

# Package ‘BioVizSeq’

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**Type** Package

**Title** Visualizing the Elements Within Bio-Sequences

**Version** 0.1.0

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**Description** Visualizing the types and distribution of elements within bio-sequences. At the same time, We have developed a geom layer, geom\_rect(), that can generate rounded rectangles. No external references are used in the development of this package.

**License** Artistic-2.0

**Encoding** UTF-8

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<i>cdd_plot</i>	<i>cdd_plot</i>
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## Description

Visualization of domain in CDD file

## Usage

```
cdd_plot(
  cdd_file,
  fasta_file,
  the_order = NULL,
  domain_select = NULL,
  shape = "RoundRect",
  r = 0.3,
  legend_size = 15,
  domain_color = NULL
)
```

## Arguments

<code>cdd_file</code>	The path of cdd file.
<code>fasta_file</code>	The path of fasta file.
<code>the_order</code>	The path of order file. A List of Gene ID , One ID Per Line.
<code>domain_select</code>	The domain ID which you want to align with.
<code>shape</code>	RoundRect or Rect.
<code>r</code>	The radius of rounded corners.
<code>legend_size</code>	The size of legend.
<code>domain_color</code>	The color set of domain.

## Value

`p`

**Author(s)**

Shiqi Zhao

**Examples**

```
hitdata_path <- system.file("extdata", "hitdata.txt", package = "BioVizSeq")
fa_path <- system.file("extdata", "idpep.fa", package = "BioVizSeq")
cdd_plot(hitdata_path, fa_path)

order_path <- system.file("extdata", "order.csv", package = "BioVizSeq")
cdd_plot(hitdata_path, fa_path, the_order = order_path)
```

---

*cdd\_to\_loc*

*cdd\_to\_loc*

---

**Description**

Extract the location information of domain from cdd file

**Usage**

```
cdd_to_loc(cdd_file)
```

**Arguments**

*cdd\_file* CDD file.

**Value**

*data.frame*

**Author(s)**

Shiqi Zhao

**Examples**

```
hitdata_path <- system.file("extdata", "hitdata.txt", package = "BioVizSeq")
cdd_file <- readLines(hitdata_path)
domain_loc <- cdd_to_loc(cdd_file)
```

fastaleng	<i>fastaleng</i>
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### Description

Statistical sequence length

### Usage

```
fastaleng(fasta_file)
```

### Arguments

fasta_file	The path of protein fasta file.
------------	---------------------------------

### Value

data.frame

### Author(s)

Shiqi Zhao

### Examples

```
fasta_path <- system.file("extdata", "idpep.fa", package = "BioVizSeq")
fastaleng(fasta_path)
```

geom_rrect	<i>geom_rrect</i>
------------	-------------------

### Description

Rounded rectangle

### Usage

```
geom_rrect(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  r = 0.2,
  ...,
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE
)
```

## Arguments

mapping	Set of aesthetic mappings created by <code>aes</code> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
data	A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame.
stat	Name of stat to modify data.
position	The position adjustment to use for overlapping points on this layer.
r	The radius of rounded corners.
...	additional parameter, e.g. color, linewidth, alpha.
na.rm	If "FALSE" (default), missing values are removed with a warning. If "TRUE", missing values are silently removed, logical.
show.legend	Whether to show legend, logical.
inherit.aes	Whether to inherit aesthetic mappings, logical, defaults to "TRUE".

## Details

draws rounded rectangle by using the locations of the four corners (`xmin`, `xmax`, `ymin` and `ymax`) like `geom_rect()`.

## Value

`ggplot` object

## Aesthetics

`geom_rrect()` understands the following aesthetics (required aesthetics are in bold):

- `xmin`
- `xmax`
- `ymin`
- `ymax`
- `alpha`
- `colour`
- `fill`
- `group`
- `linetype`

Learn more about setting these aesthetics in `vignette("ggplot2-specs")`.

