Make Interactive Complex Heatmaps in R

Zuguang Gu
German Cancer Research Center

Daniel Hübschmann
German Cancer Research Center

Abstract

Heatmap is a powerful visualization method on two-dimensional data to reveal patterns shared by subsets of rows and columns. In R, there are many packages that make heatmaps. Among them, ComplexHeatmap provides rich tools for constructing highly customizable heatmaps. It can easily establish connections between information from multiple sources by automatically concatenating and adjusting multiple heatmaps as well as complex annotations, which makes it widely applied in data analysis in various fields, especially in Bioinformatics. Nevertheless, the limit of ComplexHeatmap still exists. It only generates static plots which restricts deeper inspections on complex heatmaps, e.g., to look into a subset of rows and columns when a specific pattern of interest is observed from the heatmap. In this work, we described a new R/Bioconductor package InteractiveComplexHeatmap that brings interactivity to ComplexHeatmap. InteractiveComplexHeatmap is designed with an easy-to-use interface where static complex heatmaps can be directly exported to an interactive Shiny web application only with one extra line of code. The interactive application contains comprehensive tools for manipulating heatmaps. Besides common tools as in other interactive heatmap packages, InteractiveComplexHeatmap additionally supports, e.g., selecting over multiple heatmaps or searching heatmaps via row or column labels. Also, InteractiveComplexHeatmap provides methods to export static heatmaps from other popular heatmap functions, e.g., heatmap.2() or pheatmap(), to interactive heatmap applications. Finally, InteractiveComplexHeatmap provides functionalities for integrating interactive heatmap widgets into other Shiny applications. InteractiveComplexHeatmap provides a user interface for self-defining response to selection events on heatmaps, which helps to implement more complex Shiny web applications.

Keywords: visualization, interactive heatmap, complex heatmap, Shiny, R.

1. Introduction

Heatmap is a popular visualization method on a two-dimensional matrix where colors are the major aesthetic elements mapping to data (Wilkinson and Friendly 2009). In routine data analysis procedures, matrix for heatmap visualization is normally accompanied with row and column reordering, e.g., by hierarchical clustering or seriation (Hahsler, Hornik, and Buchta 2008), so that features with similar patterns are grouped closely and they can be
easily identified from the colors on heatmap. The reordering methods can also be chosen specifically for the topic under study. For example, the R package EnrichedHeatmap (Gu, Eils, Schlesner, and Ishaque 2018) can be used to visualize how DNA demethylation is enriched around the transcriptional start sites (TSS) of genes by a heatmap where columns are ordered by genomic distances from CpG dinucleotides to gene TSS and rows are ordered by the enrichment score defined by EnrichedHeatmap. Heatmap is widely applied in various research fields, especially popular in Bioinformatics. Since the first paper on heatmap visualization on a gene expression dataset was published (Eisen, Spellman, Brown, and Botstein 1998), heatmap has been a standard tool for visualizing e.g., gene expression, DNA methylation and other high-throughput datasets represented as matrices. In R (R Core Team 2020), there are several packages that provide functions for implementing heatmaps. The function heatmap() from package stats is the most fundamental one, but with very limited functionality. The function heatmap.2() from package gplots (Warnes, Bolker, Bonebakker, Gentleman, Huber, Liaw, Lumley, Maechler, Magnusson, Moeller, Schwartz, and Venables 2020) is an enhanced version of heatmap() which supports more graphics on heatmap, such as a color legend with value distribution and trace lines showing the difference of values to the column or row medians. There are also packages implemented with the grid graphics system such as the function pheatmap() from package pheatmap (Kolde 2019) and the function aheatmap() from package NMF (Gaujoux and Seoighe 2010). They provide more flexible controls on heatmaps.

As data emerges fast in dimensions nowadays, especially in the Genomics field, an efficient visualization for integrative analysis or multi-omics analysis is urgently needed to associate multiple types of data for easily revealing relationships between multiple objects. From the aspects of heatmap visualization, it reflects in two points. The first is the support of heatmap annotation which contains extra information to associate to the main heatmap. For example, in a typical heatmap visualization on gene expression data where rows are genes and columns are patients, it is common that patients have clinical meta-data available, such as age, gender or whether the patient has certain DNA mutations. With annotation attached to heatmap, it is easy to identify, e.g., whether a group of genes showing high expression correlate to a certain age interval or whether they have specific types of DNA mutations. Functions heatmap() and heatmap.2() only support single heatmap-like annotation for one numeric or character vector. pheatmap() and aheatmap() allow multiple heatmap-like annotations for corresponding more information to heatmap. The package superheat (Barter and Yu 2018) and heatmap3 (Zhao, Yin, Guo, Sheng, and Shyr 2021) supports more types of graphics for annotations, such as points or lines which are able to make more accurate visual representation on annotation data. The second point of visualizing multiple sources of information is to directly apply “complex heatmap visualization” by simultaneously linking multiple heatmaps. For example, in our published study (Gu et al. 2018), we applied complex heatmap visualization on gene expression, DNA methylation and various histone modifications to reveal general transcriptional regulation patterns among multiple human tissues. To implement both complex annotation and heatmap visualization, we previously developed an advanced heatmap package ComplexHeatmap (Gu, Eils, and Schlesner 2016). It supports not only the basic annotation graphics as in other packages, but also a variety of extra annotation graphics such as violin plot or horizon plot. ComplexHeatmap provides a simple syntax to link multiple heatmaps where rows or columns of all heatmaps are adjusted simultaneously. The simplicity of its user interface and comprehensiveness of the functionalities make ComplexHeatmap widely used in Bioinformatics to reveal interesting patterns from data that are
potentially biologically meaningful.

After a specific pattern of subset of rows and columns is observed from the heatmap, the next step is to extract corresponding rows and columns for downstream analysis, which requires interactivity on heatmaps. There are also several R packages that implement interactive heatmaps such as d3heatmap (Cheng and Galili 2018), heatmaply (Galili, O’Callaghan, Sidi, and Sievert 2017) and iheatmap (Schep and Kummerfeld 2017). The interactivity allows hovering on heatmap cells and selecting a region from heatmap. There are other web-based and non-R implementations, such as Morpheus (Morpheus 2021), Heatmapper (Babicki, Arndt, Marcu, Liang, Grant, Maciejewski, and Wishart 2016), Clustergrammer (Fernandez, Gundersten, Rahman, Grimes, Rikova, Hornbeck, and Ma’ayan 2017) and NG-CHM (Ryan, Stucky, Wakefield, Melott, Akbani, Weinstein, and Broom 2019) which allow users to directly upload data to the web server and manipulate heatmap with no programming knowledge.

ComplexHeatmap is popular and powerful for generating static complex heatmaps. Here we developed a new package InteractiveComplexHeatmap that brings interactivity to ComplexHeatmap. InteractiveComplexHeatmap can be very easily applied that static complex heatmaps are directly exported to an interactive Shiny web application only with one extra line of code. The interactive application contains comprehensive tools for manipulating heatmaps. Besides common tools as in other interactive heatmap packages, InteractiveComplexHeatmap additionally supports, e.g., selecting over multiple heatmaps or searching heatmaps via row or column labels. The latter would be especially useful when users already have some features of interest to look into. Also, InteractiveComplexHeatmap provides methods to export static heatmaps from other popular heatmap functions, e.g., heatmap.2() or pheatmap(), to interactive heatmap applications, which greatly expands the ability of interactive heatmap visualization in R. Finally, InteractiveComplexHeatmap provides functionalities for integrating interactive heatmap widgets into other Shiny applications. InteractiveComplexHeatmap provides a user interface for self-defining response to selection events on heatmaps, e.g., by clicking or brushing, which helps to develop more complex Shiny web applications.

The paper is structured as follows, In Section 2, we first made a brief introduction on the functionalities in ComplexHeatmap. In Section 3, we described how to export static heatmaps into an interactive Shiny web application and the tools provided by the interactive application. In Section 4, we explained how the interactivity is implemented in InteractiveComplexHeatmap. And finally, in Section 5, we explained in detail how to integrate InteractiveComplexHeatmap to Shiny application development.

