

Functional Plot Generation with purrr

Using map2 and pmap to programmatically build scatter plot grids across penguin species

Ronald ‘Ryy’ G. Thomas

2026-04-30

Table of contents

1	Introduction	2
1.1	Motivations	2
1.2	Objectives	3
2	Prerequisites and Setup	5
3	What is Functional Plot Generation?	5
4	Getting Started: Preparing the Data	6
5	Building the Plotting Function	7
6	Mapping Across Species and Variable Pairs	8
7	Assembling the Grid with Patchwork	9
7.1	Extracting Individual Plots	9
7.2	Defining the Layout	9
7.3	Composing the Final Figure	10
7.4	Things to Watch Out For	13
8	What Did We Learn?	14
8.1	Lessons Learnt	14
8.2	Limitations	14
8.3	Opportunities for Improvement	15
9	Wrapping Up	15
10	See Also	15
11	Reproducibility	16



Figure 1: Functional programming tools bring order to repetitive plotting tasks.

Generating publication-quality plot grids without writing the same code three times.

1 Introduction

I did not really know how to programmatically generate multiple plots from grouped data until I started working with the Palmer Penguins dataset. The task seemed straightforward: produce scatter plots for every pairwise combination of numeric variables, separately for each penguin species, and then assemble them into a single coherent grid.

My first instinct was to copy-paste code for each species and each variable pair. That approach collapsed quickly once I counted the combinations: three species, four numeric variables, six pairwise combinations per species – eighteen plots total. Copy-pasting eighteen ggplot calls is tedious, brittle, and nearly impossible to maintain.

The `purrr` package solved this problem cleanly. By splitting the data by species and mapping a plotting function over every variable combination, I could generate all eighteen plots with a single pipeline and assemble them into a structured grid using `patchwork`. This post documents that workflow.

1.1 Motivations

- I was tired of manually duplicating ggplot code for each subgroup in a dataset.
- I wanted to see all pairwise relationships between numeric features at a glance, stratified by species.
- I needed a workflow that could scale: if a fourth species appeared in the data, the code should handle it without modification.

- I was curious whether `purrr::pmap` could handle the three-argument case (x variable, y variable, grouping variable) cleanly.
- I wanted practice assembling complex multi-panel layouts with `patchwork` beyond simple `+` and `/` operators.

1.2 Objectives

1. Split a data frame by a categorical variable (species) and iterate over the resulting list with `map2`.
2. Use `pmap` to map a plotting function across all pairwise combinations of numeric columns.
3. Assemble the resulting list of plots into a structured grid using `patchwork::wrap_plots` with a custom design layout string.
4. Produce a final composite figure with species labels, shared legends, and collected axis titles.

I am documenting my learning process here. If you spot errors or have better approaches, please let me know in the contact section below.



Figure 2: Functional programming and visualisation.

2 Prerequisites and Setup

This post assumes familiarity with `ggplot2` and basic `tidyverse` operations. The key packages are:

- `purrr` for functional iteration (`map2`, `pmap`)
- `patchwork` for plot composition (`wrap_plots`, `plot_layout`, `plot_spacer`)
- `palmerpenguins` for the dataset
- `rlang` for tidy evaluation (`.data[[var]]` pronoun)

```
library(palmerpenguins)
library(tidyverse)
library(purrr)
library(patchwork)
library(rlang)
library(grid)
```

The Palmer Penguins dataset contains measurements for 344 penguins across three species (Adelie, Chinstrap, Gentoo) from three islands in the Palmer Archipelago, Antarctica.

```
penguins |>
  drop_na() |>
  count(species) |>
  knitr::kable(
    caption = "Observations per species after
              removing missing values."
  )
```

Table 1: Observations per species after removing missing values.

species	n
Adelie	146
Chinstrap	68
Gentoo	119

3 What is Functional Plot Generation?

Functional plot generation means treating plot creation as a function and then applying that function systematically across combinations of inputs. Instead of writing:

```
# Do not do this for 18 combinations
plot_adelie_bill_flipper <- ggplot(...)
plot_adelie_bill_body <- ggplot(...)
plot_adelie_bill_depth <- ggplot(...)
# ... 15 more times
```

you write one plotting function and let `purrr` call it for every combination of species, x-variable, and y-variable. The output is a list of `ggplot` objects that you can arrange into any layout.

Think of it like a mail merge: you define the template once and let the data fill in the blanks.

