ggalign: Bridging the Grammar of Graphics and Complex layout

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2024-11-24

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Preface

Welcome to ggalign documents. Examples in the book are generated under version 0.0.5.9000.

In the world of data visualization, aligning multiple plots in a coherent and organized layout is often a challenging task, especially when dealing with complex datasets that require precise alignment across rows, columns, and even within plot elements. While existing tools provide some solutions, they often fall short in offering the flexibility, control, and simplicity that users need to create intricate and beautiful plots. This is where ggalign comes in.

The ggalign package, built on top of the powerful ggplot2 framework, is designed to solve this very problem. It offers a suite of functions specifically crafted for aligning and organizing plots with minimal effort. Whether you need to align observations based on statistical measures, group plots by categorical factors, or fine-tune the layout to match the precise needs of your data, ggalign gives you the tools you need to create polished, publication-ready visualizations.

This book serves as both an introduction to the ggalign package and a comprehensive guide to mastering its features. Whether you're a beginner or an experienced user of ggplot2, you'll find detailed explanations, step-by-step tutorials, and real-world examples to help you leverage the full potential of ggalign in your work.

Throughout this book, we will cover everything from basic concepts to advanced layout customizations, focusing on key functions like stack_layout(), align_* series (including align_group(), align_order(), and align_hclust()), and how to combine them with other ggplot2 layers to create aligned plots. Additionally, you'll learn how to adapt ggalign for different data types and scenarios, allowing you to develop flexible, dynamic visualizations tailored to your specific needs.

By the end of this book, you will be equipped to use ggalign effectively in your own projects, whether for scientific research, data analysis, or any other field where data visualization is key. Our goal is to provide you with the knowledge and confidence to tackle complex visualization challenges and transform your datasets into clear, impactful, and visually appealing plots.

Thank you for choosing ggalign. We hope this book will inspire you to explore the endless possibilities that come with aligned data visualization.

1 Introduction

ggalign extends ggplot2 by providing advanced tools for aligning and organizing multiple plots, particularly those that automatically reorder observations, such as dendrogram. It offers fine control over layout adjustment and plot annotations, enabling you to create complex visualizations while leveraging the familiar grammar of ggplot2.

1.1 Installation

You can install ggalign from CRAN using:

```
install.packages("ggalign")
```

Alternatively, install the development version from r-universe with:

```
install.packages("ggalign",
    repos = c("https://yunuuuu.r-universe.dev", "https://cloud.r-project.org")
)
```

or from GitHub with:

```
# install.packages("remotes")
remotes::install_github("Yunuuuu/ggalign")
```

1.2 General design

The core feature of ggalign lies in its integration of the grammar of graphics into advanced visualization through its object-oriented Layout system. The package provides two main Layout classes:

- the StackLayout class: Put plots horizontally or vertically.
- the QuadLayout class: Arranges plots in the four quadrants (top, left, bottom, right) around a main plot. This layout is ideal for designs that require supplementary plots or annotations surrounding a central figure.

Both Layout classes support the alignment of observations (ordinal variable). Observations refer to data points or samples, allowing for consistent alignment of corresponding data across multiple plots when using the same axis values. Depending on whether you want to align observations across multiple plots within the layout, the following variants are available:

For StackLayout:

- stack_align(): Align the observations along the stack.
- stack_free(): Does not align the observations.

For QuadLayout:

- quad_free/ggside: Never align observations.
- quad_alignh: Align observations in the horizontal direction.
- quad_alignv: Align observations in the vertical direction.
- quad_alignb: Align observations in both horizontal and vertical directions.



Figure 1.1: General design of ggalign

1.3 Getting Started

```
library(ggalign)
```

Loading required package: ggplot2

The usage of ggalign is simple if you're familiar with ggplot2 syntax, the typical workflow includes:

- 1. Initialize the layout using:
- stack_layout(): Arrange Plots Horizontally or Vertically
 - cross_align: Arrange Plots Crosswise Horizontally or Vertically
- quad_layout(): Arrange Plots in the Quad-Side of a main plot
 - ggheatmap(): Create a Complex Heatmap.
 - ggoncoplot(): Create OncoPrint Visualizations from Genetic Alteration Data
- 2. Customize the layout with:
- align_group(): Group observations into panel with a group variable.
- align_kmeans(): Group observations into panel by kmeans.
- align_order(): Reorder layout observations based on statistical weights or by manually specifying the observation index.
- align_hclust()/align_dendro(): Reorder or group observations based on hierarchical clustering.
- 3. Adding plots with ggalign() or ggfree(), and then layer additional ggplot2 elements such as geoms, stats, or scales.

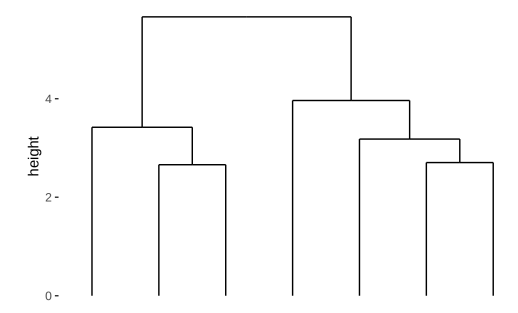
```
set.seed(123)
small_mat <- matrix(rnorm(56), nrow = 7)
rownames(small_mat) <- paste0("row", seq_len(nrow(small_mat)))
colnames(small_mat) <- paste0("column", seq_len(ncol(small_mat)))</pre>
```

Every *_layout() function accepts default data, which will be inherited by all plots within the layout.

Here's a simple example:

```
stack_alignv(small_mat) +
   align_dendro() +
   theme(axis.text.y = element_text())
   3
```

- (1) We initialize a vertical stack.
- (2) Reorder the observations based on hierarchical clustering and add a dendrogram tree.
- (3) Add y-axis text.



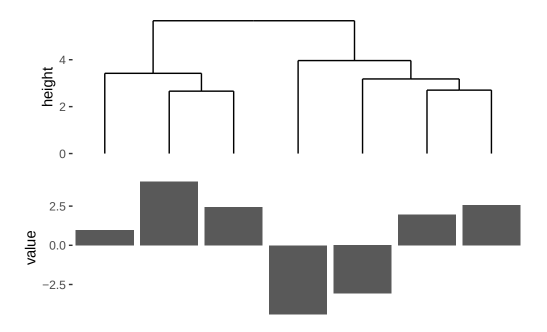
This produces a simple dendrogram. By default, stack_alignv() removes the axis text on the axis used for aligning observations. This is because it's often unclear which plot should display the axis text, as typically, we want it to appear in only one plot. However, you can easily use the theme() function to control where the axis text appears.

Internally, align_dendro() will reorder the observations based on the dendrogram, and other plots in the layout will follow this ordering.

```
stack_alignv(small_mat) +
   align_dendro() +
   ggalign(data = rowSums) +
   geom_bar(aes(.names, value), stat = "identity") +
   theme(axis.text.y = element_text())
(1)
(2)
(3)
(4)
(5)
```

- (1) We initialize a vertical stack.
- (2) Reorder the observations based on hierarchical clustering and add a dendrogram tree.

- (3) Create a new ggplot in the layout, and use data based on the sum of the layout data.
- (4) Add a bar layer.
- (5) Add y-axis text.



The data in the underlying ggplot object of ggalign() function contains at least following columns (more details will be introduced in the later chapter):

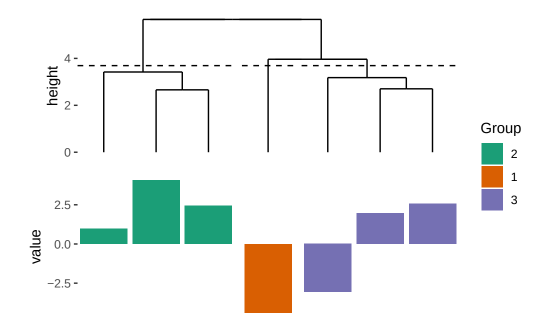
- .panel: the group panel for the aligned axis. It means x-axis for vertical stack layout (including top and bottom annotation), y-axis for horizontal stack layout (including left and right annotation).
- .x or .y: the x or y coordinates
- .names and .index: A factor of the names (only applicable when names exists) and an integer of index of the original data.
- value: the actual value (only applicable if data is a matrix or atomic vector).

