Working example for generating multiple plots inside a map2 call

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1 Introduction

Consider the problem of running a data analysis requiring a separate analysis for each of n strata. For example consider an effort to model the relationship between Bill length and Flipper length across three different species of penguins.

We can work with the dataset ${\tt penguins}$ included in the package ${\tt palmerpenguins}$



Figure 1: purrr

library(palmerpenguins)

One naive approach is to split the dataset and do three separate analyses:

```
df1 = split(penguins, penguins$species)
```

The R package purr provides a straightforward method to conduct the analyses with a single command. Assume the set of data tables are contained in a list of dataframes. Also assume the analysis is a simple visualization of a potential linear association between two features,

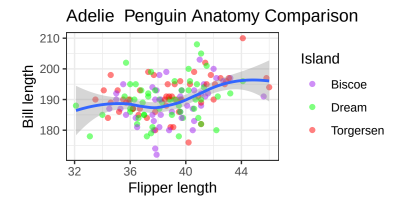


Figure 2: Scatterplot for species: Adelie

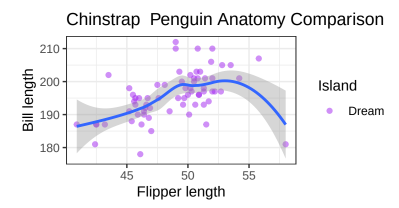


Figure 3: Scatterplot for species: Chinstrap

Gentoo Penguin Anatomy Comparison | Second Penguin Anatomy Comparison | Island | Biscoe | Penguin Anatomy Comparison | Island | Penguin Anatomy Comparison | Island | Penguin Anatomy Comparison | Island | Penguin Anatomy Comparison | Penguin Anatomy Comparison | Island | Penguin Anatomy Comparison | Penguin A

Figure 4: Scatterplot for species: Gentoo

- 2 Plots for every variable and each species map inside map see ref 2 below
- 3 combine plots in a upper triangular grid with correlation coefs
- 4 Code

```
library(pacman)
p_load(palmerpenguins, tidyverse, knitr)

opts_chunk$set(
   warning = FALSE, message = FALSE, echo = FALSE, fig.width = 4,
   fig.height = 2.1, results = "asis", dev = "pdf")

df1 = split(penguins, penguins$species)

foo <- function (df, z) {
   # df is the data frame, z is the name of the species</pre>
```

```
df |> ggplot(aes(x = bill_length_mm, y = flipper_length_mm))+
     geom_point(aes(color = island), alpha = .5) +
13
     geom_smooth() +
14
     scale_color_manual(values = c("purple", "green", "red")) +
15
     theme_bw() +
16
     labs(title = paste(z, " Penguin Anatomy Comparison"), x = "Flipper length",
17
          y = "Bill length", color = "Island")
   plotfile_name = paste0(z,".pdf")
19
   ggsave(plotfile_name)
20
   cat(paste0("\\includegraphics[]{",plotfile_name,"}"), "\n")
   cat(paste0("\\captionof{figure}{Scatterplot for species: ",z,"}"), "\n")
22
   cat("\\vspace{1cm}", "\n")
23
   }
24
25
   bar = df1 |> map2(names(df1), foo)
```

5 References

principal components analysis Automating exploratory plots with ggplot2 and purrr