2. A brief introduction on ComplexHeatmap

In this section, we briefly introduce basic functionalities of making single heatmaps and a list of heatmaps with ComplexHeatmap. For more comprehensive usages, readers are recommended to refer to the ComplexHeatmap complete reference (Gu 2021).

ComplexHeatmap is implemented with the grid graphics engine under S4 object-oriented system. There are three major classes defined in the package: a Heatmap class that defines a single heatmap, a HeatmapList class that defines a list of heatmaps, and a HeatmapAnnotation class that defines a list of heatmap annotations.

2.1. A single heatmap
The function `Heatmap()` makes a single heatmap and returns an object in `Heatmap` class. The only mandatory argument for `Heatmap()` is a matrix, either numeric or character. It provides numerous additional arguments to customize heatmap. Besides common functionalities as in other heatmap functions, `Heatmap()` has following major features:

- **Flexible controls of clustering.** The clustering can be specified in multiple ways, *i.e.*, 1. by a pre-defined distance method such as "euclidean" or "pearson", 2. by a distance function that calculates pairwise distance between two vectors or directly from a matrix, 3. by a clustering object *e.g.* a `hclust` or `dendrogram` object or object that can be coerced to `dendrogram` by a proper `as.dendrogram()` function, or 4. by a clustering function that takes matrix as input and a `dendrogram` object as output. Besides that, dendrogram can be rendered on both edges and nodes, *e.g.*, to assign different colors for dendrogram branches or to add symbols on dendrogram nodes (Figure 1A).

- **Split heatmap.** Heatmap splitting is an efficient way to highlight group-wise patterns (Figure 1D). There are multiple ways to split heatmap on rows or columns. 1. Set a number for `k`-means clustering; 2. Set a categorical variable which can be a vector or a data frame, then heatmap is split by all combinations of levels in the categorical variable; 3. If hierarchical clustering is already applied, the splitting can also be specified as a single number so that `cutree()` is internally applied to split.

- **Flexible controls of colors and legends.** It allows exact mapping between colors and values in matrix with a color mapping function by specifying breaks and corresponding colors, then remaining colors are linearly interpolated in the corresponding intervals. It also allows very flexible configurations on heatmap legends, such as multiple color scheme legends. Please refer to Section “Legends” in `ComplexHeatmap` complete reference (Gu 2021) for more examples.

- **Render heatmap body as a raster image.** For a heatmap on a huge matrix that is saved as vector graphics (*e.g.*, a `pdf` figure), rasterization helps to efficiently reduce the final file size while the loss of figure quality is ignorable. `Heatmap()` supports various methods for image rasterization. See detailed explanations and comparisons in (Gu 2020a).

- **Customize heatmap.** There are two ways for customizing heatmap with user-defined code: 1. to customize heatmap body via `cell_fun` or `layer_fun` argument to add self-defined graphics to heatmap cells when heatmap is drawing (Figure 1B), and 2. to use `decorate_*()` family functions, *e.g.*, `decorate_annotation()`, to add graphics to any heatmap component after the heatmap is drawn.

### 2.2. Heatmap annotations

Heatmap annotations are important components of heatmap that show additional information associated with rows or columns in a heatmap. `ComplexHeatmap` provides very flexible support for setting annotations and defining new annotation graphics. The annotations can be put on the four sides of the heatmap and they are automatically reordered and split according to heatmap.

There are following annotation graphics demonstrated in Figure 1C (from left to right):

- **Heatmap-like annotation.** It is named as “simple annotation” in `ComplexHeatmap`. It visualizes a vector or a matrix, either numeric or character.
Figure 1: Demonstration of **ComplexHeatmap**. A) A heatmap with both column and row annotations. The columns on the heatmap were split by a three-group $k$-means clustering and rows were split by combination of a categorical variable and a two-group $k$-means clustering. B) A heatmap with customizations. On the heatmap, horizontal neighbour cells were connected if they had the same signs. Black borders were added to the top right and bottom left heatmap slices. C) Examples of various annotation graphics supported in **ComplexHeatmap**. D) An example of complex heatmap visualization based on a real-world dataset.
• **Image annotation.** It supports images in various formats, e.g., png, svg, pdf, jpg.
• **Points annotation.** It supports a single numeric vector or a numeric matrix.
• **Lines annotation.** It supports a single numeric vector or a numeric matrix. Additionally, it supports to perform Loess smoothing over data points.
• **Barplot annotation.** It also supports stacked bar plots.
• **Boxplot annotation.**
• **Text annotation.** It supports constructing more customized text with package gridtext (Wilke 2020).
• **Histogram annotation.**
• **Violin annotation.** It visualizes a list of distributions. Alternatively, the distributions can also be visualized by normal density plots or heatmaps.
• **Joy plot annotation.**
• **Horizon annotation.**

All built-in annotation graphics are implemented by annotation functions named with `anno_` prefix, e.g., `anno_points()` for points annotation. Besides the above listed annotations, ComplexHeatmap supports more complex annotations. For example, there is a “mark annotation” (`anno_mark()`) which draws labels for a subset of rows or columns where the labels are shifted from their original positions to get rid of overlapping and lines are drawn to connect labels to their corresponding rows or columns. ComplexHeatmap provides a user interface for creating new annotation graphics. For example, package EnrichedHeatmap defined an “enriched annotation” (by `anno_enriched()`) which visualizes average enrichment of certain genomic signals over a list of genomic features, and package simplifyEnrichment (Gu 2020b) defined a “word cloud annotation” (by `anno_word_cloud()`) which visualizes the summaries of groups of rows by word clouds.

The function `HeatmapAnnotation()` accepts multiple annotations specified as name-value pairs. Simple annotations are specified as vectors or a data frame and legends are automatically generated for them. Other annotations should be specified via functions `anno_•()`.

```r
R> ha = HeatmapAnnotation(  + foo = runif(10),  + bar = sample(letters[1:4], 10, replace = TRUE),  + pt = anno_points(runif(10)),  + txt = anno_text(month.name[1:10]),  + ...  + )
```

Row annotations should be set with one additional argument `which = "row"` or by the helper function `rowAnnotation()`. Column annotations are assigned to `top_annotation` or `bottom_annotation` and row annotations are assigned to `left_annotation` or `right_annotation` in `Heatmap()`.

### 2.3. A list of heatmaps

The main feature of ComplexHeatmap is that it supports concatenating a list of heatmaps and annotations so that it is possible to visualize associations from various sources of information.
ComplexHeatmap provides a simple syntax for concatenating heatmaps with the operator `+`. The expression returns a HeatmapList object.

```
R> ht_list = Heatmap(...) + Heatmap(...) + rowAnnotation(...)
R> ht_list
```

We previously introduced annotations as components of a single heatmap, here the row annotations can also be independently concatenated to the heatmap list. Alternatively, which is less used, the heatmap lists can be vertically concatenated with the operator `%v%`.

```
R> ht_list = Heatmap(...) %v% Heatmap(...) %v% HeatmapAnnotation(...)
R> ht_list
```

The amount of heatmaps and annotations to be concatenated can be arbitrary. The ordering and the splitting of all heatmaps are adjusted to the main heatmap, which is by default the first numeric matrix, or other heatmap in the list that is specified by the user.

### 2.4. High-level plots implemented with ComplexHeatmap

The flexibility of ComplexHeatmap allows users to implement new high-level graphics with a matrix-like data structure. In ComplexHeatmap, there are following high-level graphics functions already implemented:

- `densityHeatmap()`. It visualizes a list of distributions via heatmap.
- `oncoPrint()`. It visualizes multiple genomic alteration events (e.g. single-base mutations or fragment deletions) in a group of genes and in multiple patients.
- `UpSet()`. It visualizes intersections over multiple sets. This is an enhanced version of package UpSetR (Conway, Lex, and Gehlenborg 2017). UpSet() is also able to visualize intersections over genomic intervals.