It is recommended to use .x/.y, or .names as the x/y mapping.

align_dendro() can also split the observations into groups.

```
stack_alignv(small_mat) +
    align_dendro(k = 3) +
    ggalign(data = rowSums) +
    geom_bar(aes(.names, value, fill = .panel), stat = "identity") +
    scale_fill_brewer(palette = "Dark2", name = "Group") +
    theme(axis.text.y = element_text())
(1)
(2)
(3)
(4)
(4)
(5)
(6)
```

- (1) We initialize a vertical stack.
- (2) Reorder and group the observations based on hierarchical clustering, and add a dendrogram tree.
- (3) Create a new ggplot in the layout, and use data based on the sum of the layout data.
- (4) Add a bar layer.
- (5) Add fill mapping scale.
- (6) Add y-axis text.



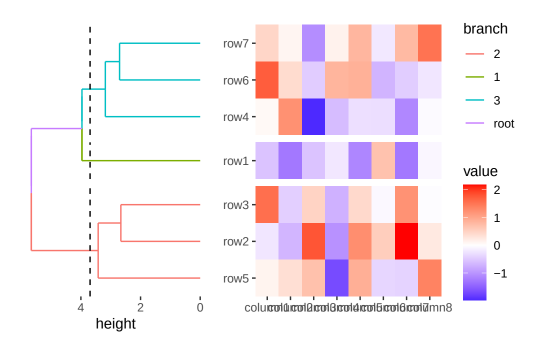
One common visualization associated with the dendrogram is the heatmap. You can use ggheatmap() to initialize a heatmap layout. When grouping the observations using align_dendro(k = 3), a special column named branch is added, which you can use to color the dendrogram tree.

```
ggheatmap(small_mat) +
   anno_left() +
   align_dendro(aes(color = branch), k = 3) +
   scale_fill_brewer(palette = "Dark2")

(1)
(2)
(3)
(4)
```

- (1) We initialize a heatmap layout (ggheatmap(small_mat)).
- (2) we initialize an annotation in the left side of the heatmap body, and set it as the active context, in this way, all following addition will be directed to the left annotation. (anno_left())
- 3 Reorder and group the observations based on hierarchical clustering, and add a dendrogram tree, coloring the tree by branch (align_dendro(k = 3)).

- (4) Add fill mapping scale (scale_fill_brewer(palette = "Dark2")).
- > heatmap built with `geom_tile()`



ggheatmap() will automatically add axis text in the heatmap body, so you don't need to manually adjust axis text visibility using theme(axis.text.x = element_text())/theme(axis.text.y
= element_text()).

Now that we've covered the basic workflow of ggalign, let's move on to exploring StackLayout, a core layout for arranging plots either horizontally or vertically. In the next chapter, we'll dive deeper into how to effectively use StackLayout for arranging and aligning your plots.

2 stack layout

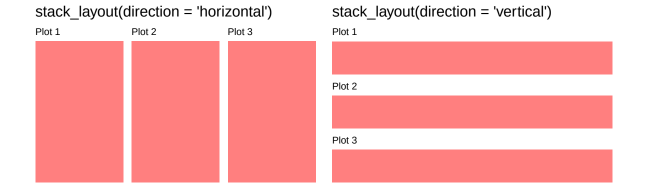
stack_layout() arranges plots either horizontally or vertically, and we can also use the alias
ggstack(). Based on whether we want to align the observations, there are two types of stack
layouts:

- stack_align(): align the observations along the stack.
- stack_free(): don't align the observations.

Several aliases are available for convenience:

- stack_alignv: Aligns the stack vertically (special case of stack_align()).
- stack_alignh: Aligns the stack horizontally (special case of stack_align()).
- stack_freev: A vertical version of stack_free().
- stack_freeh: A horizontal version of stack_free().

```
library(ggalign)
## Loading required package: ggplot2
set.seed(123)
small_mat <- matrix(rnorm(56), nrow = 7)
rownames(small_mat) <- paste0("row", seq_len(nrow(small_mat)))
colnames(small_mat) <- paste0("column", seq_len(ncol(small_mat)))</pre>
```



2.1 Input data

When aligning observations, we typically use a matrix, as it is easy to melt the matrix into a long-formatted data frame. Additionally, matrices are used to fit the observation concept, as they can be transposed (rows to columns, columns to rows), which is necessary for use in functions like quad_layout() and ggheatmap(), where observations may be aligned in both directions simultaneously.

- For stack_free(), a data frame is required, and the input will be automatically converted using fortify_data_frame() if needed.
- For stack_align(), a matrix is required, and the input will be automatically converted using fortify_matrix() if needed.

By default, fortify_data_frame() will invoke the ggplot2::fortify() function for conversion. Note, for matrix, it will be converted to a long-formatted data frame.

stack_align()/stack_free() will set up the layout, but no plot will be drawn until you add
a plot element:

```
stack_alignh(small_mat) +
    layout_annotation(theme = theme(plot.background = element_rect(color = "red")))
# the same for `stack_free()`
```

In this example, we use layout_annotation() to insert a plot background in the entire layout.

2.2 Layout Customization

When we use **stack_align()**, it aligns the observations across multiple plots along the specified direction:

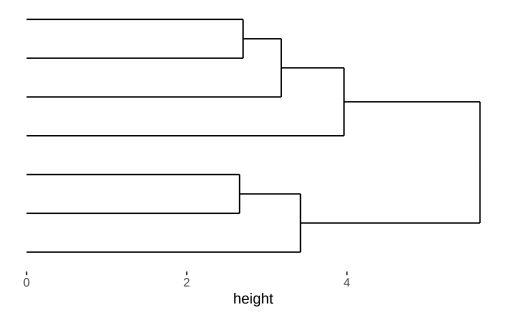
- For stack_alignh(): Alignment occurs along the horizontal direction (y-axis).
- For stack_alignv(): Alignment occurs along the vertical direction (x-axis).

The package offers a suite of align_* functions designed to give you precise control over the layout. These functions enable you to reorder the observations or partition the observations into multiple groups. Instead of detailing each align_* function individually, we will focus on the general usage and how to combine them with stack_align().

Here, we remain take align_dendro() as a example, it can reorder the observations, split them into groups, and can add a plot for visualization.

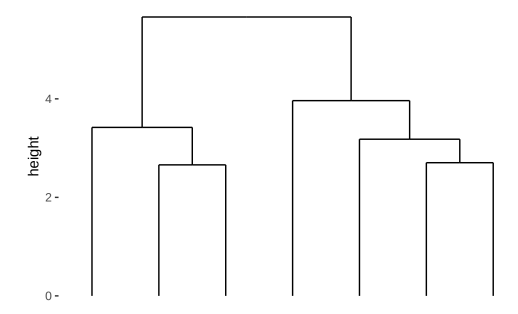
When used for stack_alignh(), the observations are aligned along the y-axis:

```
stack_alignh(small_mat) +
  align_dendro()
```



When used for stack_alignv(), the observations are aligned along the x-axis:

```
stack_alignv(small_mat) +
   align_dendro()
```

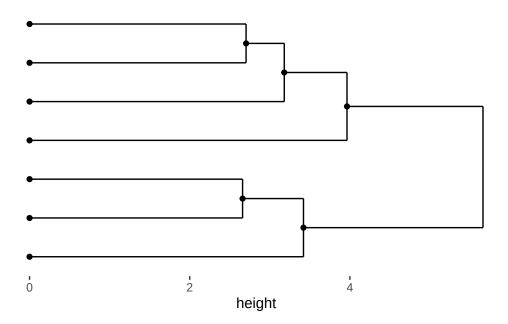


When align_dendro() is added to the layout, it performs following actions:

- 1. reorder the observations.
- 2. set the active plot to the dendrogram.

