Data visualisation - part 3: compound figures with cowplot

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1 Background

There are many add on packages for ggplot2. Here, I want to highlight and show examples for one of those packages: cowplot (short for Claus O. Wilkes plot package, initially developed for Claus's lab members). It is extensively documented on this webpage: cowplot.

cowplot is described as

provid[ing] various features that help with creating publicationquality figures, such as a set of themes, functions to align plots and arrange them into complex compound figures, and functions that make it easy to annotate plots and or mix plots with images.

2 Setting up

In the following two chunks, we will again set up our analys document by specifying the options for knitr and

load all libraries required for our analysis:

```
library("tidyverse")
library("cowplot")
library("sf")
library("rnaturalearth")
library("RColorBrewer")
```

3 Individual visualisations

In the following section, we will load the dataset we worked with in the previous session and recreate the visualisations we used to show the antigenic maps, histogram and boxplots of the clusters by time.

Unlike the previous session, where we immediately displayed the output of each ggplot2 call, we will now save each plot into a different object. To do so, we will first create the common ggplot object p, that relies on our dataset coord:

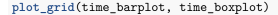
```
coord <- read csv("data/2004 Science Smith data.csv")</pre>
#> Parsed with column specification:
#> cols(
#> name = col_character(),
    year = col_double(),
#>
    cluster = col_character(),
#>
#> type = col_character(),
   x.coordinate = col_double(),
#>
#> y.coordinate = col_double(),
#> location = col_character(),
#> lat = col_double(),
#>
     lng = col\_double()
#> )
p <- ggplot(data=coord)</pre>
```

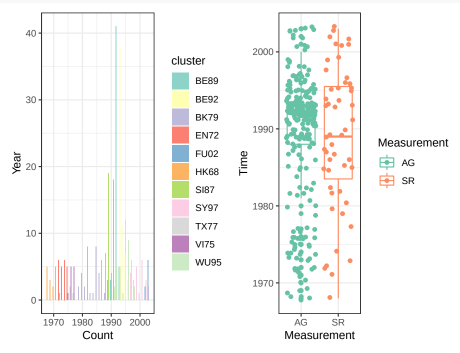
To recreate the previous plots, we save them into a new object by adding the appropriate layers to p:

cowplot's plot_grid function takes these plotting objects as input and arranges them into a grid.

4 Simple compound plots

To generate our first compound figure, we provide the bar and boxplot objects to plot_grid, which arranges them in a single row, next to one another.



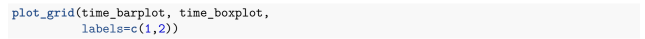


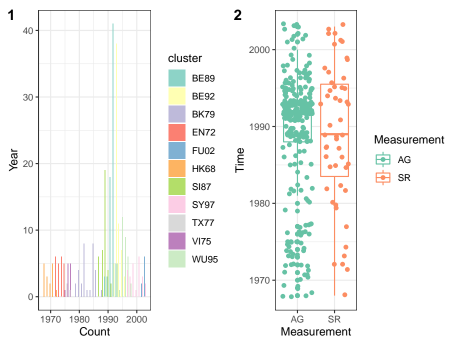
4.1 Customising plot_grid

plot_grid is highly customisable, including layout, labels, alignments and scaling.

4.1.1 Labels

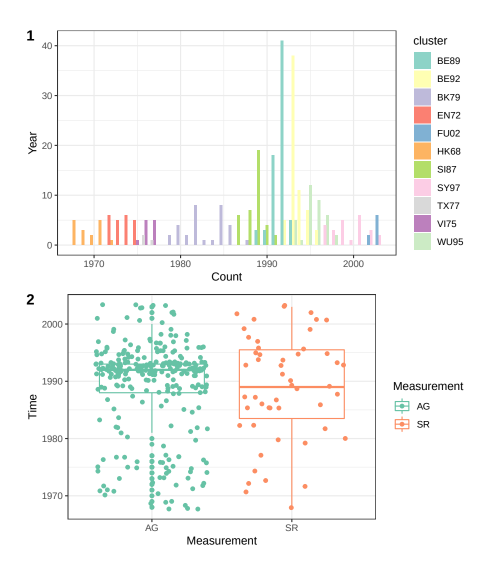
To create publication-ready figures, we can set a label for each plot, by either specifying the desired labels in a vector of the same lengths as the number of plots (as we do here for 2 plots) or by setting labels="AUTO" to auto-generate upper-case labels or labels="auto" to auto-generate lower-case labels.