More importantly, the three functions all return Heatmap objects, thus, the graphics can be concatenated to additional heatmaps or annotations to construct a complex view of the data. As an example, an oncoPrint can be concatenated to a gene expression heatmap to quickly connect the relationship between DNA mutations and the influence on gene expression. Example code is as follows:

```
R> oncoPrint(...) + Heatmap(...)
```

### 2.5. A complex example

Figure 1D demonstrates complex heatmap visualization on a dataset randomly generated but based on patterns found in an unpublished work. The figure visualizes associations between DNA methylation, gene expression, enhancers and gene-related information. In heatmaps, each row corresponds to a differentially methylated region (DMR, which is a genomic region showing significantly different methylation between control and tumor samples) or other objects that are associated with the DMR. In Figure 1D, there are following heatmaps from left to right:
1. A heatmap of methylation level in DMRs. Red corresponds to 100% methylation in the DMR and blue corresponds to no methylation.

2. A one-column heatmap showing the direction of differential methylation. hyper means methylation is higher in tumor samples than control and hypo means methylation is lower in tumor samples.

3. A heatmap of gene expression. Each gene is the nearest gene in the genome with expression negatively correlating to the methylation level in the associated DMR.

4. A one-column heatmap of p-values from Pearson correlation test between gene expression and methylation.

5. A one-column heatmap of the gene type, i.e., whether the corresponding gene is a protein-coding gene or of other type.

6. A one-column heatmap of the annotation of DMRs to genes, i.e., does the DMR locate in gene TSS or outside gene regions?

7. A one-column heatmap of the genomic distance between DMR to TSS of the associated gene.

8. A heatmap showing the overlap between enhancers and DMRs. Enhancers are genomic regions where proteins can bind to regulate gene expression.

The heatmap list is split by the combination of differential methylation directions and a two-group k-means clustering. The latter is to distinguish high-methylation groups and low methylation groups. In Figure 1D, the complex heatmaps reveal that highly methylated DMRs are enriched in intergenic and intragenic regions and rarely overlap with enhancers (row group "2,hypo" and "2,hyper"), while in contrast, lowly methylated DMRs are enriched in transcription start sites and enhancers (row group "1,hypo" and "1,hyper"). This might imply that enhancers associate with low methylation and methylation changes in enhancers might affect their transcriptional activities on related genes.

3. Export heatmap to a Shiny web application

3.1. Usage

For any single heatmap or a list of heatmaps represented as a Heatmap or a HeatmapList object produced by ComplexHeatmap, the function htShiny() in InteractiveComplexHeatmap can be simply applied to export it to a Shiny web application. To demonstrate the usage, we generated a list of two heatmaps of a numeric heatmap and a character heatmap. In the numeric heatmap, k-means clustering with two groups was applied on both rows and columns. A point annotation was put on top of the first heatmap as a heatmap component and a barplot annotation was inserted between the first and the second heatmaps.

```R
R> library(ComplexHeatmap)
R> set.seed(123)
R> mat1 = matrix(rnorm(100), 10)
R> rownames(mat1) = colnames(mat1) = paste0("a", 1:10)
R> mat2 = matrix(sample(letters[1:10], 100, replace = TRUE), 10)
R> rownames(mat2) = colnames(mat2) = paste0("b", 1:10)
R>
```
R> ht_list = Heatmap(mat1, name = "mat_a", row_km = 2, column_km = 2,  
+ top_annotation = HeatmapAnnotation(foo = anno_points(runif(10))) +  
+ rowAnnotation(bar = anno_barplot(sample(10, 10))) +  
+ Heatmap(mat2, name = "mat_b")

After the heatmap object `ht_list` is created, it can be directly sent to `htShiny()`. The interactive heatmap application is automatically opened in a web browser or in a pop-up window in RStudio IDE (Figure 2).

R> ht_list = draw(ht_list) # not necessary, but recommended
R>
R> library(InteractiveComplexHeatmap)
R> htShiny(ht_list)

To use `htShiny()`, the heatmap object is recommended to be updated with the function `draw()`. If it has not been updated, it will be applied inside `htShiny()` automatically. Updating by `draw()` speeds up loading the Shiny application because `draw()` applies clusterings which is normally the most time-consuming step in heatmap generation. After `draw()` is executed, clustering results are saved in the returned heatmap object so that repeatedly drawing heatmap can directly use the saved clustering results. If the heatmap includes randomness, such as k-means clustering by setting `row_km` or `column_km` argument or using random colors for annotations, it is necessary to execute `draw()` before sending the heatmap object to `htShiny()` to get rid of obtaining different heatmaps when executing `htShiny()` multiple times.

Executing the function `htShinyExample()` with no argument prints a list of 44 examples of various usages of interactive heatmaps. Specifying an index in `htShinyExample()` runs the corresponding interactive application as well as showing the source code that generates the application, e.g., `htShinyExample(1.5)` demonstrates an example of exporting a list of two heatmaps to an interactive web application.

### 3.2. Tools in the interactive heatmap

In the Shiny application illustrated in Figure 2, there are three main components: the original heatmap, the selected sub-heatmap and the output that prints information of the heatmap cell that was clicked or the sub-heatmap that was selected. In the original heatmap, users can click on it or select an area from it. If an area is selected, a sub-heatmap is drawn on the right side in the application. Both heatmaps can be resized by dragging from their bottom right.

Under original heatmap, there are several tools integrated:

- **Search heatmap**: If heatmaps have row or column labels, it allows users to search heatmap labels to obtain a subset of rows and columns. The keywords can be exactly matched to row or column labels or a regular expression. Once the corresponding rows or columns are found in heatmaps, a sub-heatmap is drawn in the right sub-heatmap component.

- **Configure brush**: It configures the visual style of the brush which selects the area from heatmap, i.e., border color and background color, border width and opacity of the brush.
Figure 2: A demonstration of interactive complex heatmap. On the left is the original heatmap and on the right is the sub-heatmap that was selected from left. Below both heatmaps are the tools for controlling heatmaps.
• **Save image**: The main heatmap can be saved into a file in one of the three formats: png, pdf and svg.

• **Resize image**: The size of heatmap can be precisely controlled by manually entering values for it.

Under sub-heatmap, there are also tools for controlling the sub-heatmap.

• **Configure sub-heatmap**: There are three sections of controls. 1. Basic controls such as whether to show row or column names, 2. Brushing on the original heatmap might not precisely capture rows or columns users expected. Users can manually remove a certain number of rows or columns from the four dimensions of the selected sub-heatmap. 3. Selected sub-heatmap can be further converted into a second interactive heatmap application. It will be introduced in Section 3.4.

• **Export table**: The values in sub-heatmap can be viewed and exported as a text table.

• **Save sub-heatmap**: This is similar as in the original heatmap.

• **Resize sub-heatmap**: This is similar as in the original heatmap.

At the bottom of the application, there is an output component. If a heatmap cell was clicked, the output component prints the meta information of that cell, such as the value, the row and column indices and labels. If there are annotations associated in heatmap, corresponding annotation values are also printed. If an area was selected from the original heatmap, the output component prints a runnable code that can be used to obtain row and column indices from the corresponding matrix. The output component can be self-defined to allow more complex output to respond to user’s actions on the heatmap. It will be introduced in Section 5.

### 3.3. Use last generated heatmaps

**ComplexHeatmap** is broadly used in many scripts and packages where they generate heatmaps but do not directly return the **Heatmap/HeatmapList** objects. This won’t affect the use of **InteractiveComplexHeatmap** because the last generated heatmap object is always automatically saved. Calling `htShiny()` without a heatmap object will automatically use the last one. We demonstrate this functionality with package **cola** (Gu, Schlesner, and Hübschmann 2020).