4 Getting Started: Preparing the Data

The first step is to sample from the penguins data and split it into a named list by species. We also compute all pairwise combinations of the four numeric columns (`bill_length_mm`, `bill_depth_mm`, `flipper_length_mm`, `body_mass_g`).

```
set.seed(42)
df_sampled <- penguins |>
  sample_n(50) |>
  drop_na()

df_by_species <- split(df_sampled, df_sampled$species)

numeric_cols <- names(df_sampled)[3:6]
pairs <- t(combn(numeric_cols, 2))
colnames(pairs) <- c("x_var", "y_var")
pair_grid <- as.data.frame(pairs) |>
  mutate(group_var = "sex")
```

The `pair_grid` data frame now holds six rows, one for each pairwise combination. Each row names an x-variable, a y-variable, and a grouping variable (`sex`) for colour coding within each species panel.

```
pair_grid |>
  knitr::kable(
    caption = "All pairwise variable combinations
to plot."
  )
```

Table 2: All pairwise variable combinations to plot.

x_var	y_var	group_var
bill_length_mm	bill_depth_mm	sex
bill_length_mm	flipper_length_mm	sex
bill_length_mm	body_mass_g	sex
bill_depth_mm	flipper_length_mm	sex
bill_depth_mm	body_mass_g	sex
flipper_length_mm	body_mass_g	sex

5 Building the Plotting Function

The core of the workflow is a single function that accepts variable names as strings and a data frame, then returns a ggplot object. The `.data[[var]]` pronoun from `rlang` allows column selection by string name inside `aes()`.

```
make_scatter <- function(x_var, y_var,
                        group_var, species_name,
                        df) {
  df |>
  ggplot(aes(
    x = .data[[x_var]],
    y = .data[[y_var]]
  )) +
  geom_point(
    aes(color = .data[[group_var]]),
    alpha = 0.5
  ) +
  geom_smooth(
    method = "loess",
    se = TRUE,
    linewidth = 0.7
  ) +
  scale_color_manual(
    values = c("purple", "green", "red")
  ) +
  theme_bw() +
  theme(text = element_text(size = 8))
}
```

Two design choices deserve explanation. First, `geom_smooth(method = "loess")` adds a local regression line so we can see nonlinear trends. Second, the function does not set titles – those will come from the grid layout labels. Keeping the function minimal makes it reusable across different layouts.



Figure 3: Iterating with purpose across data strata.

6 Mapping Across Species and Variable Pairs

This is the heart of the approach. We use `map2` to iterate over the list of species data frames and their names simultaneously. Inside that outer loop, `pmap` iterates over every row of `pair_grid`, calling `make_scatter` with the appropriate variable names.

```
all_plots <- df_by_species |>
  map2(names(df_by_species), function(df, spc) {
    pair_grid |>
      pmap(function(x_var, y_var, group_var) {
        make_scatter(
          x_var, y_var, group_var, spc, df
        )
      })
  })
```

The result `all_plots` is a nested list: the outer level has three elements (one per species), and each inner level has six elements (one per variable pair). That gives us eighteen ggplot objects total, organized by species.

```

cat(
  "Species:", length(all_plots), "\n",
  "Plots per species:", length(all_plots[[1]]), "\n",
  "Total plots:", sum(lengths(all_plots)), "\n"
)

```

```

Species: 3
Plots per species: 6
Total plots: 18

```

7 Assembling the Grid with Patchwork

The final step is arranging all eighteen plots into a structured grid. I want each species to occupy one row group, with a text label identifying the species. The `patchwork` package supports custom layout strings where each letter maps to a named plot element.

7.1 Extracting Individual Plots

First, we pull every plot out of the nested list and assign readable names.

```

p1 <- all_plots[[1]][[1]]
p2 <- all_plots[[1]][[2]]
p3 <- all_plots[[1]][[3]]
p4 <- all_plots[[1]][[4]]
p5 <- all_plots[[1]][[5]]
p6 <- all_plots[[1]][[6]]
p7 <- all_plots[[2]][[1]]
p8 <- all_plots[[2]][[2]]
p9 <- all_plots[[2]][[3]]
p10 <- all_plots[[2]][[4]]
p11 <- all_plots[[2]][[5]]
p12 <- all_plots[[2]][[6]]
p13 <- all_plots[[3]][[1]]
p14 <- all_plots[[3]][[2]]
p15 <- all_plots[[3]][[3]]
p16 <- all_plots[[3]][[4]]
p17 <- all_plots[[3]][[5]]
p18 <- all_plots[[3]][[6]]

```

7.2 Defining the Layout

The layout string below arranges plots in an upper triangular pattern for each species, with text labels (X, Y, Z) marking each species group. Each letter in the string maps to one named plot element in `wrap_plots`.

```

layout_string <- "
X##
ABC
#DE
##F
Y##
GHI
#JK
##L
Z##
MNO
#PQ
##R
"

```

The # characters represent empty cells. This produces a staircase pattern: three plots in the first row, two in the second, one in the third – mirroring the upper triangle of a pairwise comparison matrix.

