Drosophilidae wings — Bolstad et al. 2015

Boldstad et al. (2015) studied the allometric relationships between the length of the wings of Drosophilidae and the length of the L2 vein that runs across the wing. They measured more than 20,000 individuals, belonging to 111 species. In their Figure 1, they show regressions between the log length of the wing size and the log length of the L2 vein. They produce a regression for each species and sex. They then added points showing the average values for each species. The file data/Bolstad2015_figure1.pdf contains a simplified version of Figure 1 of the original paper. Reproduce the figure. The data is stored in CSB/ggplot2/data/Boldstad2015_data.csv. The logarithm of wing size and L2 length is already taken.

Always start by importing libraries and the data:

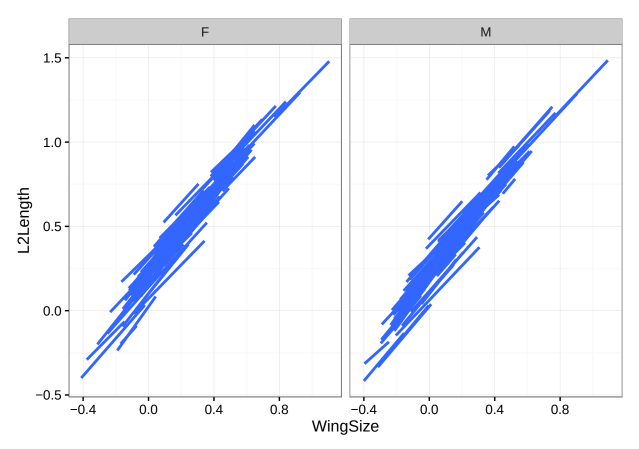
[1] 111

1

```
library(ggplot2)
library(dplyr)
##
## Attaching package: 'dplyr'
##
## The following objects are masked from 'package:stats':
##
##
       filter, lag
##
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
# read the data
b2015 <- read.csv("../data/Bolstad2015_data.csv")
# explore dimensions of the data
dim(b2015)
## [1] 20345
# explore structure of the data
head(b2015)
##
       Species
                        ID
                                Date Sex WingSize L2Length
## 1 D_acutila ACU1006.TIF 24_Jul_01  F 0.1311220 0.4972620
## 2 D_acutila ACU1009.TIF 24_Jul_01
                                       F 0.1360881 0.4879716
## 3 D_acutila ACU1010.TIF 24_Jul_01
                                      F 0.1953933 0.5401366
## 4 D_acutila ACU1013.TIF 24_Jul_01
                                       F 0.2773285 0.6463595
## 5 D_acutila ACU1018.TIF 24_Jul_01
                                       F 0.1515312 0.4977579
## 6 D_acutila ACU1021.TIF 24_Jul_01
                                       F 0.1751865 0.4919342
# number of species
dim(distinct(select(b2015, Species)))
```

We want to produce a regression per Species, and facet by Sex. We could use the aes colour, but that would produce many different colors. Alternatively, we can define the group directly:

```
# draw a regression line for each species and sex
pl <- ggplot(data = b2015, aes(x = WingSize, y = L2Length, group = Species)) +
   geom_smooth(alpha = 0.1, method = "lm", se = FALSE) + facet_grid(.~Sex) + theme_bw()
pl</pre>
```

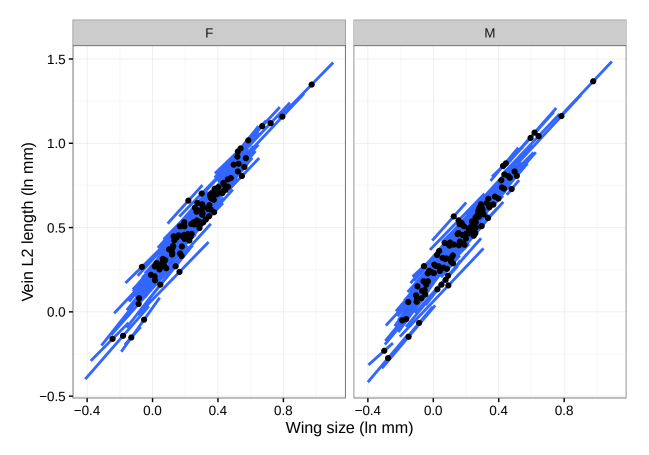


Now we want to add another layer, with a point for each Species/Sex combination. We create a new data frame using the function summarise of dplyr:

```
## Source: local data frame [220 x 4]
## Groups: Species [?]
##
##
          Species
                     Sex
                           WingSize L2Length
           (fctr) (fctr)
                                         (db1)
##
                               (dbl)
## 1
     C procnemis
                       M 0.07531273 0.1904114
## 2
        D_acutila
                       F 0.19390128 0.5319778
## 3
        D_acutila
                       M 0.07009950 0.4083030
```

```
## 4
       D_{affinis}
                     F 0.19271682 0.4567225
## 5
       D_affinis
                       M 0.07317315 0.3128763
                       F 0.30348696 0.6383540
## 6
       D_albostr
## 7
       D_albostr
                      M 0.18746097 0.5073995
## 8
       D_algonqu
                       F 0.34727439 0.6329803
## 9
       D_algonqu
                       M 0.21745274 0.4923027
## 10
                       F 0.47858859 0.7935839
       D_america
## ..
```

```
# add points to current plot
pl <- pl + geom_point(data = by_species_sex)
# add labels for x- and y-axis
pl <- pl + xlab("Wing size (ln mm)") + ylab("Vein L2 length (ln mm)")
# plot the result
pl</pre>
```



```
# save the graph
ggsave(pl, filename = "../data/Bolstad2015_figure1.pdf", width = 9, height = 4)
```