

Package ‘g3viz’

March 9, 2021

Type Package

Title Interactively Visualize Genetic Mutation Data using a Lollipop-Diagram

Version 1.1.4

Maintainer Xin Guo <g3viz.group@gmail.com>

Description R interface for 'g3-lollipop' JavaScript library.
Visualize genetic mutation data using an interactive lollipop diagram in RStudio or your browser.

License MIT + file LICENSE

Encoding UTF-8

LazyData true

Depends R (>= 3.5.0)

Imports jsonlite, cgdSr, stringr, htmlwidgets

Suggests shiny (>= 1.0.0), knitr, rmarkdown, kableExtra

URL <https://github.com/G3viz/g3viz>

BugReports <https://github.com/G3viz/g3viz/issues>

RoxygenNote 7.1.1

VignetteBuilder knitr, rmarkdown

NeedsCompilation no

Author Xin Guo [aut, cre]

Repository CRAN

Date/Publication 2021-03-09 09:00:23 UTC

R topics documented:

g3Lollipop	2
g3Lollipop-shiny	3
g3Lollipop.options	4
g3Lollipop.theme	8
getMutationsFromCbiportal	8

guessMAFColumnName	10
hgnc2pfam	10
hgnc2pfam.df	11
hgnc2uniprot	12
mapMutationTypeToMutationClass	13
mutation.table.df	14
parseProteinChange	15
readMAF	16
uniprot2pfam	17

Index	18
--------------	-----------

g3Lollipop	<i>Render g3lollipop diagram for the given mutation data</i>
------------	--

Description

Render g3lollipop diagram for the given mutation data

Usage

```
g3Lollipop(
  mutation.dat,
  gene.symbol,
  uniprot.id = NA,
  gene.symbol.col = "Hugo_Symbol",
  aa.pos.col = "AA_Position",
  protein.change.col = c("Protein_Change", "HGVS_Short"),
  factor.col = "Mutation_Class",
  plot.options = g3Lollipop.options(),
  save.png.btn = TRUE,
  save.svg.btn = TRUE,
  btn.style = NA,
  output.filename = "output"
)
```

Arguments

mutation.dat	Input genomic mutation data frame
gene.symbol	HGNC primary gene symbol
uniprot.id	UniProt ID, in case that the specified gene symbol links to multiple UniProt entries (isoforms). For example, <i>AKAP7</i> gene has two isoforms in UniProt, O43687 and Q9P0M2 .
gene.symbol.col	Column name of Hugo gene symbols (e.g., <i>TP53</i>). Default <i>Hugo_Symbol</i> .
aa.pos.col	Column name of the parsed amino-acid change position. Default <i>AA_Position</i> .

protein.change.col	Column name of protein change information (e.g., p.K960R, G658S, L14Sfs*15). Default is a list of <i>Protein_Change</i> , <i>HGVSp_Short</i> .
factor.col	column of classes in the plot legend. IF NA, use parsed <i>Mutation_Class</i> column, otherwise, use specified. Default NA.
plot.options	g3lollipop diagram options in list format. Check g3Lollipop.options
save.png.btn	If add <i>save-as-png</i> button to the diagram. Default TRUE.
save.svg.btn	If add <i>save-as-svg</i> button to the diagram. Default TRUE.
btn.style	button style, including browser default button style, and two built-in styles, <i>blue</i> or <i>gray</i> . Default NA, indicating browser default.
output.filename	Specify output file name.

Examples

```
# system mutation data
maf.file <- system.file("extdata", "TCGA.BRCA.varscan.somatic.maf.gz", package = "g3viz")
# read in MAF file
mutation.dat <- readMAF(maf.file)

# use built-in chart theme
chart.options <- g3Lollipop.theme(theme.name = "default",
                                  title.text = "PIK3CA gene (default theme)")

# generate chart
g3Lollipop(mutation.dat,
           gene.symbol = "PIK3CA",
           plot.options = chart.options,
           btn.style = "blue",
           output.filename = "default_theme")
```

g3Lollipop-shiny

Shiny bindings for g3Lollipop

Description

Output and render functions for using g3viz lollipop diagram within Shiny applications and interactive Rmd documents.