The active plot refers to the plot that subsequent ggplot2 components will target. In this case, the active plot is the dendrogram, and any new layers added will be applied to it. For instance, we can add additional layers to visualize the dendrogram's structure or data. The default data underlying the ggplot object of align_dendro() consists of the dendrogram node data. It is also possible to use the dendrogram's edge data for customization, which I will introduce in a later chapter.

```
stack_alignh(small_mat) +
  align_dendro() +
  geom_point()
```



The active argument controls whether a plot is set as the active plot. It accepts an active() object with the use argument to specify if the plot should be active.

```
stack_alignh(small_mat) +
   align_dendro(active = active(use = FALSE)) +
   geom_point()
```

```
Error in `stack_layout_add()`:
! Cannot add `geom_point()` to `stack_align()`
i No active plot component
i Did you forget to initialize a <ggplot> object with `ggalign()` or `ggfree()`?
```

Usually, you don't need to set this manually, as the active context is automatically applied only for functions that add plot areas. You can inspect whether a align_* function will add a plot by print it:

```
align_dendro()
```

```
`align_dendro()` object:
  plot: yes
  reorder: yes
  split: no
```

You might find it confusing that we mentioned align_dendro() will split observations into groups, while the print output shows split = "no". This happens because we haven't specified the k/h argument in align_dendro().

```
align_dendro(k = 3L)

`align_dendro()` object:
  plot: yes
  reorder: yes
  split: yes
```

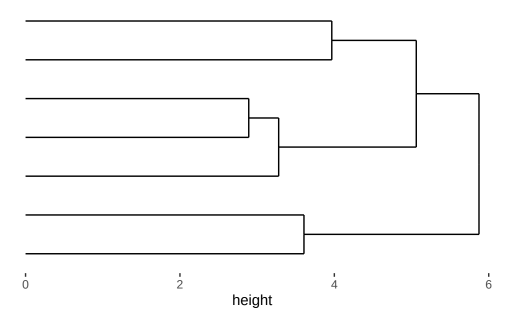
You don't need to explicitly provide data to align_dendro(). By default, it inherits data from the layout. However, you can always provide another data source, but note that this package uses the concept of "number of observations" (NROW()). When aligning the observations, you must ensure the number of observations is consistent across all plots.

```
set.seed(123)
stack_alignh(small_mat) +
    align_dendro(data = matrix(rnorm(56), nrow = 8)) +
    theme(axis.text.y = element_text())

Error in `align()`:
! `align_dendro(data = matrix(rnorm(56), nrow = 8))` (nobs: 8) is not
    compatible with the `stack_align()` (nobs: 7)

set.seed(123)
stack_alignh(small_mat) +
    align_dendro(data = matrix(rnorm(70), nrow = 7)) +
```

theme(axis.text.y = element_text())



Alternatively, you can provide a function (or purrr-lambda) that will be applied to the layout's matrix. Note that, for layouts that align observations, a matrix is always required, so the data input must be in matrix form.

```
set.seed(123)
stack_alignh(small_mat) +
   align_dendro(data = ~ .x[sample(nrow(.x)), ]) +
   theme(axis.text.y = element_text())
```



Without adding another plot, it's difficult to appreciate the benefits. Let's now explore how to incorporate a plot.

2.3 Plot initialize

There are two primary functions for adding plots:

- align_gg()/ggalign(): Create a ggplot object and align with the layout.
- free_gg()/ggfree(): Create a ggplot object without aligning.

Both functions initialize a ggplot object and, by default, set the active plot when added to the layout.

For stack_align(), plots can be added regardless of whether they need to align observations.

```
stack_alignh(small_mat) +
   align_dendro() +
   ggalign(data = rowSums) +
   geom_bar(aes(value, .names), stat = "identity") +
   theme(axis.text.y = element_text())
```



You can build the plot layer separately and then add it to the layout:

```
my_bar <- ggalign(data = rowSums) +
    geom_bar(aes(value, .names), stat = "identity") +
    theme(axis.text.y = element_text())
stack_alignh(small_mat) +
    align_dendro() +
    my_bar</pre>
```



The active argument can also control the place of the plot area to be added. It accepts an active() object with the order argument to specify the order of the plot area.

```
stack_alignh(small_mat) +
    align_dendro() +
    ggalign(data = rowSums, active = active(order = 1)) +
    geom_bar(aes(value, .names), stat = "identity") +
    theme(axis.text.y = element_text())
```



You can also stack plots vertically using stack_alignv():

```
stack_alignv(small_mat) +
  align_dendro() +
  ggalign(data = rowSums) +
  geom_bar(aes(value, .names), stat = "identity") +
  theme(axis.text.y = element_text())
```



stack_align() can also add plot without aligning observations. free_gg() focuses on layout
integration without enforcing strict axis alignment. ggfree() is an alias for free_gg.

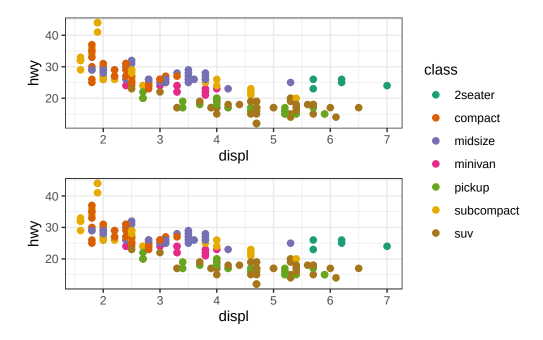
```
stack_alignv() +
    ggfree(mpg, aes(displ, hwy, colour = class)) +
    geom_point(size = 2) +
    ggfree(mpg, aes(displ, hwy, colour = class)) +
    geom_point(size = 2) &
    scale_color_brewer(palette = "Dark2") &
    theme_bw()
```



The & operator applies the added element to all plots in the layout, similar to its functionality in the patchwork package.

For stack_free(), only free plots (ggfree()) can be added. This layout arranges plots in one row or column without enforcing axis alignment:

```
stack_freev(mpg) +
    ggfree(mapping = aes(displ, hwy, colour = class)) +
    geom_point(size = 2) +
    ggfree(mapping = aes(displ, hwy, colour = class)) +
    geom_point(size = 2) &
    scale_color_brewer(palette = "Dark2") &
    theme_bw()
```

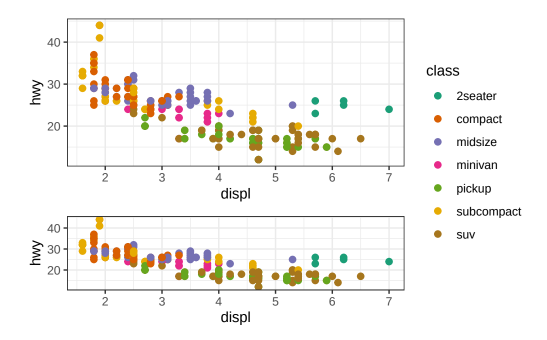


By default, ggfree() will also inherit data from the layout and call fortify_data_frame() to convert the data to a data frame. So, note that if the layout data is a matrix, it will be converted into a long-formatted data frame.

2.4 Plot Size

Both ggalign() and ggfree() functions have a size argument to control the relative width (for horizontal stack layout) or height (for vertical stack layout) of the plot's panel area.

```
stack_freev(mpg) +
    ggfree(mapping = aes(displ, hwy, colour = class), size = 2) +
    geom_point(size = 2) +
    ggfree(mapping = aes(displ, hwy, colour = class), size = 1) +
    geom_point(size = 2) &
    scale_color_brewer(palette = "Dark2") &
    theme_bw()
```



Alternatively, you can define an absolute size by using a unit() object:

```
stack_freev(mpg) +
    ggfree(mapping = aes(displ, hwy, colour = class), size = unit(1, "cm")) +
    geom_point(size = 2) +
    ggfree(mapping = aes(displ, hwy, colour = class)) +
    geom_point(size = 2) &
    scale_color_brewer(palette = "Dark2") &
    theme_bw()
```



2.5 active plot

As mentioned earlier, the active plot refers to the plot that subsequent ggplot2 components will target. The package provide two functions to work with active plot.

- stack_switch(): switch the active context
- stack_active: An alias for stack_switch(), which sets what = NULL

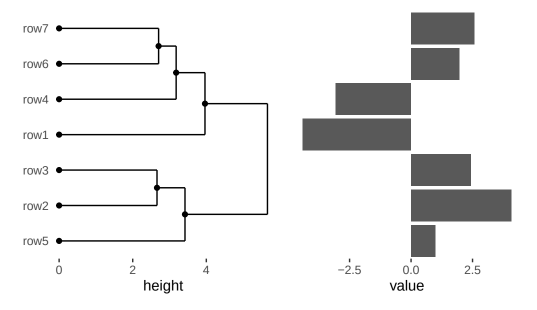
The stack_switch() function accepts the what argument, which can either be the index of the plot added (based on its adding order) or the plot name specified via the active() object using the name argument.