**cola** heavily uses **ComplexHeatmap** to implement various customized heatmaps to visualize consensus clustering results as well as results from downstream analysis. As an example, the function `get_signatures()` extracts signatures that are significantly different between the predicted subgroups. `get_signatures()` makes a list of heatmaps for visualizing the patterns of signatures and returns a data frame of the signatures as well as various statistics for the statistical test. When `get_signatures()` draws the heatmaps, the heatmap object is saved internally, then directly calling `htShiny()` without any argument converts the signature heatmap into an interactive application (Figure 3).

```r
R> library(cola) # from Bioconductor
R> data(golub_cola)
```
Make Interactive Complex Heatmaps in R

Figure 3: Interactivating last generated heatmap. The original heatmap is generated by `get_signatures()` from package `cola`.

```r
R> get_signatures(golub_cola["ATC:skmeans"], k = 3) # this makes the heatmap
R> ht_shiny()
```

Note the functionality of automatically saving the last heatmap object is only turned on when `InteractiveComplexHeatmap` is loaded, i.e., `library(InteractiveComplexHeatmap)` should be executed before making heatmap, or users simply execute `ComplexHeatmap::ht_opt(save_last = TRUE)` to manually turn it on.

3.4. Recursive interactive heatmap application

When the original heatmap visualizes a huge matrix, in the interactive heatmap application, a small selection rectangle would cover a dense subset of rows and columns where single cells are still hard to identify in the sub-heatmap component. In this case, the sub-heatmap can be continually exported to another independent interactive heatmap widget which is in a new layer above current one, just by clicking the button “Interactivate sub-heatmap” in the tool under sub-heatmap (Figure 4). This process can be recursively applied until users are satisfied with the details seen in the sub-heatmap.

3.5. Applications on other heatmap functions

`InteractiveComplexHeatmap` automatically interactivates static heatmaps generated by `ComplexHeatmap`. To facilitate analysis from users who still use other heatmap functions and also want to convert their heatmaps into interactive, in the companion package `ComplexHeatmap`, we implemented three “translation functions” `ComplexHeatmap:::heatmap()`, `ComplexHeatmap:::heatmap.2()` and `ComplexHeatmap:::pheatmap()` to replace `stats::heatmap()`, `gplots::heatmap.2()` and `pheatmap::pheatmap()`. The three translation functions have the
Figure 4: Recursively generate interactive heatmaps. The second interactive heatmap (Figure B) is generated from the sub-heatmap in the first interactive heatmap by clicking the button "Interactivate sub-heatmap" (highlighted in orange rectangle in Figure A).
same set of arguments as original ones and corresponding arguments are translated to proper settings in `ComplexHeatmap` so that they make heatmaps visually as same as possible to original heatmaps. Note in order not to dilute user’s namespace, `heatmap()` and `heatmap.2()` are not exported from `ComplexHeatmap` and users must add `ComplexHeatmap:::` prefix to use them. While for `pheatmap()`, if package `pheatmap` is already loaded ahead of `ComplexHeatmap`, there is a message printed to inform `ComplexHeatmap::pheatmap()` overwrites `pheatmap::pheatmap()`. Since the translation functions have the same sets of arguments as original ones, no further modification on code that users need to apply. `ComplexHeatmap:::heatmap()`, `ComplexHeatmap:::heatmap.2()` and `ComplexHeatmap:::pheatmap()` all generate `Heatmap` objects, thus heatmaps from these three functions can be converted into interactive. The returned object by e.g., `ComplexHeatmap:::heatmap()` can be sent to `htShiny()`, or `htShiny()` can be directly called with no argument after the heatmap is drawn. Example code is as follows:

```r
R> ht = ComplexHeatmap:::heatmap(...)  
R> htShiny(ht)  
R> # or even simpler  
R> ComplexHeatmap:::heatmap(...)  
R> htShiny()  
R> ComplexHeatmap:::heatmap.2(...)  
R> htShiny()  
R> ComplexHeatmap:::pheatmap(...)  
R> htShiny()  
```

Demonstrations are in Figure 5 and live examples on the three heatmap functions can be obtained from `htShinyExample(2.4)`, `htShinyExample(2.5)` and `htShinyExample(2.6)`.  

3.6. Applications on other high-level plots

`ComplexHeatmap` supports implementing high-level plots as long as the data is represented in a matrix-like structure. The following listed functions all generate `Heatmap` objects, thus, they can be easily exported to interactive applications.

- `ComplexHeatmap::oncoPrint()`: It visualizes various types of genomic alterations for a list of genes in multiple samples and generates a so-called oncoPrint plot. The example of an interactive oncoPrint can be found in Figure 6A and a live example can be run with `htShinyExample(2.2)`.  
- `ComplexHeatmap::UpSet()`: It implements the UpSet plot (Conway et al. 2017) which is an efficient way for visualizing intersections between a large number of sets. Additionally `UpSet()` is able to visualize intersections between multiple lists of genomic intervals. The example of an interactive UpSet plot can be found in Figure 6B and a live example can be run with `htShinyExample(2.3)`.  
- `ComplexHeatmap::densityHeatmap()`: It visualizes a list of distributions by heatmap. The example of an interactive density heatmap can be found in Figure 6C and a live example can be run with `htShinyExample(2.1)`.
Figure 5: Applications on heatmaps generated from other heatmap functions. The three interactive heatmaps are from A) `heatmap()`, B) `heatmap.2()` and C) `pheatmap()`.
• EnrichedHeatmap::EnrichedHeatmap(): The package EnrichedHeatmap visualizes the enrichment of certain genomic signals (e.g., DNA methylation) on specific genomic features (e.g., gene TSS) by a special heatmap. EnrichedHeatmap inherits ComplexHeatmap and the constructor function EnrichedHeatmap() returns a Heatmap object, thus, an “enriched heatmap” can be exported as an interactive application as well. One simple example is in Figure 6D and runnable examples can be found in htShinyExample(3.1), htShinyExample(3.2) and htShinyExample(3.3).

As we have demonstrated in Section 3.3, for some other high-level plots generated by functions which internally use ComplexHeatmap while do not directly return Heatmap or HeatmapList objects (e.g., cola::get_signatures() or cola::consensus_heatmap()), users can always first generate heatmap in a certain graphics device (on-screen or off-screen), then execute htShiny() with no heatmap object to obtain the interactive applications (see examples in htShinyExample(1.6) and htShinyExample(1.7)).

3.7. Interactivate heatmaps indirectly generated by heatmap(), heatmap.2() and pheatmap()

In Section 3.3, we introduced how to export heatmaps indirectly generated from ComplexHeatmap to interactive applications. There is another scenario where heatmaps are indirectly generated by stats::heatmap(), gplots::heatmap.2() or pheatmap::pheatmap(), i.e., they are generated by functions which internally use the three heatmap functions. How can we turn these heatmaps into interactive? The solution is simple. We just need to go to e.g. pheatmap namespace and replace pheatmap with ComplexHeatmap::pheatmap right there.