## Author(s)

Shiqi Zhao

## Examples

```
library(ggplot2)
df <- data.frame(
  xmin = c(1, 2, 3),
  xmax = c(2, 3, 4),
  ymin = c(1, 2, 3),
  ymax = c(2, 3, 4),
  category = c("A", "B", "C")
)

p <- ggplot(df) +
  geom_rect(aes(xmin = xmin, xmax = xmax,
                 ymin = ymin, ymax = ymax, fill = category),
            r = 0.4, linewidth = 1, colour = "black")

print(p)
```

*get\_motif\_location*      *get\_motif\_location*

## Description

Extract the location information of motif from mast or meme file

## Usage

```
get_motif_location(motif_file)
```

## Arguments

**motif\_file**      The motif data of mast or meme file.

## Value

list

## Author(s)

Shiqi Zhao

## Examples

```
meme_path <- system.file("extdata", "meme.xml", package = "BioVizSeq")
meme_file <- readLines(meme_path)
motif_loc <- get_motif_location(meme_file)

mast_path <- system.file("extdata", "mast.xml", package = "BioVizSeq")
mast_file <- readLines(mast_path)
```

```
motif_loc <- get_motif_location(mast_file)
```

---

*gff\_plot**gff\_plot*

---

## Description

Visualization of element in gff or gtf file

## Usage

```
gff_plot(  
  gff_file,  
  the_order = NULL,  
  shape = "Rect",  
  r = 0.3,  
  legend_size = 15,  
  element_color = NULL  
)
```

## Arguments

<code>gff_file</code>	The path of gff file.
<code>the_order</code>	The path of order of mRNA. It is also the mRNA you want to showcase. A List of Gene ID , One ID Per Line.
<code>shape</code>	RoundRect or Rect.
<code>r</code>	The radius of rounded corners.
<code>legend_size</code>	The size of legend.
<code>element_color</code>	The color set of element.

## Value

`p`

## Author(s)

Shiqi Zhao

## Examples

```
gff_path <- system.file("extdata", "test.gff", package = "BioVizSeq")  
gff_plot(gff_path)
```

`gff_to_loc`                  *gff\_to\_loc*

### Description

Extract the location information of element from gff or gtf file

### Usage

```
gff_to_loc(gff_data, mRNA_ID = NULL)
```

### Arguments

`gff_data`                  gff file.

`mRNA_ID`                  The mRNA you selected. If NULL, it means selecting all mRNAs.

### Value

list

### Author(s)

Shiqi Zhao

### Examples

```
gff_path <- system.file("extdata", "test.gff", package = "BioVizSeq")
gff_data <- read.table(gff_path, header = FALSE, sep = '\t')
gff_loc <- gff_to_loc(gff_data)

ID_path <- system.file("extdata", "ID_select.csv", package = "BioVizSeq")
mRNA_ID <- readLines(ID_path)
gff_loc <- gff_to_loc(gff_data, mRNA_ID=mRNA_ID)
```

`meme_plot`                  *meme\_plot*

### Description

Visualization of motif in meme file or mast file

**Usage**

```
meme_plot(  
  meme_file,  
  the_order = NULL,  
  motif_select = NULL,  
  shape = "RoundRect",  
  show_motif_id = FALSE,  
  r = 0.3,  
  legend_size = 15,  
  motif_color = NULL  
)
```

**Arguments**

<code>meme_file</code>	The path of meme file or mast file.
<code>the_order</code>	The path of order file. A List of Gene ID , One ID Per Line.
<code>motif_select</code>	The motif ID which you want to align with.
<code>shape</code>	RoundRect or Rect.
<code>show_motif_id</code>	Display the name of the motif.
<code>r</code>	The radius of rounded corners.
<code>legend_size</code>	The size of legend.
<code>motif_color</code>	The color set of motif.

**Value**

`p`

**Author(s)**

Shiqi Zhao

**Examples**

```
meme_path <- system.file("extdata", "meme.xml", package = "BioVizSeq")  
meme_plot(meme_path)  
  
mast_path <- system.file("extdata", "mast.xml", package = "BioVizSeq")  
meme_plot(mast_path)  
  
meme_plot(meme_path, motif_select="1", show_motif_id = TRUE)  
  
order_path <- system.file("extdata", "order.csv", package = "BioVizSeq")  
meme_plot(meme_path, the_order=order_path, motif_select="1")
```

<code>motif_plot</code>	<i>motif_plot</i>
-------------------------	-------------------

## Description

Draws multiple rounded rectangle.

## Usage

```
motif_plot(
  motif_loc,
  gene_length,
  the_order = NULL,
  motif_select = NULL,
  shape = "RoundRect",
  show_motif_id = FALSE,
  r = 0.3,
  legend_size = 15,
  motif_color = NULL
)
```

## Arguments

<code>motif_loc</code>	A data.frame contains for columuns: ID, motif, start, end.
<code>gene_length</code>	A data.fram of the length of biosequences. Two columns: ID, length.
<code>the_order</code>	A List of Gene ID , One ID Per Line.
<code>motif_select</code>	The motif ID which you want to align with.
<code>shape</code>	RoundRect or Rect.
<code>show_motif_id</code>	Display the name of the motif.
<code>r</code>	The radius of rounded corners.
<code>legend_size</code>	The size of legend.
<code>motif_color</code>	The color set of motif.