Note that the what argument must be explicitly named, as it is placed second in the function signature. This is because, in most cases, we don't need to switch the active plot manually—adjusting the order of plot additions typically suffices.

```
stack_alignh(small_mat) +
   align_dendro() +
   ggalign(data = rowSums) +
   geom_bar(aes(value, .names), stat = "identity") +
   # switch to the `align_dendro()` plot area
   stack_switch(what = 1) +
```

```
geom_point() +
theme(axis.text.y = element_text()) +
layout_title(title = "switch by integer")
```

switch by integer



```
stack_alignh(small_mat) +
    align_dendro(active = active(name = "tree")) +
    ggalign(data = rowSums) +
    geom_bar(aes(value, .names), stat = "identity") +
    # switch to the `align_dendro()` plot area
    stack_switch(what = "tree") +
    geom_point() +
    theme(axis.text.y = element_text()) +
    layout_title(title = "switch by string")
```

switch by string



In the example, we use layout_title() to insert a title for the entire layout. Alternatively, you can add a title to a single plot with ggtitle().

By setting what = NULL (or alias stack_active()), we remove the active plot. This is particularly useful when the active plot is a nested Layout object, as any additions would otherwise be directed to that nested Layout. By removing the active plot, you can continue adding components directly to the StackLayout.

Now, let's move on to the next chapter where I'll introduce a new Layout.

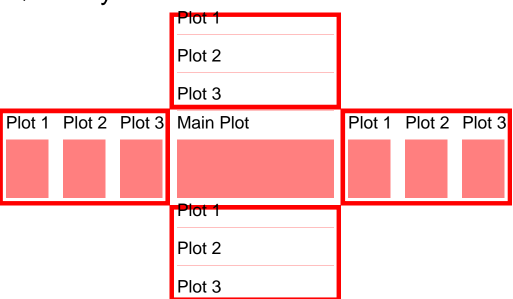
3 heatmap layout

The heatmap_layout() function provides a powerful way to create customizable heatmaps in R using ggplot2. This chapter will guide you through its usage.

heatmap_layout() is a specialized version of quad_alignb(), which itself is a specific variant of QuadLayout (quad_layout()) designed to align observations both horizontally and vertically. We introduce heatmap_layout() directly, as it is more familiar to many users, especially those experienced with popular heatmap packages like pheatmap and ComplexHeatmap.

```
library(ggalign)
## Loading required package: ggplot2
set.seed(123)
small_mat <- matrix(rnorm(56), nrow = 7)
rownames(small_mat) <- paste0("row", seq_len(nrow(small_mat)))
colnames(small_mat) <- paste0("column", seq_len(ncol(small_mat)))</pre>
```

QuadLayout



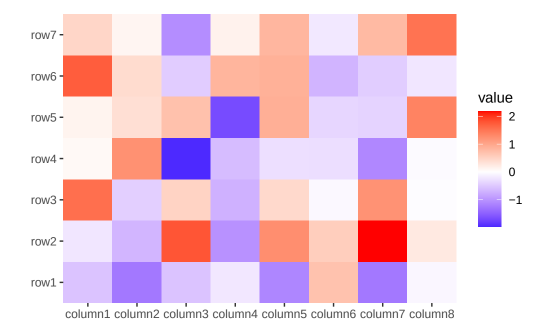
heatmap_layout() simplifies the creation of heatmap plots by integrating essential elements for a standard heatmap layout, ensuring that the appropriate data mapping and visualization layers are automatically applied. ggheatmap() is an alias for heatmap_layout().

3.1 input data

As mentioned in Section 2.1, we typically require a matrix for the Layout which need align observations. Internally, fortify_matrix() will be used to process the data. You can provide a numeric or character vector, a data frame, or any other data type that can be converted into a matrix using as.matrix().

ggheatmap(small_mat)

> heatmap built with `geom_tile()`



3.2 heatmap body

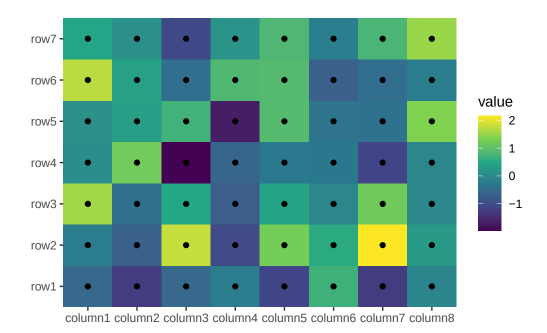
A ggplot object will be automatically created for the heatmap body, the matrix input will be converted into a long formated data frame when drawing. The data in the underlying ggplot object contains following columns:

- .xpanel and .ypanel: the column and row panel
- .x and .y: the x and y coordinates
- .row_names and .column_names: A factor of the row and column names of the original matrix (only applicable when names exist).
- .row_index and .column_index: the row and column index of the original matrix.
- value: the actual matrix value.

The default mapping will use aes(.data\$.x, .data\$.y), but can be customized using mapping argument.

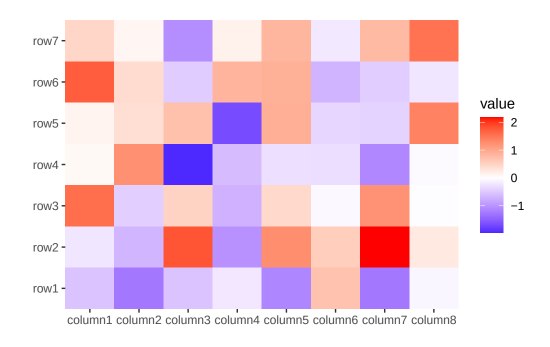
By default, the heatmap body is regarded as the active plot, meaning you can add ggplot2 elements directly to the heatmap body.

> heatmap built with `geom_tile()`

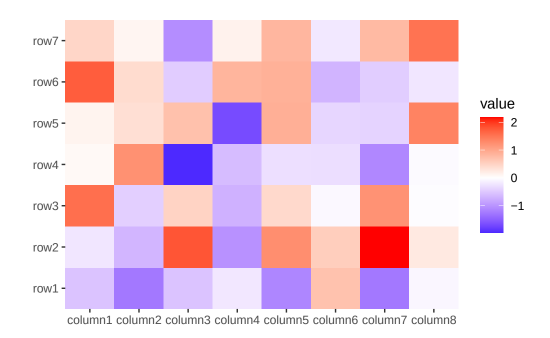


By default, ggheatmap()/heatmap_layout() adds a heatmap layer. If the matrix has more than 20,000 cells (nrow * ncol > 20000), it uses geom_raster() for performance efficiency; for smaller matrices, geom_tile() is used. You can explicitly choose the layer by providing a single string ("raster" or "tile") in the filling argument.

ggheatmap(small_mat, filling = "raster")

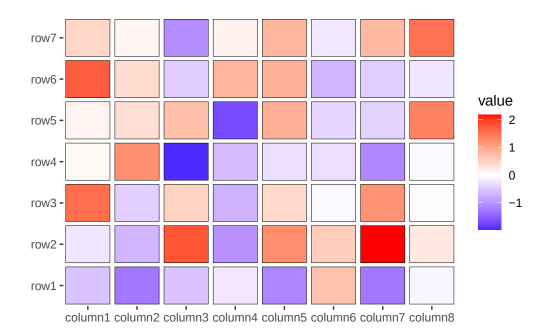


ggheatmap(small_mat, filling = "tile")



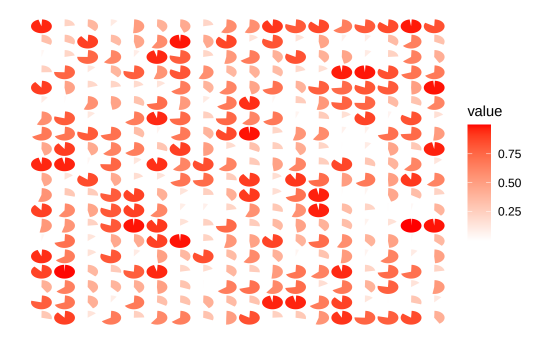
Note, the filling layer will always use mapping of aes(.data\$.x, .data\$.y), if you want to customize filling, you can set filling = NULL, which will remove the filling layer and allow you to add custom filling geoms.

```
ggheatmap(small_mat, filling = NULL) +
  geom_tile(aes(fill = value), color = "black", width = 0.9, height = 0.9)
```



A heatmap pie charts can be easily drawn:

```
set.seed(123)
ggheatmap(matrix(runif(360L), nrow = 20L), filling = NULL) +
    geom_pie(aes(angle = value * 360, fill = value))
```



For more complex customizations of pie charts, you can try using ggforce::geom_arc_bar() instead.