Usage

```
g3LollipopOutput(outputId, width = "100%", height = "520px")

renderG3Lollipop(expr, env = parent.frame(), quoted = FALSE)
```

Arguments

outputId	output variable to read from
width, height	Must be a valid CSS unit (like '100%', '400px', 'auto') or a number, which will be coerced to a string and have 'px' appended.
expr	An expression that generates a g3-lollipop
env	The environment in which to evaluate expr.
quoted	Is expr a quoted expression (with quote())? This is useful if you want to save an expression in a variable.

g3Lollipop.options *G3Lollipop plot options*

Description

G3Lollipop plot options

Usage

```
g3Lollipop.options(
  chart.width = 800,
  chart.type = "circle",
  chart.margin = list(left = 40, right = 20, top = 15, bottom = 25),
  chart.background = "transparent",
  transition.time = 600,
  y.axis.label = "# of mutations",
  axis.label.font = "normal 12px Arial",
  axis.label.color = "#4f4f4f",
  axis.label.alignment = "middle",
  axis.label.dy = "-2em",
  y.axis.line.color = "#c4c8ca",
  y.axis.line.style = "dash",
  y.axis.line.width = 1,
  y.max.range.ratio = 1.1,
  legend.margin = list(left = 10, right = 0, top = 5, bottom = 5),
  legend.interactive = TRUE,
  legend.title = NA,
  lollipop.track.height = 420,
  lollipop.track.background = "rgb(233,233,233)",
  lollipop.pop.min.size = 2,
  lollipop.pop.max.size = 12,
  lollipop.pop.info.limit = 8,
  lollipop.pop.info.color = "#EEE",
  lollipop.pop.info.dy = "0.35em",
  lollipop.line.color = "rgb(42,42,42)",
  lollipop.line.width = 0.5,
```

```

    lollipop.circle.color = "wheat",
    lollipop.circle.width = 0.5,
    lollipop.label.ratio = 1.4,
    lollipop.label.min.font.size = 10,
    lollipop.color.scheme = "accent",
    highlight.text.angle = "90",
    title.text = "",
    title.font = "normal 16px Arial",
    title.color = "#424242",
    title.alignment = "middle",
    title.dy = "0.35em",
    anno.height = 30,
    anno.margin = list(top = 4, bottom = 0),
    anno.background = "transparent",
    anno.bar.fill = "#e5e3e1",
    anno.bar.margin = list(top = 2, bottom = 2),
    domain.color.scheme = "category10",
    domain.margin = list(top = 0, bottom = 0),
    domain.text.font = "normal 11px Arial",
    domain.text.color = "#f2f2f2",
    brush = TRUE,
    brush.selection.background = "#666",
    brush.selection.opacity = 0.2,
    brush.border.color = "#969696",
    brush.handler.color = "#333",
    brush.border.width = 1,
    legend = TRUE,
    tooltip = TRUE,
    zoom = TRUE
)

```

Arguments

`chart.width` chart width. Default 800.

`chart.type` *pie* or *circle*. Default *circle*.

`chart.margin` specify chart margin in `_list_` format.
Default `list(left = 40, right = 20, top = 15, bottom = 25)`.

`chart.background` chart background. Default *transparent*.

`transition.time` animation transition time when clicking lollipop pops to show labels (in milisecond). Default 600.

`y.axis.label` Y-axis label text. Default *"# of mutations"*.

`axis.label.font` css font style shorthand (*font-style font-variant font-weight font-size/line-height font-family*). Default *"normal 12px Arial"*.

`axis.label.color` axis label text color. Default *#4f4f4f*.

`axis.label.alignment`
 axis label text alignment (start/end/middle). Default *middle*.

`axis.label.dy` text adjustment of axis label text. Default *-2em*.

`y.axis.line.color`
 color of y-axis in-chart lines (ticks). Default *#c4c8ca*.

`y.axis.line.style`
 style of y-axis in-chart lines (ticks), "dash" or "line". Default *dash*.

`y.axis.line.width`
 width of y-axis in-chart lines (ticks). Default 1.

`y.max.range.ratio`
 ratio of y-axis range to data value range. Default 1.1.