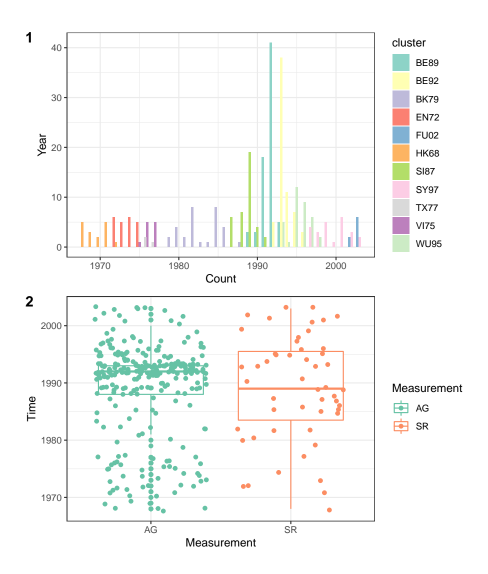
4.1.2 Layout

Per default, plot_grid chooses a 'one row, two column' layout for the plots we specified. To specify a 'two row, one column' layout we can either specify the number of column ncol or number or rows nrow argument:



4.1.3 Alignment

You might notice that in the one column layout, the plots are slightly shifted with respect to their y-axis and legends alignment. Per default, plot_grid keeps the *axis titles* aligned. To align by *actual axis*, we can use a combination of the align and axis arguments. Here we want the plots to be vertically aligned along the left and right margins of the plot panel:



4.2 Exercises

- 1. Autogenerate lower-case labels for the compound plot of time_boxplot and time_barplot.
- 2. Change the default layout by specifying the number of columns.
- 3. Read the help function for plot_grid and experiment with other axis options.

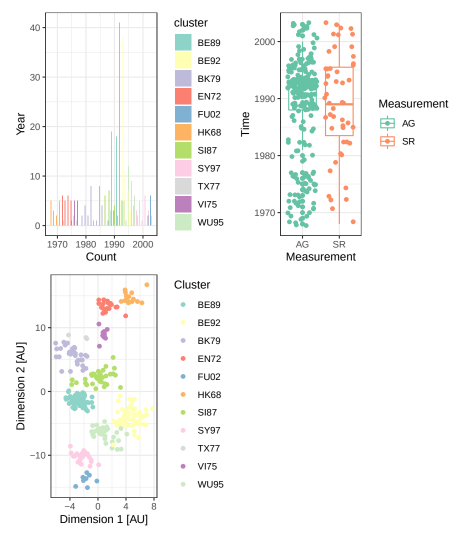
5 Nested compound plots

So far, we have only considered the two time-related plots we generated to visualise the antigenetic cluster data. In the following few chunks, we will see how we can add a third plot to the grid that has different dimensions than then previous two.

5.1 Arranging plots

Let's start with the default behaviour:

plot_grid(time_barplot, time_boxplot, antigenic_map)

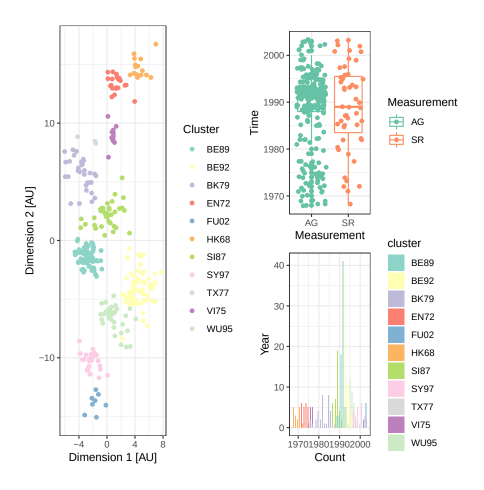


This layout does not look great, for several reason:

- 1. We have a void in the lower right corner;
- 2. The heights of both rows are the same, which unnessecarily stretches the bar and boxplot.

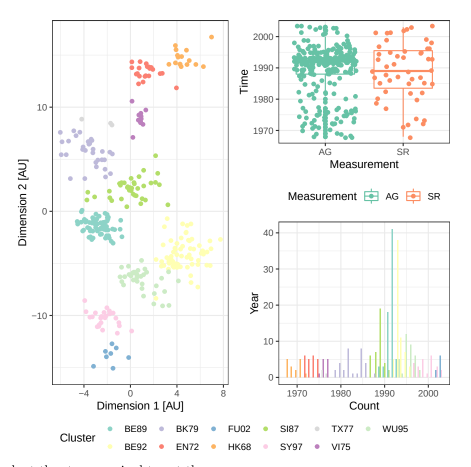
To address both issues, we can arrange a two-column layout, with the first column containing the antigenic map only, spanning over two rows, and the bar and boxplot depicted in the first and second row of the second column, respectively.

To achieve this, we will use two calls to plot_grid, the first to generate our composite for the right-hand column, the second to put this composite together with the antigenic map.



5.2 Shared legends

In the visualisation above, we have color-coded both the antigenic map and the bar plot by cluster. Below, we will have a look how we can add one shared legend as an additional row to plot_grid and remove the individual legends. This is what our final figure will look like:



Let's have a look at the steps required to get there:

First, we will create the individual components of our panel, this time adding legend.position via theme. We create the right_column, similar as above, but, in addition, specify that in this plot_grid, the boxplot legend should move below the plot (theme(legend.position = "bottom")) and the legend from the barplot should be removed with theme(legend.position = "none"). Similarly, we remove the the legend from the antigenic_map that we pass to plot_grid.