3.3 rasterization

When working with large heatmaps, it's often beneficial to rasterize the heatmap body layer. You can achieve this by using the raster_magick() function. The res argument controls the resolution of the raster image. By default, the res argument matches the resolution of the current device, but specifying a different value can help reduce the resolution of the rasterized heatmap body.

```
ggheatmap(small_mat, filling = NULL) +
  raster_magick(geom_tile(aes(fill = value)), res = 50)
```



By leveraging raster_magick(), you can also perform image post-processing using the magick package. This allows for custom image resizing with filters.



Note: When using magick::image_resize(), you should specify the geometry argument to resize the image. If only the filter is specified, it will only distort the image data (though subtle). For more information on image resizing, refer to ImageMagick's resize documentation.

You can also rasterize all plots in the layout directly with raster_magick(). This method is defined for both ggheatmap()/quad_layout() and stack_layout() objects.

Additionally, You can use external packages like ggrastr or ggfx to rasterize the heatmap body.

```
ggheatmap(small_mat, filling = FALSE) +
    ggrastr::rasterise(geom_tile(aes(fill = value)), dev = "ragg")
```



Likewise, you can also rasterize all plots in the layout directly with ggrastr::rasterise() for both ggheatmap()/quad_layout() and stack_layout().

```
ggrastr::rasterise(ggheatmap(small_mat), dev = "ragg")
```



Furthermore, ggfx offers many image filters that can be applied to ggplot2 layers. See the package for the details.

3.4 annotations

In ggheatmap()/quad_layout(), annotations are handled by a stack_layout() object and can be positioned at the top, left, bottom, or right of the main plot (heatmap body).

By default, ggheatmap()/quad_layout() do not activate an annotation, You can use quad_anno() to activate an annotation, directing all subsequent additions to the specified annotation position. The quad anno() function has the following aliases:

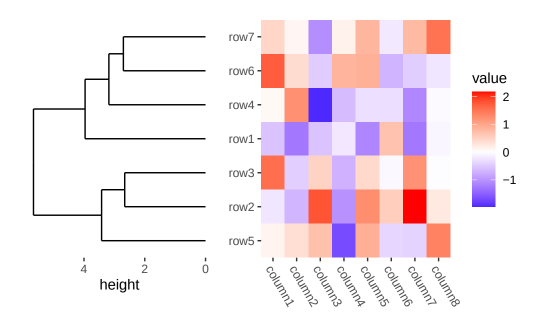
- anno_top: A special case of quad_anno() with position = "top".
- anno left: A special case of quad anno() with position = "left".
- anno_bottom: A special case of quad_anno() with position = "bottom".
- anno right: A special case of quad anno() with position = "right".

When quad_anno() is added to a ggheatmap()/quad_layout(), it will try to automatically create a new stack_layout() (either stack_align() or stack_free()) depending on whether you want to align observations in the specified direction. For top and bottom annotations, stack_alignv() or stack_freev() will be used; for left and right annotations, stack_alignh() or stack_freeh() will be applied.

Additionally, quad_anno() will set the active context to the annotation. This means that subsequent additions will be directed to the annotation rather than the main plot. We use the term active context in contrast to active plot (as described in Chapter 2), since the annotation is a Layout object.

```
ggheatmap(small_mat) +
    theme(axis.text.x = element_text(angle = -60, hjust = 0)) +
    # we set the active context to the left annotation
    anno_left() +
    align_dendro()
```

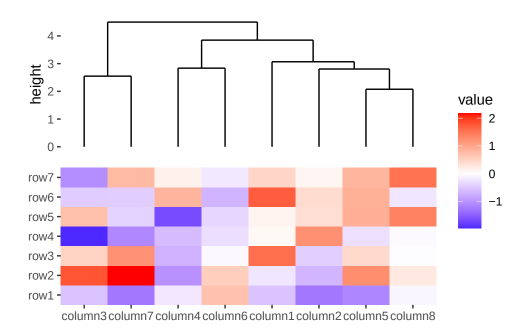
> heatmap built with `geom_tile()`



By default, the annotation stack_layout() will inherit data from ggheatmap()/quad_layout(). If the observations require alignment vertically, this means the data from ggheatmap()/quad_layout() should be a matrix, the column annotations will also require a matrix and the matrix from ggheatmap()/quad_layout() will be transposed for use in the column annotations.

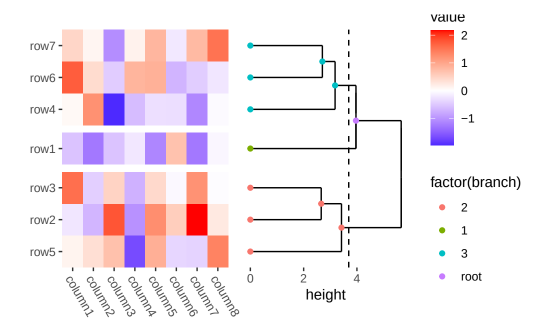
```
ggheatmap(small_mat) +
    # we set the active context to the top annotation
    anno_top() +
    align_dendro()
```

> heatmap built with `geom_tile()`



You can further customize the layout design or add new plots in the annotation stack, as described in Chapter 2.

```
ggheatmap(small_mat) +
    # in the heatmap body, we set the axis text theme
    theme(axis.text.x = element_text(angle = -60, hjust = 0)) +
    # we set the active context to the right annotation
    anno_right() +
    # in the right annotation, we add a dendrogram
    align_dendro(k = 3L) +
    # in the dendrogram, we add a point layer
    geom_point(aes(color = factor(branch)))
```



In this example:

- anno_right() initialize the right annotation stack, and change the active context to the right of the heatmap.
- $align_dendro(k = 3L)$ adds a dendrogram to the annotation and sets itself as the active plot in the annotation stack.
- geom_point(aes(color = factor(branch))) is then added to this active plot within the annotation stack, here, it means the align_dendro() plot.

ggheatmap() aligns observations both horizontally and vertically, so it's safe to always use quad_anno() directly, as all annotations require a matrix, and the layout data is also a matrix. However, for quad_alignh() and quad_alignv(), which only align observations in one direction, the data in the layout may not fit the data for the annotation (when the layout requires alignment of observations, we typically use a matrix, regardless of whether alignment is needed in one or two directions) - quad_alignh(): aligning observations in horizontal direction, for column annotations, we ll need a data frame for stack_free(). - quad_alignv(): aligning observations in vertical direction, for row annotations, we ll need a data frame for stack_free().

In both conditions, quad_anno() won't initialize the annotation by default, instead, you must provide the annotation stack_layout() manually.

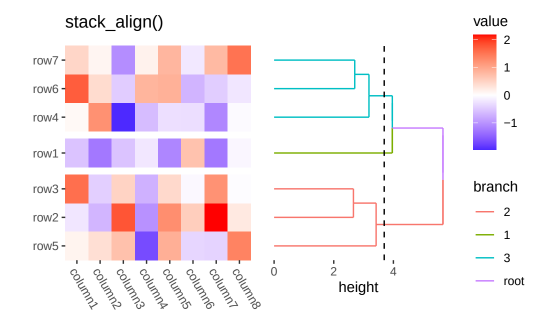
3.5 Adding stack layout

Like adding plot in stack_layout() (Chapter 2), when the direction need alignment, you can add a stack_layout() regardless of whether it need to align observations.

