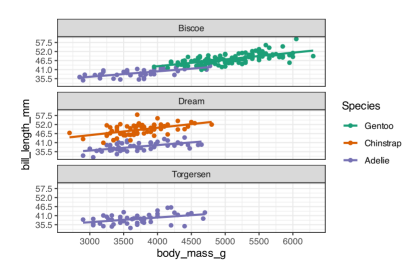
```
geom_smooth() using formula = 'y ~ x'
```



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

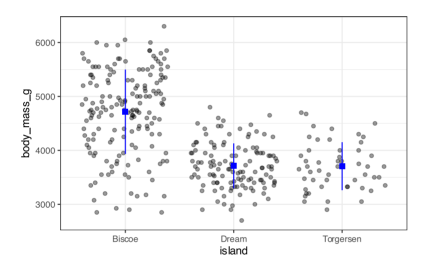
```
p_new +
    scale_x_continuous(n.breaks = 10) +
    scale_y_continuous(breaks = seq(from = 30, to = 65, by = 5.5))
```

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



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

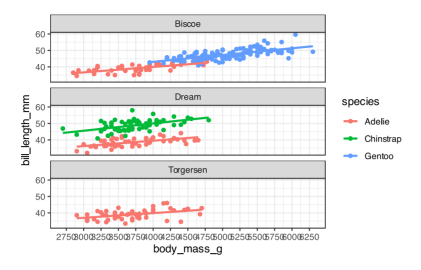
Warning: Removed 2 rows containing missing values or values outside the scale range (`geom\_point()`).



You can also use scale\_x\_log10() to create breaks for a log scale

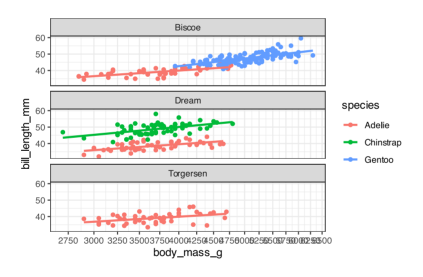
```
p + scale_x_continuous(n.breaks = 15)
```

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

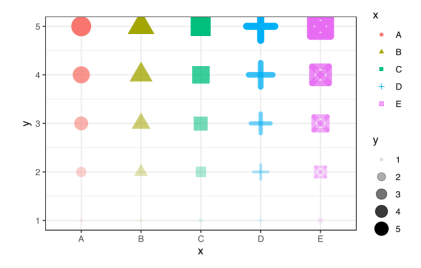


```
p + scale_x_log10(n.breaks = 15)
```

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



# Geoms



## **Stats**

## stat\_summary()

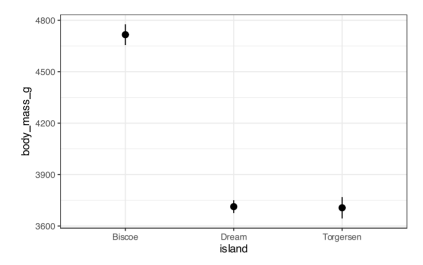
stat\_summary() calculates some summary statistics as y, ymin, and ymax (and possibly other aesthetic mappings) and supplies them to a geom (default = "pointrange"). This is a shortcut for doing the sort of plot we did in the "Data" section without having to create a separate dataset.

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:

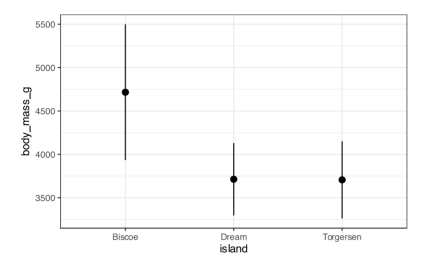
```
mean_sdl(rnorm(100), mult = 1)
```

```
y ymin ymax
1 -0.1080539 -1.157794 0.9416861
```

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.

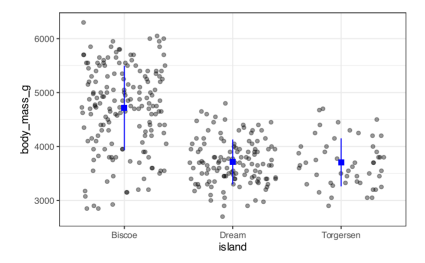
```
ggplot(penguins, aes(x = island, y = body_mass_g)) +
   stat_summary(fun.data = mean_sdl, fun.args = list(mult = 1))
```

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()`).