Note: We only move and remove the legends from the objects that we pass to plot_grid, the original objects remain unchanged.

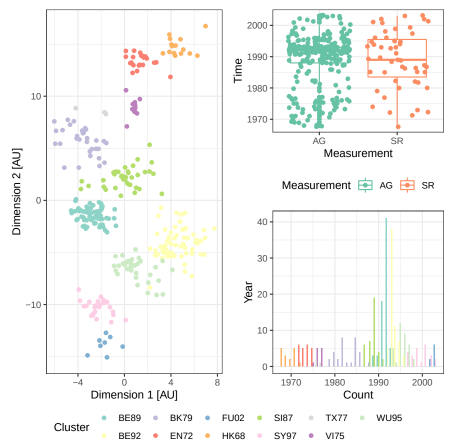
We then create a new map object from which we extract the legend. We will place the shared legend as an additional row at the bottom of the our panel plot, so we want a legend with horizontal layout, as if it was at the bottom of a plot. Thus, we specify in this new object, that the legend should be at the bottom of the plot (as we saw above) and, in addition, specify that the color legend should have two rows for its elements with guides(color=guide_legend(nrow=2)).

We then extract the legend from the new_map object with get_legend:

```
new_map <- antigenic_map +
  guides(color=guide_legend(nrow=2)) +
  theme(legend.position = "bottom")

cluster_legend <- get_legend(new_map)</pre>
```

Finally, we will put it all together in a new call to plot_grid. In addition to the arguments that we have already seen for plot_grid, we also specify rel_heights here, which says that the ratio of the first row (i.e. our compound panel) to the second row (the shared legend) should be 10:1.



5.2.1 Exercises

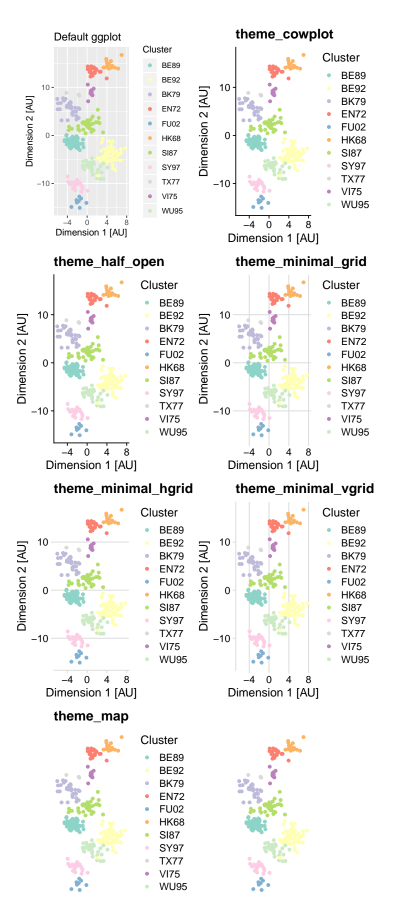
- 1. What would be a good labeling strategy for this nested plot? Label accordingly.
- 2. Play with the ratios provided to rel_heights. Why does it not make sense to provide rel_widths in this example?

6 cowplot beyond plot_grid

Many people come to cowplot in the search of a function for composite figures (plot_grid). cowplot has other things to offer, most notably for me, it extends the range of different themes provided by ggplot.

Below an overview of available cowplot themes, all applied to the antigenic_map scatter plot:

```
plot_grid(antigenic_map + theme_gray() + ggtitle("Default ggplot"),
    antigenic_map + theme_cowplot() + ggtitle("theme_cowplot"),
    antigenic_map + theme_half_open() + ggtitle("theme_half_open"),
    antigenic_map + theme_minimal_grid() + ggtitle("theme_minimal_grid"),
    antigenic_map + theme_minimal_hgrid() + ggtitle("theme_minimal_hgrid"),
    antigenic_map + theme_minimal_vgrid() + ggtitle("theme_minimal_vgrid"),
    antigenic_map + theme_map() + ggtitle("theme_map"),
    antigenic_map + theme_nothing() + ggtitle("theme_nothing"),
    nrow=4,
    align="vh",
    axis="tblr")
```



All themes are cowplot themes apart from the default ggplot2 theme shown for reference. The lower right plot shows theme_nothing by cowplot.

Note: Not all themes are equally well suited for each plot type. The ggplot2 default for boxplots chooses a simply horizontal grid for instance, where as the same theme applied to the scatter plot display horizontal and vertical grids.

6.1 Exercises

- 1. Discuss which other plot type might benefit from a simpler grid than the default one chosen?
- 2. What happens if you add a theme to plot_grid?
- 3. Apply different themes to the three individual plots and visualise in grid.

7 Additional material

The ggpubr package provides some easy-to-use functions for creating and customizing ggplot2- based publication ready plots: ggplot2 Based Publication Ready Plots