To add a stack_layout() to the ggheatmap(), we must prevent the automatical creation of annotation by quad_anno() by setting initialize = FALSE

```
my_stack_align <- stack_alignh(small_mat) +
    align_dendro(aes(color = branch), k = 3L)

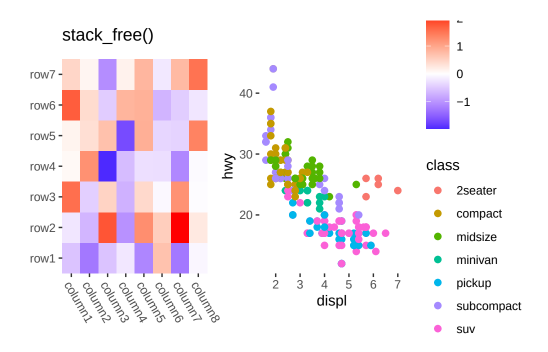
ggheatmap(small_mat) +
    theme(axis.text.x = element_text(angle = -60, hjust = 0)) +
    anno_right(initialize = FALSE) +
    my_stack_align +
    layout_title("stack_align()")</pre>
```



```
my_stack_free <- stack_freeh(mpg) +
    ggfree(mapping = aes(displ, hwy, colour = class)) +
    geom_point(size = 2)
ggheatmap(small_mat) +</pre>
```

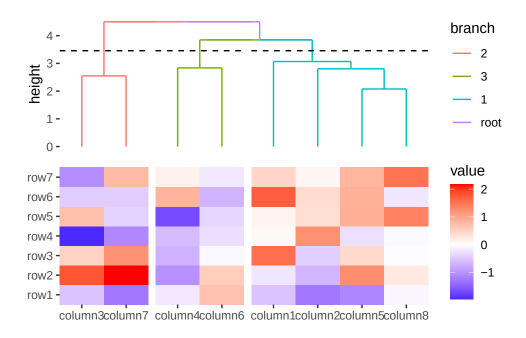
```
theme(axis.text.x = element_text(angle = -60, hjust = 0)) +
anno_right(initialize = FALSE) +
my_stack_free +
layout_title("stack_free()")
```

> heatmap built with `geom_tile()`



Note when aligning the observations, you must ensure the number of observations is consistent in the direction. So for column annotations, you need transpose the data manually.

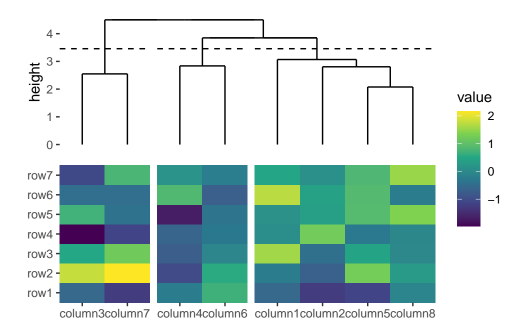
```
my_stack <- stack_alignv(t(small_mat)) +
    align_dendro(aes(color = branch), k = 3L)
ggheatmap(small_mat) +
    anno_top(initialize = FALSE) +
    my_stack</pre>
```



3.6 quad_active()

To remove the active context and redirect additions back to the heatmap body, you can use quad_active().

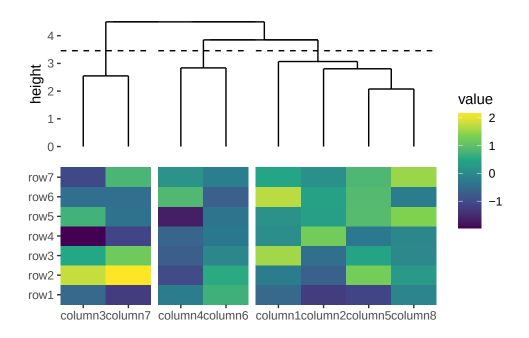
```
ggheatmap(small_mat) +
    # we set the active context to the top annotation
    anno_top() +
    # we split the observations into 3 groups by hierarchical clustering
    align_dendro(k = 3L) +
    # remove any active annotation
    quad_active() +
    # set fill color scale for the heatmap body
    scale_fill_viridis_c()
```



3.7 quad_switch()/hmanno()

We also provide quad_switch()/hmanno() (heatmap annotation) which integrates quad_active() and quad_anno() into one function for ease of use. Feel free to use any of these functions to streamline your annotation process.

```
ggheatmap(small_mat) +
    # we set the active context to the top annotation
    quad_switch("t") +
    # we split the observations into 3 groups by hierarchical clustering
    align_dendro(k = 3L) +
    # remove any active annotation
    quad_switch() +
    # set fill color scale for the heatmap body
    scale_fill_viridis_c()
```



```
ggheatmap(small_mat) +
    # we set the active context to the top annotation
    hmanno("t") +
    # we split the observations into 3 groups by hierarchical clustering
    align_dendro(k = 3L) +
    # remove any active annotation
    hmanno() +
    # set fill color scale for the heatmap body
    scale_fill_viridis_c()
```



3.8 Plot Size

3.8.1 Heatmap Body Size

You can specify the relative sizes of the heatmap body using the width and height arguments in the ggheatmap() function.

```
ggheatmap(small_mat, height = 2) +
  anno_top() +
  align_dendro()
```



Alternatively, the quad_active() function allows you to control the heatmap body sizes.

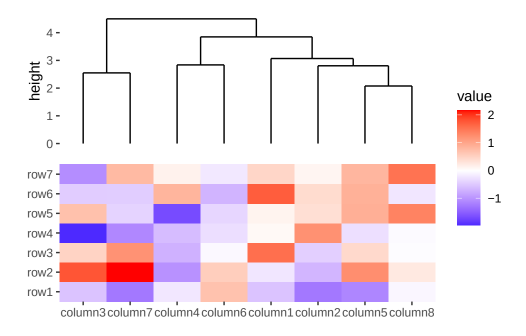
```
ggheatmap(small_mat) +
  quad_active(height = 2) +
  anno_top() +
  align_dendro()
```



3.8.2 Annotation Stack Size

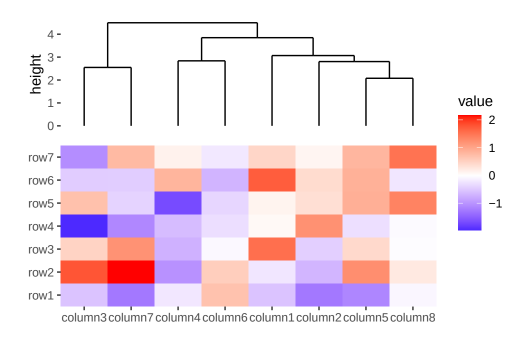
The quad_anno() function allows you to control the total annotation stack size. The size argument controls the relative width (for left and right annotations) or height (for top and bottom annotations) of the whole annotation stack.

```
ggheatmap(small_mat) +
  anno_top(size = 1) +
  align_dendro()
```



You can also specify it as an absolute size using unit():

```
ggheatmap(small_mat) +
  anno_top(size = unit(30, "mm")) +
  align_dendro()
```



4 Layout customize

For layouts that can align observations, the package provides a suite of align_* functions designed to give you precise control over the layout. These functions allow you to reorder observations or partition them into multiple groups.

Currently, there are four key align_* functions available for layout customization:

- align_group: Group and align plots based on categorical factors.
- align_order: Reorder layout observations based on statistical weights or allows for manual reordering based on user-defined ordering index.
- align_kmeans: Group observations by k-means clustering results.
- align_hclust: Reorder or group observations based on hierarchical clustering.

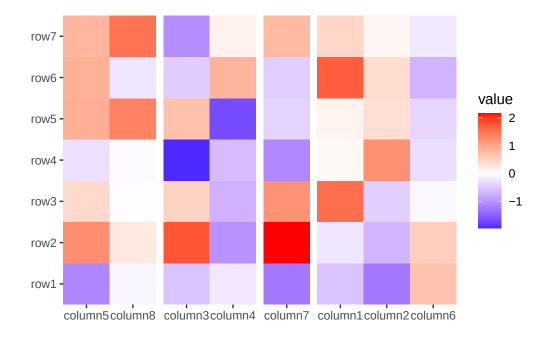
Note that none of these functions add a plot area or set the active context, meaning you cannot add ggplot2 elements to these objects.

```
library(ggalign)
## Loading required package: ggplot2
set.seed(123)
small_mat <- matrix(rnorm(56), nrow = 7)
rownames(small_mat) <- paste0("row", seq_len(nrow(small_mat)))
colnames(small_mat) <- paste0("column", seq_len(ncol(small_mat)))</pre>
```

4.1 align_group()

The align_group() function allows you to split the observations into groups.

```
set.seed(1234)
ggheatmap(small_mat) +
   anno_top() +
   align_group(sample(letters[1:4], ncol(small_mat), replace = TRUE))
```



Note that all align_* functions which split observations into groups must not break the previous established groups. This means the new groups must nest in the old groups, usually they cannot be used if groups already exist.

```
set.seed(1234)
ggheatmap(small_mat) +
   anno_top() +
   align_group(sample(letters[1:4], ncol(small_mat), replace = TRUE)) +
   align_group(sample(letters[1:5], ncol(small_mat), replace = TRUE))
```

Error in `align()`:
! align_group(sample(letters[1:5], ncol(small_mat), replace = TRUE))
 disrupt the previously established panel groups of the top annotation
 `stack_align()`

4.2 align_order()

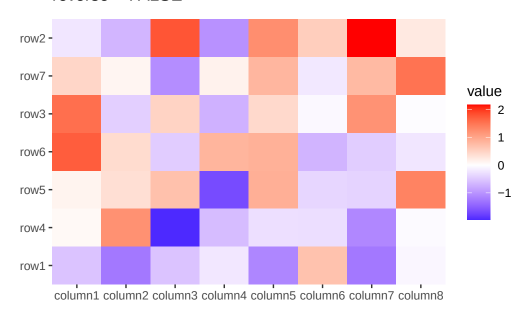
The align order() function reorder the observations based on the summary weights.