7.3 Composing the Final Figure

We create text labels for each species using `grid::textGrob` and pass everything to `wrap_plots`.

```

label_sp1 <- grid::textGrob(
  names(df_by_species)[1],
  gp = gpar(fontsize = 10, fontface = "bold")
)
label_sp2 <- grid::textGrob(
  names(df_by_species)[2],
  gp = gpar(fontsize = 10, fontface = "bold")
)
label_sp3 <- grid::textGrob(
  names(df_by_species)[3],
  gp = gpar(fontsize = 10, fontface = "bold")
)

composite <- wrap_plots(
  X = label_sp1,
  A = p1, B = p2, C = p3,
  D = p4, E = p5, F = p6,
  Y = label_sp2,
  G = p7, H = p8, I = p9,
  J = p10, K = p11, L = p12,
  Z = label_sp3,
  M = p13, N = p14, O = p15,
  P = p16, Q = p17, R = p18,

```

```
design = layout_string
) +
plot_layout(
  guides = "collect",
  axis_titles = "collect"
) +
theme(
  legend.position = "bottom",
  legend.direction = "horizontal",
  text = element_text(size = 8)
)

composite
```

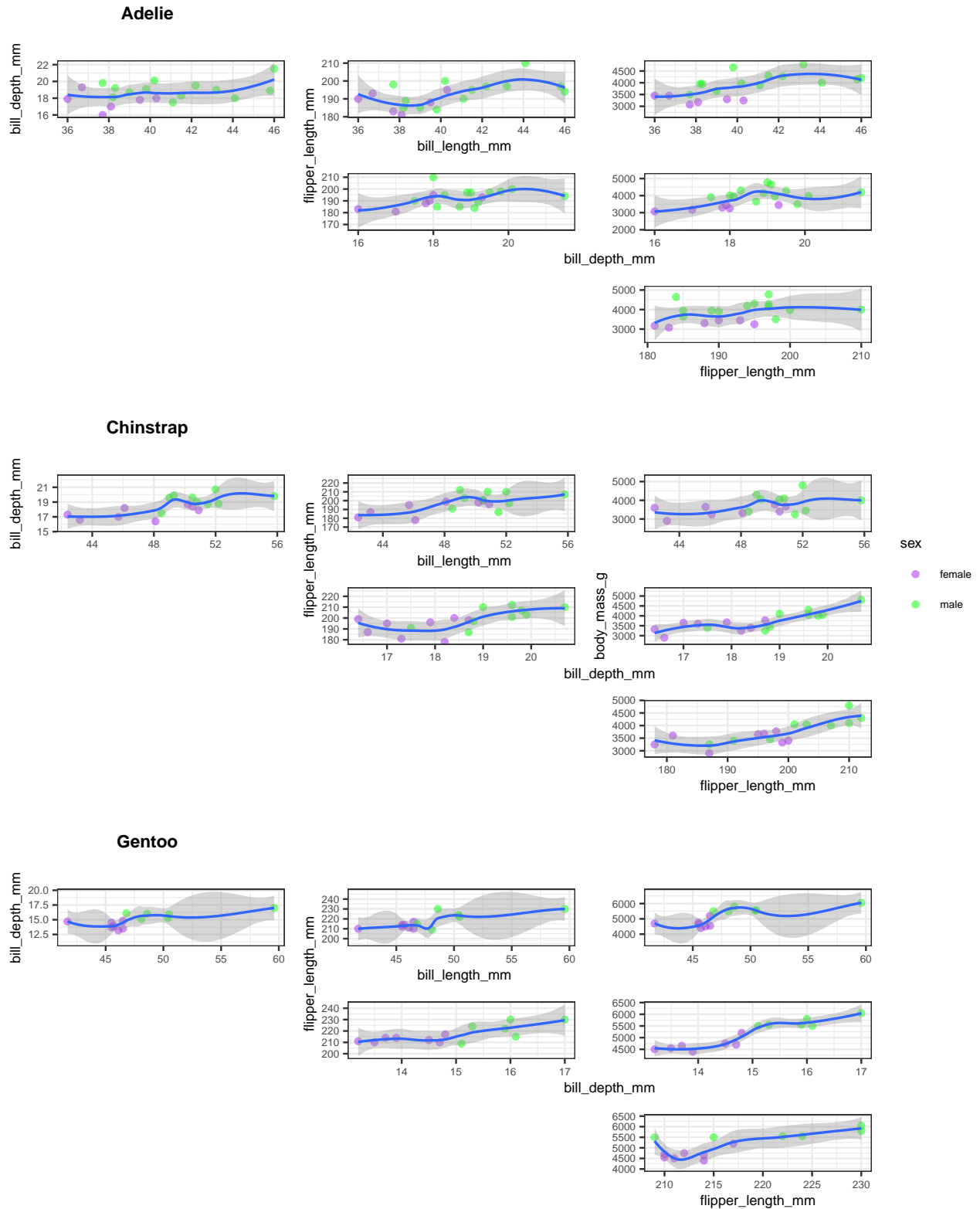


Figure 4: Pairwise scatter plots for four numeric penguin measurements, stratified by species. Each row group shows one species with a LOESS smooth and points coloured by sex.

The `guides = "collect"` argument consolidates duplicate legends into a single shared legend at the bottom. The `axis_titles = "collect"` argument prevents repeated axis labels across panels. Together, these two settings produce a clean, readable composite figure.

7.4 Things to Watch Out For

1. **Variable types matter.** The `.data[[var]]` pronoun works only when the column name is a string. If you pass a symbol instead, you will get cryptic subscription errors.
2. **Missing data propagates.** If you forget `drop_na()` before splitting, some species subsets will contain NA rows, and `geom_smooth` will warn about removed observations.
3. **Layout string alignment.** Every row in the `patchwork` layout string must have the same number of characters. A misaligned row silently produces an incorrect layout.
4. **Global environment side effects.** The original draft used `assign(..., envir = .GlobalEnv)` inside the plotting function. This is fragile and pollutes the workspace. Returning the plot object and storing it in a list is cleaner and more predictable.
5. **Sample size per species.** When sampling 50 rows from the full dataset, some species may end up with very few observations. For production analyses, use the full dataset or stratified sampling to ensure adequate representation.



Figure 5: Assembling the pieces into a coherent whole.

8 What Did We Learn?

8.1 Lessons Learnt

Conceptual Understanding:

- Pairwise scatter plot matrices provide a rapid visual summary of relationships between continuous features, but they scale quadratically: four variables produce six pairs, five produce ten.