The following example is from package SC3 (Kiselev, Kirschner, Schaub, Andrews, Yiu, Chandra, Natarajan, Reik, Barahona, Green, and Hemberg 2017) where function sc3_plot_expression() draws a heatmap internally using pheatmap().

```
R> library(SingleCellExperiment)
R> library(SC3)
R> library(scater)
R>
R> sce = SingleCellExperiment(
+   assays = list(counts = as.matrix(yan),
+                  logcounts = log2(as.matrix(yan) + 1)),
+   colData = ann
+ )
R>
R> rowData(sce)$feature_symbol = rownames(sce)
R> sce = sce[!duplicated(rowData(sce)$feature_symbol),]
R> sce = runPCA(sce)
R> sce = sc3(sce, ks = 2:4, biology = TRUE)
R>
R> sc3_plot_expression(sce, k = 3)
```

To replace the internal use of pheatmap::pheatmap() with ComplexHeatmap::pheatmap(), the function assignInNamespace() can be used to directly change the value of pheatmap.
Figure 6: Applications on other high-level plots. A) An interactive oncoPrint. B) An interactive UpSet plot. C) An interactive density heatmap. D) An interactive enriched heatmap.
in `pheatmap` namespace. After doing that, recalling `sc3_plot_expression()` always uses `ComplexHeatmap::pheatmap()` and now it is possible to use `htShiny()` to export it to an interactive application.

```r
R> assignInNamespace("pheatmap", ComplexHeatmap::pheatmap, ns = "pheatmap")
R> library(InteractiveComplexHeatmap)
R> sc3_plot_expression(sce, k = 3)
R> htShiny()
```

To safely change `stats::heatmap()`, `gplots::heatmap.2()` or `pheatmap::pheatmap()` to `ComplexHeatmap:::heatmap()`, `ComplexHeatmap:::heatmap.2()` or `ComplexHeatmap::pheatmap()`, it is recommended to execute one of the following commands at the beginning of user’s R session.

```r
R> assignInNamespace("heatmap", ComplexHeatmap:::heatmap, ns = "stats")
R> assignInNamespace("heatmap.2", ComplexHeatmap:::heatmap.2, ns = "gplots")
R> assignInNamespace("pheatmap", ComplexHeatmap::pheatmap, ns = "pheatmap")
```

## 4. Implementation of interactivity

Being different from interactive heatmap packages based on JavaScript, e.g., `iheatmapr`, `heatmaply` or `d3heatmap`, `InteractiveComplexHeatmap` has a special way to capture positions that users selected on heatmaps and to extract values from corresponding matrices. To demonstrate it, we still use the object `ht_list` previously generated in Section 3.1 which includes a list of two heatmaps and k-means clustering was applied on the numeric heatmap.

`InteractiveComplexHeatmap` implements two types of interactivity: 1. on the interactive graphics device, and 2. in the Shiny web application. The interactivity on the interactive graphics device is the basis of the interactivity of the Shiny application, thus, in following sections, we will first introduce how the interactivity is implemented with the interactive graphics device.

### 4.1. On the interactive graphics device

Here the “interactive graphics device” refers to the window for generating plots if R is directly used in the terminal, or the figure panel in Rstudio IDE.

`InteractiveComplexHeatmap` first captures physical positions of all heatmap slices, i.e. the distance to the bottom left of the graphics device, by the function `htPositionsOnDevice()`. Internally, the function goes to the viewport of every heatmap slice and utilizes the function `grid::deviceLoc()` to capture positions measured in the graphics device. Before executing `htPositionsOnDevice()`, the heatmap should be already drawn on the device so that `htPositionsOnDevice()` can access various viewports.

```r
R> pos = htPositionsOnDevice(ht_list)
R> pos
```
DataFrame with 6 rows and 8 columns

<table>
<thead>
<tr>
<th>heatmap</th>
<th>slice</th>
<th>row_slice</th>
<th>column_slice</th>
<th>x_min</th>
</tr>
</thead>
<tbody>
<tr>
<td>mat_a</td>
<td>mat_a</td>
<td>mat_a_heatmap_body_1_1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>mat_a</td>
<td>mat_a</td>
<td>mat_a_heatmap_body_1_2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>mat_a</td>
<td>mat_a</td>
<td>mat_a_heatmap_body_2_1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>mat_a</td>
<td>mat_a</td>
<td>mat_a_heatmap_body_2_2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>mat_b</td>
<td>mat_b</td>
<td>mat_b_heatmap_body_1_1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>mat_b</td>
<td>mat_b</td>
<td>mat_b_heatmap_body_2_1</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>x_max</th>
<th>y_min</th>
<th>y_max</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.66inches</td>
<td>0.948inches</td>
<td>2.83inches</td>
</tr>
<tr>
<td>2.63inches</td>
<td>0.948inches</td>
<td>2.83inches</td>
</tr>
<tr>
<td>1.66inches</td>
<td>0.437inches</td>
<td>0.909inches</td>
</tr>
<tr>
<td>2.63inches</td>
<td>0.437inches</td>
<td>0.909inches</td>
</tr>
<tr>
<td>5.08inches</td>
<td>0.948inches</td>
<td>2.83inches</td>
</tr>
<tr>
<td>5.08inches</td>
<td>0.437inches</td>
<td>0.909inches</td>
</tr>
</tbody>
</table>

Note, the calculation of heatmap slice positions relies on the device size. In previous example code, pos was calculated in a device with 6 inches width and 4 inches height.

The returned object pos is a DataFrame object that contains the positions of all heatmap slices. The DataFrame class defined in package S4Vectors (Pagès, Lawrence, and Aboyoun 2020) is very similar to a data frame, but it can store more complex data types, such as the simpleUnit vectors generated by grid::unit(). Figure 7A contains the original heatmaps and black rectangles in Figure 7B were drawn based on the positions that were captured.

Next, InteractiveComplexHeatmap captures the physical position of the point that the user clicked on the heatmap. This is simply done by function grid::grid.locator() that is applied in the ROOT viewport. With knowing positions of heatmap slices and the point that the user clicked, it is possible to calculate which row and column in the original matrix user’s click corresponds to.

In Figure 7B, blue point with coordinate \((a, b)\) was clicked by the user. The heatmap slice where the user clicked into can be easily identified by comparing the position of the click and the positions of every heatmap slice. Assume the heatmap slice where user’s click is in has a range \((x_1, x_2)\) on \(x\) direction and a range \((y_1, y_2)\) on \(y\) direction. There are \(n_r\) rows \((n_r = 8)\) and \(n_c\) columns \((n_c = 5)\) in this heatmap slice and they are marked by dashed lines in Figure 7B. Note all coordinate values \((a, b, x_1, y_1, x_2, y_2)\) are measured as the distances to the bottom left of the graphics device.

In this heatmap slice, the row index \(i_r\) and column index \(i_c\) of the cell where the click is in can be calculated as (assume the left bottom corresponds to the index of 1 for both rows and columns):

\[
i_c = \left\lfloor \frac{a - x_1}{x_2 - x_1} \cdot n_c \right\rfloor
\]

\[
i_r = \left\lfloor \frac{b - y_1}{y_2 - y_1} \cdot n_r \right\rfloor
\]
Figure 7: Demonstration of the implementation of interactivity. A) The original heatmap. B) The black borders are drawn based on the positions captured for all heatmap slices. Figure A and B were both drawn in a device with 6 inches width and 4 inches height. Dashed rectangles correspond to image borders.
where the symbol $\lceil x \rceil$ means the ceiling of the numeric value $x$. In ComplexHeatmap, the row with index 1 is always put on the top of the heatmap, then $i_r$ should be adjusted as:

$$i_r = n_r - \left\lceil \frac{b - y_1}{y_2 - y_1} \cdot n_r \right\rceil + 1$$

The subset of row and column indices of the original matrix that belongs to the selected heatmap slice is already stored in `ht_list` object (they can be retrieved by `row_order()` and `column_order()` function), thus, we can obtain the row and column index in the original matrix that corresponds to user’s click easily with $i_r$ and $j_c$.

Denote the matrix for the complete heatmap (without splitting) as $M$, and denote the subset of row and column indices in the heatmap slice where user’s click is in as $o_r, o_c$. Note, $o_r$ and $o_c$ can be reordered due to e.g. clustering. Then row and column indices ($j_r$ and $j_c$) for the clicked point in $M$ are calculated as follows:

$$j_r = o_r \cdot i_r$$

$$j_c = o_c \cdot j_c$$

And the corresponding value in $M$ is $M_{j_r,j_c}$.