In this example, we order the rows based on their means. By default, the ordering is in ascending order according to the summary weights. You can reverse the order by setting reverse = TRUE.

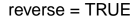
```
ggheatmap(small_mat) +
  anno_left() +
  align_order(rowMeans) +
  layout_title(title = "reverse = FALSE")
```

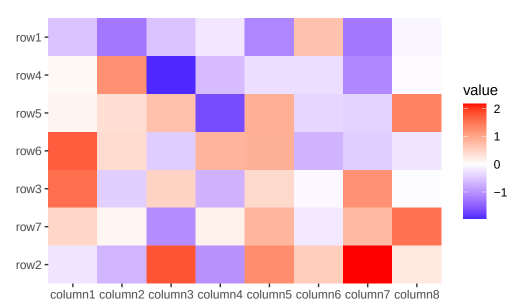
> heatmap built with `geom_tile()`

reverse = FALSE



```
ggheatmap(small_mat) +
  anno_left() +
  align_order(rowMeans, reverse = TRUE) +
  layout_title(title = "reverse = TRUE")
```



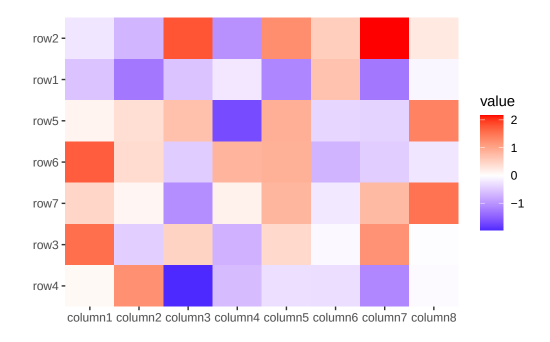


Additionally, you can provide the ordering integer index or character index directly:

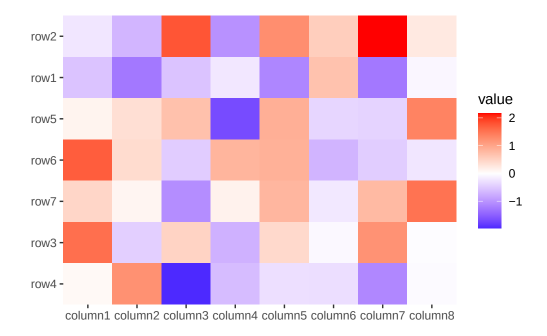
```
my_order <- sample(nrow(small_mat))
print(rownames(small_mat)[my_order])</pre>
```

```
[1] "row4" "row3" "row7" "row6" "row5" "row1" "row2"
```

```
ggheatmap(small_mat) +
  anno_left() +
  align_order(my_order)
```



```
ggheatmap(small_mat) +
  anno_left() +
  align_order(rownames(small_mat)[my_order])
```



Some align_* functions also accept a data argument. It's important to note that all align_* functions treat rows as the observations. This means NROW(data) must match the number of observations along the axis used for alignment. The data argument can also accept a function (supporting purrr-like lambda syntax), which will be applied to the layout matrix.

As mentioned in Chapter 3, for top and bottom annotations, the data matrix of quad_layout()/ggheatmap() is transposed to create the annotation stack_layout(). Therefore, you can use rowMeans() to calculate the mean value across all columns.

```
ggheatmap(small_mat) +
  anno_top() +
  align_order(rowMeans)
```



Some align_* functions that reorder observations include an argument called strict. This argument is especially useful when previous groups have already been established. If previous groups have been created and strict = FALSE, the function will reorder the observations within each group.

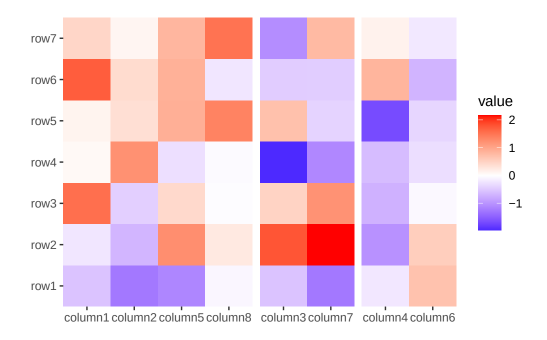
```
set.seed(1234)
ggheatmap(small_mat) +
   anno_top() +
   align_group(sample(letters[1:4], ncol(small_mat), replace = TRUE))+
   align_order(rowMeans, strict = FALSE)
```



4.3 align_kmeans()

The align_kmeans() function split the observations into groups based on k-means clustering.

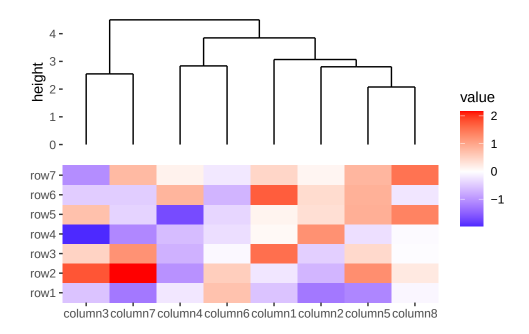
```
ggheatmap(small_mat) +
  anno_top() +
  align_kmeans(3L)
```



4.4 align_hclust()

The <code>align_dendro()</code> function adds a dendrogram to the layout and can also reorder or split the layout based on hierarchical clustering. This is particularly useful for working with heatmap plots.

```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro()
```



Hierarchical clustering is performed in two steps: calculate the distance matrix and apply clustering. You can use the distance and method argument to control the dendrogram builind process.

There are two ways to specify distance metric for clustering:

- specify distance as a pre-defined option. The valid values are the supported methods in dist() function and coorelation coefficient "pearson", "spearman" and "kendall". The correlation distance is defined as 1 cor(x, y, method = distance).
- a self-defined function which calculates distance from a matrix. The function should only contain one argument. Please note for clustering on columns, the matrix will be transposed automatically.

```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro(distance = "pearson") +
  patch_titles(top = "pre-defined distance method (1 - pearson)")
```

pre-defined distance method (1 – pearson) 1.5 - to be a second or second or

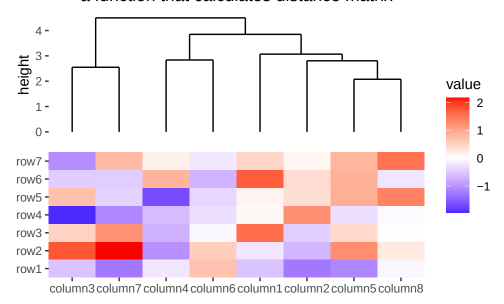
column1 column4 column2 column8 column6 column3 column5 column7

ggheatmap(small_mat) +
 anno_top() +
 align_dendro(distance = function(m) dist(m)) +
 patch_titles(top = "a function that calculates distance matrix")