- Stratification by species reveals within-group patterns that pooled analyses obscure (Simpson's paradox).
- LOESS smoothers are useful for initial exploration but carry no inferential interpretation without further modeling.
- The upper-triangular layout avoids redundancy by showing each pair exactly once.

Technical Skills:

- `purrr::map2` iterates over two lists in parallel – ideal for pairing data frames with their names.
- `purrr::pmap` generalises to arbitrary numbers of arguments by iterating over rows of a data frame.
- `rlang::.data[[var]]` enables column selection by string name inside `aes()`, which is essential for programmatic ggplot construction.
- `patchwork::wrap_plots` with a custom `design` string provides fine-grained control over multi-panel layouts.

Gotchas and Pitfalls:

- Forgetting `drop_na()` before `split()` produces species-level data frames with missing rows that cause warnings in `geom_smooth`.
- The `plot_spacer()` function is useful for empty cells, but the `#` character in the design string is more readable for fixed layouts.
- `assign()` to `.GlobalEnv` inside mapped functions creates hard-to-debug side effects. Always return values from functions and collect them in lists.
- `scale_color_manual` requires the correct number of colour values matching the levels of the grouping variable.

8.2 Limitations

- The sample of 50 penguins is small. Some species may have fewer than 10 observations, making LOESS fits unreliable.
- The grouping variable (`sex`) is hard-coded. A more general approach would accept the grouping variable as a parameter.
- The layout string is manually crafted for exactly three species and six variable pairs. Adding a fourth species or fifth numeric variable requires rewriting the layout.
- No statistical tests accompany the visual exploration. The plots show associations but do not quantify significance or effect size.
- The colour palette (purple, green, red) is not colourblind-friendly. Production code should use a palette validated for accessibility.

8.3 Opportunities for Improvement

1. Replace the hard-coded layout string with a function that generates the design string dynamically based on the number of species and variable pairs.
2. Add correlation coefficients as text annotations inside each scatter plot panel.
3. Use `GGally::ggpairs` as a comparison point – it handles pairwise plots natively, though with less layout flexibility.
4. Implement stratified sampling with `dplyr::slice_sample(n, by = species)` to ensure balanced representation.
5. Replace the manual plot extraction (p1 through p18) with a programmatic approach using `list_flatten` and named assignment.
6. Switch to a colourblind-friendly palette such as `viridis` or the Okabe-Ito palette.