InteractiveComplexHeatmap has two functions `selectPosition()` and `selectArea()` which allow users to pick single positions or to select areas from heatmaps. Under the interactive graphics device, users do not need to run `htPositionsOnDevice()` explicitly. The positions of heatmaps are automatically calculated, cached and reused if heatmaps are the same and the device has not changed its size. If users change the device size, `htPositionsOnDevice()` will be automatically re-executed.
Figure 8 demonstrates an example of using `selectPosition()` function. Interactively, the function asks the user to click one position on the heatmap. The function returns a DataFrame which contains the heatmap name, slice name and the row and column index of the corresponding matrix. An example output from `selectPosition()` is as follows:

```r
## DataFrame with 1 row and 6 columns
## heatmap slice row_slice column_slice row_index
## <character> <character> <numeric> <numeric> <integer>
## 1 mat_a mat_a_heatmap_body_1_2 1 2 4
## column_index
## <integer>
## 1 2
```

The output means the position that the user clicked is in a heatmap named “mat_a” and in its first row slice and in the second column slice. Assume `mat` is the matrix for heatmap “mat_a”, then the clicked point corresponds to the value `mat[4, 2]`.

Similarly, the function `selectArea()` asks the user to click two positions on the heatmap which define an area. Note since the selected area may overlap over multiple heatmaps and slices, the function returns a DataFrame with multiple rows. An example output is as follows.

```r
## DataFrame with 4 rows and 6 columns
## heatmap slice row_slice column_slice row_index
## <character> <character> <numeric> <numeric> <IntegerList>
## 1 mat_a mat_a_heatmap_body_1_2 1 2 7,5,2,...
## 2 mat_a mat_a_heatmap_body_2_2 2 2 6,3
## 3 mat_b mat_b_heatmap_body_1_1 1 1 7,5,2,...
## 4 mat_b mat_b_heatmap_body_2_1 2 1 6,3
## column_index
## <IntegerList>
## 1 2,4,1,...
## 2 2,4,1,...
## 3 1,2,3,...
## 4 1,2,3,...
```

The columns `row_index` and `column_index` are stored in a class of `IntegerList` which is simply a list of integer vectors. To get the row indices in e.g. `mat_a_heatmap_body_1_2` (in the first row), user can use either one of the following command (assume the DataFrame object is named `df`):

```r
R> df[1, "row_index"][[1]]
R> unlist(df[1, "row_index"])
R> df$row_index[[1]]
```

### 4.2. On off-screen graphics devices
It is also possible to use `selectPosition()` and `selectArea()` on off-screen graphics devices, such as `pdf` or `png`. Now the positions cannot be selected interactively, but instead argument `pos` in `selectPosition()` or `pos1/pos2` in `selectArea()` need to be specified to simulate clicks. The values for `pos`, `pos1` and `pos2` all should be `unit` objects of length two which corresponds to the $x$ and $y$ coordinates of the positions relative to the bottom left of the device.

```r
R> png(...)  
R> ht_list = draw(ht_list)  
R> pos = selectPosition(ht_list, pos = unit(c(3, 3), "cm"))  
R> dev.off()
```

Users do not need to use this functionality directly with an off-screen graphics device, but it is very useful when implementing the interactivity in a Shiny application where the plot is actually generated under an off-screen graphics device.

### 4.3. Shiny web application

With the three functions `htPositionsOnDevice()`, `selectPosition()` and `selectArea()`, it is possible to implement Shiny applications for interactively working with heatmaps. Now the problem is how does the server side capture the positions that the user clicked on the web page. Luckily, there is a solution for this. The output heatmap is normally put within a `shiny::plotOutput()` and `plotOutput()` provides two actions: `click` and `brush`, so that the server can receive the information of the positions as soon as the user performs clicking or brushing on heatmaps. With knowing the positions, they can be set to `selectPosition()` and `selectArea()` via `pos` or `pos1/pos2` arguments for correctly corresponding to the original matrices.

### 5. Shiny web application development

`htShiny()` exports heatmaps as stand-alone Shiny web applications. `InteractiveComplexHeatmap` also provides general solutions for integrating interactive heatmap widgets in other Shiny application development. It allows dynamically generating interactive heatmap widgets according to different configurations on heatmaps, and customizing the output that responds to user’s actions on heatmaps.

#### 5.1. Usage

There are following two main functions for Shiny application development:

- `InteractiveComplexHeatmapOutput()`: for building user interface (UI).
- `makeInteractiveComplexHeatmap()`: for processing on the server side.

As already demonstrated in Section 3.2, the interactive heatmap widget contains three components, i.e., the original heatmap, the sub-heatmap and the output that prints information of cells that users selected. The first two heatmap components contain various tools for controlling both heatmaps. `makeInteractiveComplexHeatmap()` defines a list of responses to user’s actions that were performed on heatmaps and in the tools.
My first interactive ComplexHeatmap Shiny app

This is an interactive heatmap visualization on a random matrix.

**Output**

Selected over 1 heatmap with 27 rows and 21 columns. Row and column indices can be obtained by copying following code:

`show/hide code`

---

Figure 9: A simple Shiny application directly from `ui` and `server`.
The usage for implementing a Shiny application with these two functions is rather simple. Following is an example that can be directly copied and pasted to an R session (Figure 9).

```r
R> library(ComplexHeatmap)
R> library(InteractiveComplexHeatmap)
R> library(shiny)
R>
R> data(rand_mat) # simply a random matrix
R> ht1 = Heatmap(rand_mat, name = "mat",
+ show_row_names = FALSE, show_column_names = FALSE)
R> ht1 = draw(ht1)
R>
R> ui = fluidPage(
+ h3("My first interactive ComplexHeatmap Shiny app"),
+ p("This is an interactive heatmap visualization on a random matrix."),
+ InteractiveComplexHeatmapOutput()
+ )
R> server = function(input, output, session) {
+ makeInteractiveComplexHeatmap(input, output, session, ht1)
+ } 
R> shinyApp(ui, server)
```

Multiple interactive heatmap widgets can be integrated into one single Shiny application. Now a “heatmap ID” must be assigned to each widget, so that `makeInteractiveComplexHeatmap()` can find the correct heatmap to respond. If there is only one heatmap widget in the Shiny application, such as in the previous example, the heatmap ID is internally generated if it is not specified, and the UI is automatically linked to the server in the two functions. In the following example, we created a second widget which visualizes a random character matrix (Figure 10). The two interactive heatmap widgets are independent in the application. See `htShinyExample(1.7)` and `htShinyExample(5.1)` for live examples.

```r
R> set.seed(88)
R> mat2 = matrix(sample(letters[1:10], 100, replace = TRUE), 10)
R> ht2 = draw(Heatmap(mat2, name = "mat2"))
R>
R> ui = fluidPage(
+ h3("The first heatmap"),
+ InteractiveComplexHeatmapOutput("heatmap_1",
+ height1 = 300, height2 = 300),
+ hr(),
+ h3("The second heatmap"),
+ InteractiveComplexHeatmapOutput("heatmap_2",
+ height1 = 300, height2 = 300)
+ )
R> server = function(input, output, session) {
+ makeInteractiveComplexHeatmap(input, output, session, ht1, "heatmap_1")
+ makeInteractiveComplexHeatmap(input, output, session, ht2, "heatmap_2")
}```
Make Interactive Complex Heatmaps in R

The first heatmap

Output
Selected over 1 heatmap with 24 rows and 21 columns. Row and column indices can be obtained by copying following code:

show/hide code

The second heatmap

Output
Selected over 1 heatmap with 4 rows and 4 columns. Row and column indices can be obtained by copying following code:

show/hide code

Figure 10: A Shiny application with two interactive heatmap widgets.
5.2. Customize the widgets

The original heatmap and sub-heatmap components can be resized by dragging from bottom right of the two boxes, but still, `InteractiveComplexHeatmapOutput()` provides arguments of `width1`, `width2`, `height1` and `height2` to precisely control the initial sizes of the two components. They can be manually set to make sure the heatmaps are well visualized and aligned when the application is initialized.

The layout of the three components are controlled by argument `layout`. It supports following values:

- "(1-2)|3": Original heatmap and sub-heatmap are in the same row, and output is in a second row. This is the default layout.
- "1|(2-3)": Original heatmap is in a single row, while sub-heatmap and output are in a second row.
- "1-2-3": All three components are in the same row.
- "1|2|3": Each component is in a single row.
- "1-(2|3)": Being different from the other four layouts, this is a two-column layout. Original heatmap is in a single column. Sub-heatmap and output are vertically aligned and the two are in the second column. An example can be found at htShinyExample(4.1).

Note the values for `layout` are in a special format to help to understand the layout, where the three code 1, 2 and 3 correspond to original heatmap, sub-heatmap and output respectively, symbol "-" corresponds to horizontal alignment and "|" corresponds to vertical alignment. With different layouts, different default values are assigned to widths and heights of the three components to make sure they are well aligned.

By default, a "click" action is used to retrieve the information of a single cell in a heatmap.
In `InteractiveComplexHeatmapOutput()`, argument `action` accepts other values of "hover" or "dblclick" so that hovering or double clicking is used instead.

The argument `response` can be set as one of "click", "hover", "dblclick" or "brush" so that the server only responds to one action on the heatmap. For example, if `response` is set to "click", there will be no response for the “brush action” in the interactive heatmap, also the sub-heatmap component is removed from the application. Runnable examples are in htShinyExample(1.9) and htShinyExample(9.3).

5.3. Work with R Markdown documents

It is straightforward to integrate `InteractiveComplexHeatmap` in an interactive R Markdown document, just in the same way as integrating normal Shiny applications. Examples can be found in htShinyExample(7.1) and htShinyExample(7.2).

5.4. Self-define the output
Both click and brush actions trigger an output associated to the heatmap. The output gives the information of rows and columns selected by users. The response for the two actions can be self-defined.

In `makeInteractiveComplexHeatmap()`, there are two arguments: `click_action` and `brush_action` which accept self-defined functions to define how to respond after the heatmap was clicked or brushed. The two functions should accept two arguments, the first is a DataFrame object which contains the information of which rows and columns are selected by the user, and the second argument should always be `output` which is used in the Shiny server function.

To use `click_action` or `brush_action`, a `htmlOutput` (or other similar *Output) should be first set up in the UI, then the Shiny application knows where to update the output. The output UI can replace the default output by directly assigning to argument `output.ui` in `InteractiveComplexHeatmapOutput()`.