> heatmap built with `geom_tile()`

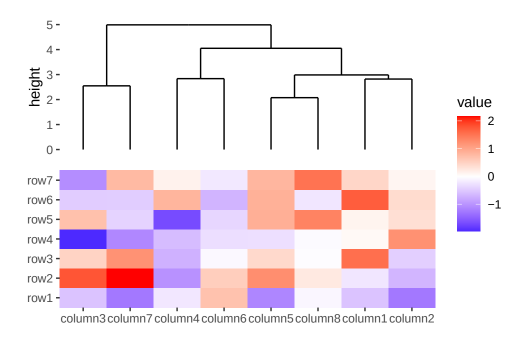
row1 -

a function that calculates distance matrix



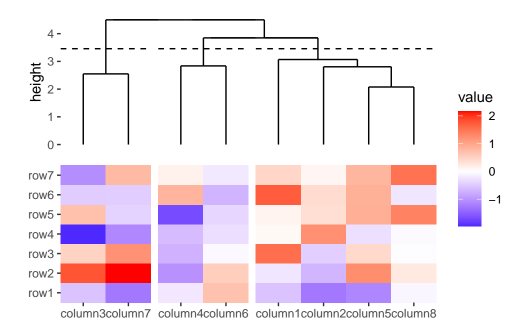
Method to perform hierarchical clustering can be specified by method. Possible methods are those supported in hclust() function. And you can also provide a self-defined function, which accepts the distance object and return a hclust object.

```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro(method = "ward.D2")
```



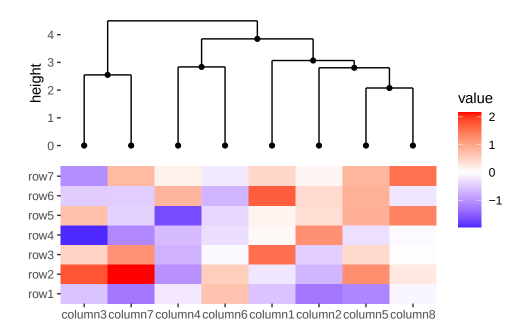
The dendrogram can also be used to cut the columns/rows into groups. You can specify k or h, which work similarly to cutree():

```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro(k = 3L)
```



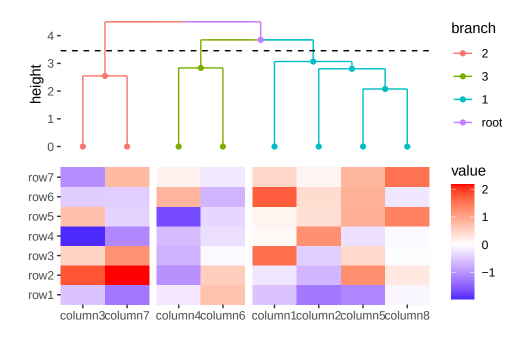
In contrast to align_group(), align_kmeans(), and align_order(), align_dendro() is capable of drawing plot components. So it has a default set_context value of TRUE, meaning it will set the active context of the annotation stack layout. In this way, we can add any ggplot elements to this plot area.

```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro() +
  geom_point(aes(y = y))
```



The align_dendro() function creates default node data for the ggplot. See ggplot2 specification in ?align_dendro for details. Additionally, edge data is added to the ggplote::geom_segment() layer directly, used to draw the dendrogram tree. One useful variable in both node and edge data is the branch column, corresponding to the cutree result:

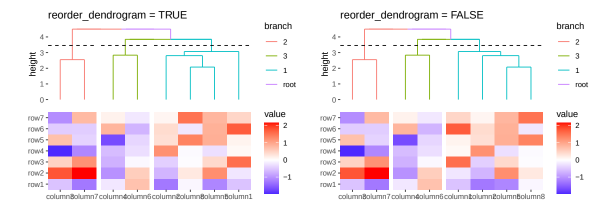
```
ggheatmap(small_mat) +
  anno_top() +
  align_dendro(aes(color = branch), k = 3) +
  geom_point(aes(color = branch, y = y))
```



You can reorder the dendrogram based on the mean values of the observations by setting reorder_dendrogram = TRUE.

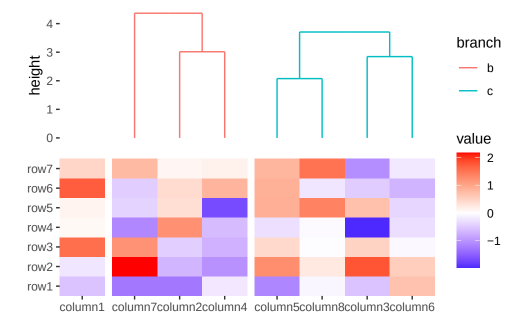
```
h1 <- ggheatmap(small_mat) +
    anno_top() +
    align_dendro(aes(color = branch), k = 3, reorder_dendrogram = TRUE) +
    ggtitle("reorder_dendrogram = TRUE")
h2 <- ggheatmap(small_mat) +
    anno_top() +
    align_dendro(aes(color = branch), k = 3) +
    ggtitle("reorder_dendrogram = FALSE")
align_plots(h1, h2)</pre>
```

- > heatmap built with `geom_tile()`
- > heatmap built with `geom_tile()`



align_dendro() can also perform clustering between groups, meaning it can be used even if there are existing groups present in the layout, in this way, you cannot specify k or h:

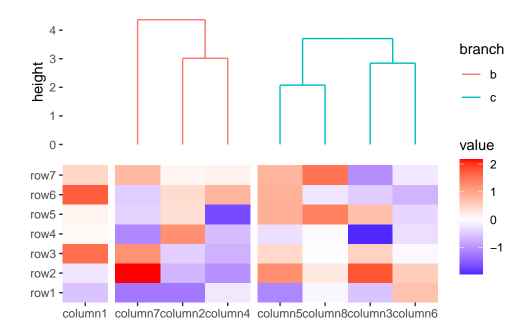
```
set.seed(3L)
column_groups <- sample(letters[1:3], ncol(small_mat), replace = TRUE)
ggheatmap(small_mat) +
   anno_top() +
   align_group(column_groups) +
   align_dendro(aes(color = branch))</pre>
```



You can reorder the groups by setting reorder_group = TRUE.

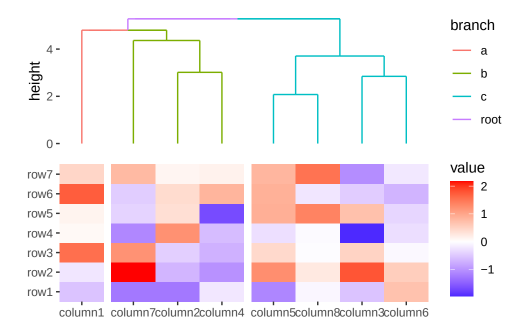
```
ggheatmap(small_mat) +
  anno_top() +
  align_group(column_groups) +
  align_dendro(aes(color = branch), reorder_group = TRUE)
```

> heatmap built with `geom_tile()`



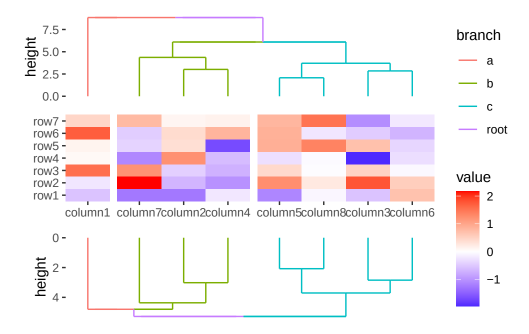
You can merge the sub-tree in each group by settting merge_dendrogram = TRUE.

```
ggheatmap(small_mat) +
  anno_top() +
  align_group(column_groups) +
  align_dendro(aes(color = branch), merge_dendrogram = TRUE)
```



You can reorder the dendrogram and merge simutaneously.

```
ggheatmap(small_mat) +
   anno_top() +
   align_group(column_groups) +
   align_dendro(aes(color = branch),
        reorder_group = TRUE,
        merge_dendrogram = TRUE
) +
   anno_bottom() +
   align_dendro(aes(color = branch),
        reorder_group = FALSE,
        merge_dendrogram = TRUE
)
```



If you specify k or h, this will always turn off sub-clustering. The same principle applies to align_dendro(), where new groups must be nested within the previously established groups.

```
ggheatmap(small_mat) +
  anno_top() +
  align_group(column_groups) +
  align_dendro(k = 2L)
```

Error in `align()`:

! align_dendro(k = 2L) disrupt the previously established panel groups
 of the top annotation `stack_align()`