## Details

`motif_plot()` draws multiple rounded rectangle to represent the above elements of biosequences, but not limited to biosequences

## Value

`p`

## Author(s)

Shiqi Zhao

## Examples

```
df <- data.frame(
  ID = rep(c("geneA", "geneB", "geneC"), each = 3),
  motif = rep(c("1", "2", "3"), times = 3),
  start = c(1, 3, 6, 1, 6, 10, 10, 7, 17),
  end = c(3, 5, 11, 3, 8, 15, 12, 9, 22)
)

length_data <- data.frame(
  ID = c("geneA", "geneB", "geneC"),
  length = c(15, 27, 30)
)

order_data <- c("geneB", "geneA", "geneC")

motif_plot(df, length_data)
motif_plot(df, length_data, the_order = order_data)
```

**motif\_seq**

*motif\_seq*

## Description

Get motif sequence from meme file or mast file

## Usage

```
motif_seq(meme_file)
```

## Arguments

`meme_file` The path of meme file or mast file.

## Value

`data.frame`

## Author(s)

Shiqi Zhao

## Examples

```
meme_path <- system.file("extdata", "meme.xml", package = "BioVizSeq")
meme_file <- readLines(meme_path)
motifseq<- motif_seq(meme_file)

mast_path <- system.file("extdata", "mast.xml", package = "BioVizSeq")
```

```
mast_file <- readLines(mast_path)
motifseq<- motif_seq(mast_file)
```

**pfam\_plot**

*pfam\_plot*

## Description

Visualization of domain in pfam result file

## Usage

```
pfam_plot(
  pfam_file,
  the_order = NULL,
  domain_select = NULL,
  shape = "RoundRect",
  r = 0.3,
  legend_size = 15,
  domain_color = NULL
)
```

## Arguments

<code>pfam_file</code>	The path of meme file or mast file.
<code>the_order</code>	The path of order file. A List of Gene ID , One ID Per Line.
<code>domain_select</code>	The domain ID which you want to align with.
<code>shape</code>	RoundRect or Rect.
<code>r</code>	The radius of rounded corners.
<code>legend_size</code>	The size of legend.
<code>domain_color</code>	The color set of domain.

## Value

`p`

## Author(s)

Shiqi Zhao

## Examples

```
pfam_path <- system.file("extdata", "iprscan.tsv", package = "BioVizSeq")
order_path <- system.file("extdata", "order.csv", package = "BioVizSeq")
pfam_plot(pfam_path)
pfam_plot(pfam_path, the_order=order_path)
```

---

*pfam\_to\_loc**pfam\_to\_loc*

---

**Description**

Extract the location information of domain from pfam result

**Usage**

```
pfam_to_loc(pfam_data)
```

**Arguments**

pfam\_data      The result file (.tsv) of pfam (via InterPro).

**Value**

list

**Author(s)**

Shiqi Zhao

**Examples**

```
pfam_path <- system.file("extdata", "iprscan.tsv", package = "BioVizSeq")
pfam_file <- read.table(pfam_path, sep='\t', header = FALSE)
motif_loc <- pfam_to_loc(pfam_file)
```

---

---

*plantcare\_classify*      *plantcare\_classify*

---

**Description**

Classify the functions of cis element from Plantcare

**Usage**

```
plantcare_classify(plantcare_file)
```

**Arguments**

plantcare\_file    The result file (.tab) of Plantcare.

**Value**

data.frame

**Author(s)**

Shiqi Zhao

**Examples**

```
plantcare_path <- system.file("extdata", "plantCARE_output.tab", package = "BioVizSeq")
plantcare_file <- read.table(plantcare_path, header = FALSE, sep = '\t', quote="")
plantcare_data <- plantcare_classify(plantcare_file)
```

plantcare_plot	<i>plantcare_plot</i>
----------------	-----------------------

**Description**

Visualization of cis-element in plantcare result file

**Usage**

```
plantcare_plot(
  plantcare_file,
  promoter_length = 2000,
  the_order = NULL,
  shape = "Rect",
  r = 6,
  legend_size = 15,
  element_color = NULL
)
```

**Arguments**

plantcare_file	The path of plantcare result file (.tab).
promoter_length	The promoter length.
the_order	The path of order file. A List of Gene ID , One ID Per Line.
shape	RoundRect or Rect.
r	The radius of rounded corners.
legend_size	The size of legend.
element_color	The color set of cis-element.