```r
R> ui = fluidPage(
+   InteractiveComplexHeatmapOutput(output.ui = htmlOutput("info"))
+ )

Or to create a new output UI independent to the interactive heatmap widget.

```r
R> ui = fluidPage(
+   InteractiveComplexHeatmapOutput(),
+   htmlOutput("info")
+ )
```r

The `click_action` or `brush_action` is normally defined as follows (assume the ID set in `htmlOutput()` is "info"):

```r
R> function(df, output) {
+   outputiteralOutput{{
+     if(is.null(df)) {
+       ...
+     } else {
+       ...
+     }
+   }}
+ }
```r

If users didn’t click or brush inside the heatmap body (e.g., clicked in the dendrogram), `df` that is passed to the functions will be NULL. A sanity check might be performed here to print specific output when heatmap was not selected.

The format of `df` is slightly different between click and brush. If it is a click action, `df` has the same format as the returned object of function `selectPosition()` (introduced in Section 4), which looks like the following chunk. It always has one row if the user clicked into the heatmap.
### DataFrame with 1 row and 6 columns

<table>
<thead>
<tr>
<th>heatmap</th>
<th>slice</th>
<th>row_slice</th>
<th>column_slice</th>
<th>row_index</th>
<th>column_index</th>
</tr>
</thead>
<tbody>
<tr>
<td>mat_a</td>
<td>mat_a_heatmap_body_1_2</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

If it is a brush action, `df` has the same format as the returned object of function `selectArea()` (introduced in Section 4), which looks like the following chunk. Each line contains row and column indices of the selected sub-matrix in a specific heatmap slice of a specific heatmap.

### DataFrame with 4 rows and 6 columns

<table>
<thead>
<tr>
<th>heatmap</th>
<th>slice</th>
<th>row_slice</th>
<th>column_slice</th>
<th>row_index</th>
<th>column_index</th>
</tr>
</thead>
<tbody>
<tr>
<td>mat_a</td>
<td>mat_a_heatmap_body_1_2</td>
<td>1</td>
<td>2</td>
<td>7,5,2,...</td>
<td></td>
</tr>
<tr>
<td>mat_a</td>
<td>mat_a_heatmap_body_2_2</td>
<td>2</td>
<td>2</td>
<td>6,3</td>
<td></td>
</tr>
<tr>
<td>mat_b</td>
<td>mat_b_heatmap_body_1_1</td>
<td>1</td>
<td>1</td>
<td>7,5,2,...</td>
<td></td>
</tr>
<tr>
<td>mat_b</td>
<td>mat_b_heatmap_body_2_1</td>
<td>2</td>
<td>1</td>
<td>6,3</td>
<td></td>
</tr>
</tbody>
</table>

Note as demonstrated above, after a brush action, values in column `row_index` and `column_index` might be duplicated due to the fact that the selected heatmap slices are in the same row or column. E.g., in the previous example, the first and the third rows correspond to the selection in the first row slice, but in two different column slices, thus they have the same values for `row_index`. To safely get row indices and column indices of the selected heatmap, users might need to perform:

```r
R> unique(unlist(df$row_index))
R> unique(unlist(df$column_index))
```

If argument `action` in `InteractiveComplexHeatmapOutput()` was set to "hover" or "dblclick", the corresponding argument in `makeInteractiveComplexHeatmap()` is `hover_action` or `dblclick_action` respectively. Their usages are the same as `click_action`.

We demonstrate how to self-define `click_action` in the next example which visualizes a correlation matrix based on `mtcars` dataset. In the UI, the default output is replaced with a new `plotOutput`. On the server side, a self-defined `click_action` is defined to draw a scatter plot of the two corresponding variables that were selected from heatmap (Figure 11). Note in the example `response = "click"` is set to simplify Figure 11 so that the sub-heatmap component is not included in the application. The runnable example can be found with `htShinyExample(5.6)`.
Figure 11: A demonstration of self-defining the output. The heatmap visualizes a correlation matrix based on `mtcars` dataset. Clicking on a cell draws a scatter plot of the two corresponding variables.

```r
R> data(mtcars)
R> cor_mat = cor(mtcars)
R>
R> library(circlize)
R> col_fun = colorRamp2(c(-1, 0, 1), c("darkgreen", "white", "red"))
R> ht = Heatmap(cor_mat, name = "Correlation",
R> +   col = col_fun, show_row_dend = FALSE, show_column_dend = FALSE)
R>
R> ui = fluidPage(
R>   +   InteractiveComplexHeatmapOutput(
R>   +     output_ui = plotOutput("scatterplot", width = 400, height = 400),
R>   +     response = "click"
R>   +   )
R>   
R>   +   click_action = function(df, output) {
R>   +     output$scatterplot = renderPlot({
R>   +       nm = colnames(mtcars)
R>   +       i1 = df$column_index
R>   +       i2 = df$row_index
R>   +       x = mtcars[, i1]
R>   +       y = mtcars[, i2]
R>   +       plot(x, y, xlab = nm[i1], ylab = nm[i2],
R>   +         main = paste0("Correlation = ", sprintf("%.3f", cor(x, y)))
R>   +     })
R>   +   })
```
Figure 12: Floating output to mouse positions. The heatmap visualizes a GO similarity matrix from an analysis by package `simplifyEnrichment`. The output for click was self-defined to show the description of the two GO IDs and argument `output_ui_float` in `InteractiveComplexHeatmapOutput()` was set to `TRUE` to float the output to mouse positions. Top left of the output box corresponds to the cell that was clicked.

```r
R> server = function(input, output, session) {
+   makeInteractiveComplexHeatmap(input, output, session, ht,
+       click_action = click_action)
+ }
R> shinyApp(ui, server)
```

There are several other examples in `InteractiveComplexHeatmap` that demonstrate the use of self-defining output:

- **htShinyExample(5.3)**: A gene expression matrix is visualized and clicking on the heatmap prints the corresponding gene ID and other annotations related to this gene.