9 Wrapping Up

This post demonstrated how `purrr::map2` and `purrr::pmap` can replace copy-pasted ggplot code with a single functional pipeline. The key insight is that plots are objects: you can store them in lists, pass them to functions, and assemble them programmatically.

The approach worked well for the Palmer Penguins data. Starting from a single plotting function and a table of variable combinations, we generated eighteen scatter plots and arranged them into a structured grid that reveals within-species patterns. The total code is compact enough to fit in a single script, yet flexible enough to adapt to new datasets.

If you are working with grouped data and finding yourself copying ggplot code, try this pattern:

- **Split** the data by your grouping variable.
- **Define** a single plotting function that accepts column names as strings.
- **Map** the function across all combinations using `pmap`.
- **Assemble** the resulting list with `patchwork`.

10 See Also

- [Automating exploratory plots with ggplot2 and purrr](#) – Ariel Muldoon’s tutorial on the same pattern with detailed examples.
- [Principal components and penguins](#) – PCA as an alternative to pairwise scatter plots for high-dimensional penguin data.
- [purrr documentation](#) – Official reference for `map`, `map2`, and `pmap`.
- [patchwork documentation](#) – Guide to plot composition and layout design.
- [Programming with dplyr](#) – Background on `.data[[var]]` and tidy evaluation.

11 Reproducibility

This analysis uses the `palmerpenguins` built-in dataset and requires no external data files. To reproduce:

```
quarto render index.qmd
```

Session information:

```
sessionInfo()
```

```
R version 4.5.3 (2026-03-11)
```

```
Platform: aarch64-apple-darwin20
```

```
Running under: macOS Tahoe 26.4.1
```

```
Matrix products: default
```

```
BLAS: /Library/Frameworks/R.framework/Versions/4.5-arm64/Resources/lib/libRblas.0.dylib
```

```
LAPACK: /Library/Frameworks/R.framework/Versions/4.5-arm64/Resources/lib/libRlapack.dylib; LAPACK version
```

```
locale:
```

```
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
```

```
time zone: America/Los_Angeles
```

```
tzcode source: internal
```

```
attached base packages:
```

```
[1] grid      stats      graphics  grDevices  utils      datasets  methods
```

```
[8] base
```

```
other attached packages:
```

```
[1] rlang_1.1.7      patchwork_1.3.2    lubridate_1.9.4
```

```
[4] forcats_1.0.0    stringr_1.6.0      dplyr_1.2.1
```

```
[7] purrr_1.2.1      readr_2.2.0        tidyr_1.3.2
```

```
[10] tibble_3.3.1     ggplot2_4.0.2      tidyverse_2.0.0
```

```
[13] palmerpenguins_0.1.1
```

```
loaded via a namespace (and not attached):
```

```
[1] Matrix_1.7-4      gtable_0.3.6      jsonlite_2.0.0    compiler_4.5.3
```

```
[5] tinytex_0.59      tidyselect_1.2.1  parallel_4.5.3    splines_4.5.3
```

```
[9] scales_1.4.0      yaml_2.3.12       fastmap_1.2.0     lattice_0.22-9
```

```
[13] R6_2.6.1          labeling_0.4.3    generics_0.1.4    knitr_1.51
```

```
[17] pillar_1.11.1     RColorBrewer_1.1-3 tzdb_0.5.0         stringi_1.8.7
```

```
[21] xfun_0.57         S7_0.2.1          otel_0.2.0        timechange_0.3.0
```

```
[25] cli_3.6.5         mgcv_1.9-4        withr_3.0.2       magrittr_2.0.5
```

```
[29] digest_0.6.39     hms_1.1.4         nlme_3.1-168      lifecycle_1.0.5
```

```
[33] vctrs_0.7.2       evaluate_1.0.5    glue_1.8.0        farver_2.1.2
```

```
[37] rmarkdown_2.31    tools_4.5.3       pkgconfig_2.0.3   htmltools_0.5.9
```

12 Let's Connect

- **GitHub:** [rgt47](#)
- **Twitter/X:** [@rgt47](#)
- **LinkedIn:** [Ronald Glenn Thomas](#)
- **Email:** [rgtlab.org/contact](mailto:rgt@rgtlab.org)

I would enjoy hearing from you if:

- You spot an error or a better approach to any of the code in this post.
- You have suggestions for topics you would like to see covered.
- You want to discuss R programming, data science, or reproducible research.
- You have questions about anything in this tutorial.
- You just want to say hello and connect.