**Value**

p

**Author(s)**

Shiqi Zhao

**Examples**

```
plantcare_path <- system.file("extdata", "plantCARE_output.tab", package = "BioVizSeq")
plantcare_plot(plantcare_path, promoter_length = 2000)
```

---

*plantcare\_statistic1* *plantcare\_statistic1*

---

**Description**

Count the number of cis element from Plantcare for heatmap

**Usage**

```
plantcare_statistic1(plantcare_data)
```

**Arguments**

*plantcare\_data* The result of plantcare\_classify().

**Value**

*data.frame*

**Author(s)**

Shiqi Zhao

**Examples**

```
plantcare_path <- system.file("extdata", "plantCARE_output.tab", package = "BioVizSeq")
plantcare_file <- read.table(plantcare_path, header = FALSE, sep = '\t', quote="")
plantcare_data <- plantcare_classify(plantcare_file)
statistic_data1 <- plantcare_statistic1(plantcare_data)
```

---

*plantcare\_statistic2* *plantcare\_statistic2*

---

**Description**

Count the number of cis element from Plantcare for Bar chart

**Usage**

```
plantcare_statistic2(plantcare_data)
```

**Arguments**

`plantcare_data` The result of `plantcare_classify()`.

**Value**

`data.frame`

**Author(s)**

Shiqi Zhao

**Examples**

```
plantcare_path <- system.file("extdata", "plantCARE_output.tab", package = "BioVizSeq")
plantcare_file <- read.table(plantcare_path, header = FALSE, sep = '\t', quote="")
plantcare_data <- plantcare_classify(plantcare_file)
statistic_data2 <- plantcare_statistic2(plantcare_data)
```

---

`plantcare_to_loc`      *plantcare\_to\_loc*

---

**Description**

Extract the location information of cis-element from Plantcare

**Usage**

`plantcare_to_loc(plantcare_data)`

**Arguments**

`plantcare_data` The result of `plantcare_classify()`.

**Value**

`data.frame`

**Author(s)**

Shiqi Zhao

**Examples**

```
plantcare_path <- system.file("extdata", "plantCARE_output.tab", package = "BioVizSeq")
plantcare_file <- read.table(plantcare_path, header = FALSE, sep = '\t', quote="")
plantcare_data <- plantcare_classify(plantcare_file)
plantcare_loc <- plantcare_to_loc(plantcare_data)
```

---

smart\_plot

---

*smart\_plot*

---

## Description

Visualization of domain in SMART result file

## Usage

```
smart_plot(  
  fasta_file,  
  the_order = NULL,  
  domain_select = NULL,  
  shape = "RoundRect",  
  r = 0.3,  
  legend_size = 15,  
  domain_color = NULL  
)
```

## Arguments

fasta_file	The path of protein fasta file.
the_order	The path of order file. A List of Gene ID , One ID Per Line.
domain_select	The domain ID which you want to align with.
shape	RoundRect or Rect.
r	The radius of rounded corners.
legend_size	The size of legend.
domain_color	The color set of domain.

## Value

p

## Author(s)

Shiqi Zhao

---

smart_to_loc	<i>smart_to_loc</i>
--------------	---------------------

---

**Description**

Extract the location information of domain from SMART result

**Usage**

```
smart_to_loc(input_file, do_pfam = TRUE)
```

**Arguments**

input_file	The path of protein fasta file.
do_pfam	Include the pfam domain or not.

**Value**

list

**Author(s)**

Shiqi Zhao

---

upload_fa_to_plantcare	<i>upload_fa_to_plantcare</i>
------------------------	-------------------------------

---

**Description**

Upload the promoter file to Plantcare database

**Usage**

```
upload_fa_to_plantcare(fasta_file, email)
```

**Arguments**

fasta_file	The path of promoter file.
email	e-mail address.

**Details**

Due to the file size limitation of plantcare on fasta, upload\_fa\_to\_plantcare() first splits fasta file. Then uploads the splitted fasta files to the plantcare database, and automatically returns the results to the email provided by the user.

```
upload_fa_to_plantcare("the path/test.fasta", "your e-mail address")
```

**Value**

plantcare\_result

**Author(s)**

Shiqi Zhao

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