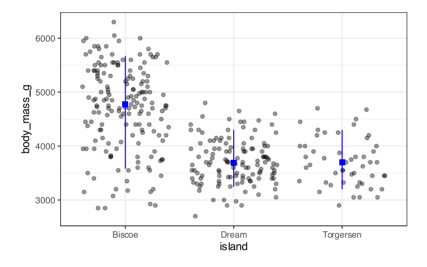
We could instead use our own custom function that plots the median and the middle 80% of data points, for example.

```
median_80 <- function(vals) {
   tibble(y = median(vals, na.rm = TRUE),
        ymin = quantile(vals, 0.1),
        ymax = quantile(vals, 0.9))
}

ggplot(penguins, aes(x = island, y = body_mass_g)) +
   geom_jitter(alpha = 0.4, height = 0) +
   stat_summary(fun.data = "median_80", color = "blue", shape = "square")</pre>
```

```
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()

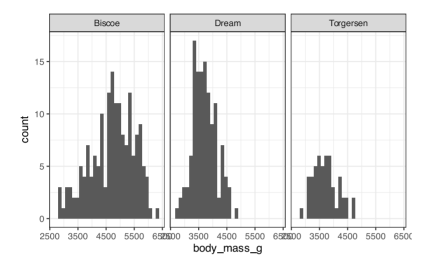
Some "stats" calculate multiple values available with after\_stat(). For example, geom\_histogram() uses the count variable calculated by stat\_bin() to plot the number of data points in each bit on the y-axis.

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()`).



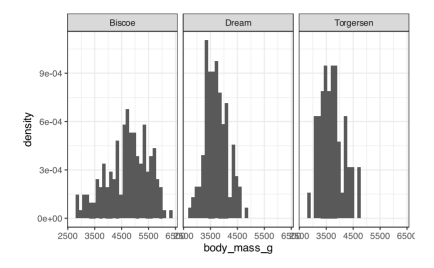
Torgersen island clearly just has fewer penguins, making it somewhat difficult to compare the relative *distribution* of body mass among the islands.

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  $(\dot stat_bin()\dot )$ .

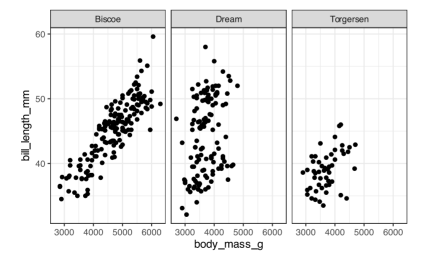


Now the bars for each island add up to 1, taking sample size out of the equation.

#### **Facets**

facet\_wrap() facets by a single variable

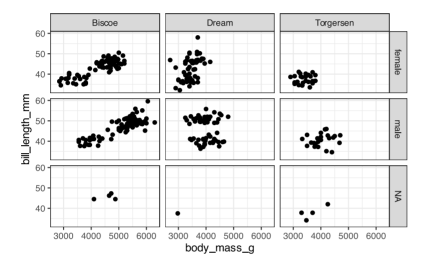
```
ggplot(penguins, aes(x = body_mass_g, y = bill_length_mm)) +
  facet_wrap(vars(island)) +
  geom_point()
```



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()`).

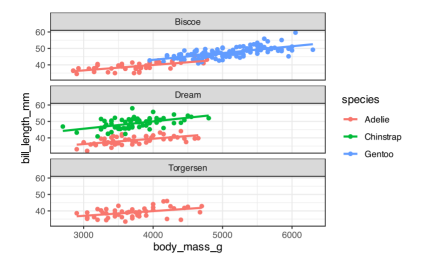


### Coords

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
```

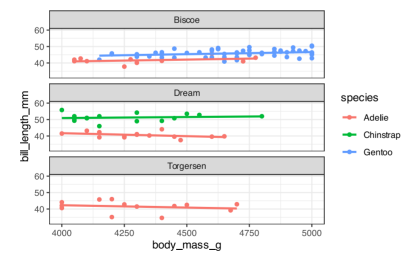
```
`geom_smooth()` using formula = 'y ~ x'
```



```
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 ( $`stat\_smooth()`)$ .

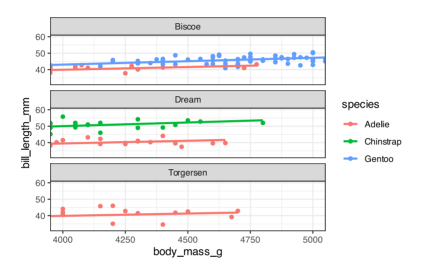


You can see in the warning message printed that 222 rows have been **removed** before drawing this plot. You can tell because the trend lines produced by <code>geom\_smooth()</code> now have different slopes because they are fit to only a subset of data!

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

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

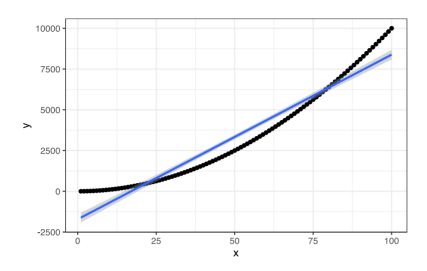
```
geom_smooth() using formula = 'y ~ x'
```



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.

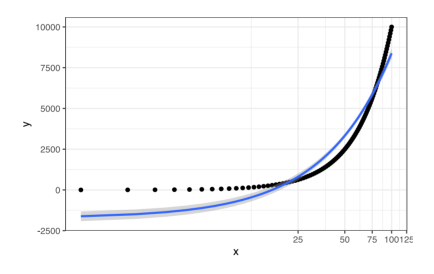
We can use coord\_trans() to warp the coordinate system to plot things on a log scale, for example. This does something slightly different than scale\_x\_log10().

```
`geom_smooth()` using formula = 'y \sim x'
```



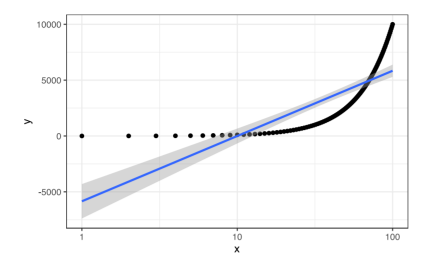
p2 + coord\_trans(x = "log10")

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



p2 + scale\_x\_log10()

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



You can see that  $coord\_trans()$  warps the coordinate space because the regression line, which we specified as being straight with method = "lm", is now curved. But with  $scale\_x\_log10()$  the line stays straight—the scale transforms the data before  $geom\_smooth()$  calculates its statistics.