- **htShinyExample(5.4)**: The heatmap visualizes correlations of a list of Gene Ontology (GO) terms. In this example, the click and brush actions are self-defined so that the selected GO IDs as well as their detailed descriptions are printed.

- Section 5.6 demonstrates a complex Shiny example of visualizing results from a differential expression analysis where the output is self-defined to link selected genes on heatmaps to a MA-plot as well as a table of results only for these selected genes.

Instead of occupying static space in the application, the output component can be floated to the mouse positions by setting `output_ui_float = TRUE` in `InteractiveComplexHeatmapOutput()`, then clicking, hovering or brushing from the heatmap opens a frame that contains the output. **htShinyExample(9.1)** demonstrates output floating when mouse action is set to...
"hover", "click" or "dblclick", respectively. Figure 12 demonstrates a slightly complex example where clicking on the GO similarity heatmap opens a floating frame that contains the information of the two corresponding GO terms.

5.5. Dynamically generate interactive heatmap widget

In previous examples, heatmaps are already generated before making the interactive applications. There are also scenarios where the heatmaps are generated on the fly. There might be following scenarios where heatmap is dynamically generated:

- The heatmap is based on a subset of matrix which is filtered by users, *e.g.*, the expression matrix for differentially expressed genes filtered by different cutoffs.
- The annotations are dynamically provided by users.
- The heatmap parameters are changed by users, *e.g.*, the clustering method or the splitting variable.
- If there are multiple heatmaps, which heatmaps are going to be drawn is dynamically selected.

Since now the heatmap is dynamically generated, thus code that generates the heatmap according to user's configurations as well as `makeInteractiveComplexHeatmap()` can be wrapped in `shiny::observe()` or `shiny::observeEvent()`, so that changes on heatmap trigger the update of the interactive heatmap. In next example, the interactive heatmap widget is dynamically generated according to the number of heatmaps that the user selected.

```r
R> ui = fluidPage(
+   sliderInput("n_heatmap", label = "How many heatmaps?",
+     value = 1, min = 1, max = 5),
+   InteractiveComplexHeatmapOutput()
+ )

R> generate_heatmap_list = function(n) {
+   ht_list = NULL
+   for(i in 1:n) {
+     ht_list = ht_list + Heatmap(matrix(rnorm(100), 10),
+       name = paste0("mat_", i),
+       column_title = paste0("heatmap_", i))
+   }
+   ht_list
+ }

R> server = function(input, output, session) {
+   observe({
+     ht_list = generate_heatmap_list(input$n_heatmap)
+     makeInteractiveComplexHeatmap(input, output, session, ht_list)
+   })
+ }
R> shiny::shinyApp(ui, server)
```

The previous example can also be implemented by the function `InteractiveComplexHeatmapWidget()`. To use this function, a placeholder by `htmlOutput()` should be first created on
Figure 13: Dynamically generate the interactive heatmap widget. The widget visualizes a user-specified number of heatmaps.
the UI. On the server side, `InteractiveComplexHeatmapWidget()` dynamically creates the complete interactive heatmap widget and inserts it to the place defined by `htmlOutput()`. One advantage of using `InteractiveComplexHeatmapWidget()` is it supports removing the interactive heatmap widget from the application, which would be useful if the interactive widget is only for temporary use.

```r
R> ui = fluidPage(
+   sliderInput("n_heatmap", label = "How many heatmaps?",
+               value = 1, min = 1, max = 5),
+   htmlOutput("heatmap_widget")
+ )

R> server = function(input, output, session) {
+   observe({
+     ht_list = generate_heatmap_list(input$n_heatmap)
+     InteractiveComplexHeatmapWidget(input, output, session, ht_list,
+       output_id = "heatmap_widget")
+   })
+ }

R> shiny::shinyApp(ui, server)
```

In previous examples, the interactive heatmap widget is immediately generated when the Shiny application is loaded. However, there are scenarios where the widget does not need to be firstly initialized, while it is triggered by users’ action. The function `InteractiveComplexHeatmapModal()` dynamically generates an interactive heatmap widget which locates in a new layer above the current web page. Following code demonstrates an application which dynamically creates interactive widgets of a numeric heatmap or a character heatmap. Live examples can be found in `htShinyExample(6.3)`, `htShinyExample(6.4)` and `htShinyExample(6.5)`. The recursive interactive heatmap introduced in Section 3.4 is also implemented with `InteractiveComplexHeatmapModal()`.

```r
R> ui = fluidPage(
+   radioButtons("select", "Select a matrix",
+               choices = c("Numeric" = 1, "Character" = 2)),
+   actionButton("show_heatmap", "Generate heatmap"),
+ )

R> get_heatmap = function(i) {
+   mat_list = list(
+     matrix(rnorm(100), 10),
+     matrix(sample(letters[1:10], 100, replace = TRUE), 10)
+   )
+   Heatmap(mat_list[[i]])
+ }

R> server = function(input, output, session) {
+   observeEvent(input$show_heatmap, {
+     i = as.numeric(input$select)
+     ht = get_heatmap(i)
+     InteractiveComplexHeatmapModal(input, output, session, ht)
+   })
+ }
```
5.6. Case study

Figure 14 demonstrates an interactive heatmap application which visualizes results of differential gene expression analysis on the airway dataset performed by package DESeq2 (Love, Huber, and Anders 2014). In the application, heatmap visualizes the expression of genes that are significantly different from a two-condition comparison, i.e., trt vs untrt (treated vs. untreated). In the original heatmap list, there are two additional one-column heatmaps which visualize the absolute expression level of genes (baseMean) and the fold change between the two conditions (log2foldChange). To get more comprehensive information of how genes are differentially expressed when selecting from heatmap, we defined extra output for the brush event on the original heatmaps. As shown in Figure 14, when a subset of genes are selected from the original heatmap, they are highlighted in a MA-plot which is a scatterplot of log2foldChange against baseMean where baseMean is in log10 scale. Also a table of the statistics for the selected genes from DESeq2 analysis is printed below MA-plot. Users can also search in heatmaps to obtain a subset of genes to generate corresponding MA-plot and table. The live example can be found in htShinyExample(5.5).

6. Conclusion

Interactivity on heatmaps allows users to capture features which have specific patterns on heatmaps. In this paper, we described a new R package InteractiveComplexHeatmap that supports interactive visualization on complex heatmaps. It can easily export static heatmaps into interactive Shiny applications and it also provides functions for implementing more complex Shiny applications. We believe it will be a useful tool for effectively interpreting data and developing new tools.

7. Acknowledgments

This work was supported by the German Cancer Research Center-Heidelberg Center for Personalized Oncology (DKFZ-HIPO) and by the Molecular Diagnostics Program of the National Center for Tumor Diseases (NCT) Heidelberg.

References


Figure 14: An interactive heatmap for visualizing results from a differential gene expression analysis.


Affiliation:

Zuguang Gu
German Cancer Research Center
Computational Oncology, Molecular Diagnostics Program, National Center for Tumor Diseases (NCT)

and

DKFZ-HIPO (Heidelberg Center for Personalized Oncology)
Im Neuenheimer Feld 280
69120 Heidelberg, Germany
E-mail: z.gu@dkfz-heidelberg.de
URL: http://jokergoo.github.io

Daniel Hübbschmann
German Cancer Research Center
Computational Oncology, Molecular Diagnostics Program, National Center for Tumor Diseases (NCT)

and

Heidelberg Institute of Stem cell Technology and Experimental Medicine (HI-STEM)

and

German Cancer Consortium (DKTK)
Im Neuenheimer Feld 280
69120 Heidelberg, Germany

and

Department of Pediatric Immunology, Hematology and Oncology, University Hospital Heidelberg
69120 Heidelberg, Germany
E-mail: d.huebschmann@dkfz-heidelberg.de