`legend.margin` legend margin in *list*. Default `list(left = 10, right = 0, top = 5, bottom = 5)`.

`legend.interactive`
 legend interactive mode. Default TRUE.

`legend.title` legend title. If NA, *factor.col* in `g3Lollipop` is used. Default is NA.

`lollipop.track.height`
 height of lollipop track. Default 420.

`lollipop.track.background`
 background of lollipop track. Default *rgb(244,244,244)*

`lollipop.pop.min.size`
 lollipop pop minimal size. Default 2.

`lollipop.pop.max.size`
 lollipop pop maximal size. Default 12.

`lollipop.pop.info.limit`
 threshold of lollipop pop size to show count information in middle of pop. Default 8.

`lollipop.pop.info.color`
 lollipop pop information text color. Default *#EEE*.

`lollipop.pop.info.dy`
 y-axis direction text adjustment of lollipop pop information. Default *-0.35em*.

`lollipop.line.color`
 lollipop line color. Default *rgb(42,42,42)*.

`lollipop.line.width`
 lollipop line width. Default 0.5.

`lollipop.circle.color`
 lollipop circle border color. Default *wheat*.

`lollipop.circle.width`
 lollipop circle border width. Default 0.5.

`lollipop.label.ratio`
 lollipop click-out label font size to circle size ratio. Default 1.4.

`lollipop.label.min.font.size`
 lollipop click-out label minimal font size. Default 10.

`lollipop.color.scheme`
 color scheme to fill lollipop pops. Default *accent*.

highlight.text.angle	pop-on-click highlight text angle. Default 90.
title.text	title of chart. Default is empty.
title.font	font of chart title. Default <i>normal 16px Arial</i> .
title.color	color of chart title. Default #424242.
title.alignment	text alignment of chart title (start/middle/end). Default <i>middle</i> .
title.dy	text adjustment of chart title. Default <i>0.35em</i> .
anno.height	height of protein structure annotation track. Default 30.
anno.margin	margin of protein structure annotation track. Default <code>list(top = 4, bottom = 0)</code> .
anno.background	background of protein structure annotation track. Default <i>transparent</i> .
anno.bar.fill	background of protein bar in protein structure annotation track. Default #e5e3e1.
anno.bar.margin	margin of protein bar in protein structure annotation track. Default <code>list(top = 2, bottom = 2)</code> .
domain.color.scheme	color scheme of protein domains. Default <i>category10</i> .
domain.margin	margin of protein domains. Default <code>list(top = 0, bottom = 0)</code> .
domain.text.font	domain label text font in shorthand format. Default <i>normal 11px Arial</i> .
domain.text.color	domain label text color. Default #f2f2f2.
brush	if show brush. Default TRUE.
brush.selection.background	background color of selection brush. Default #666.
brush.selection.opacity	background opacity of selection brush. Default 0.2.
brush.border.color	border color of selection brush. Default #969696.
brush.handler.color	color of left and right handlers of selection brush. Default #333.
brush.border.width	border width of selection brush. Default 1.
legend	if show legend. Default TRUE.
tooltip	if show tooltip. Default TRUE.
zoom	if enable zoom feature. Default TRUE.

Value

a list with g3Lollipop plot options

g3Lollipop.theme *G3Lollipop chart options of built-in themes.*

Description

G3Lollipop chart options of built-in themes.

Usage

```
g3Lollipop.theme(
  theme.name = "default",
  title.text = "",
  y.axis.label = "# of mutations",
  legend.title = NA
)
```

Arguments

theme.name	theme name, including <i>default</i> , <i>cbioportal</i> , <i>nature</i> , <i>nature2</i> , <i>dark</i> , <i>blue</i> , <i>ggplot2</i> , and <i>simple</i> . Default <i>default</i> .
title.text	title of chart. Default is empty.
y.axis.label	Y-axis label text. Default <i>"# of mutations"</i> .
legend.title	legend title. If NA, <i>factor.col</i> in <code>g3Lollipop</code> is used. Default is NA.

Value

a list with g3Lollipop plot options

getMutationsFromCbioportal
Query cancer genomic mutation data from cBioPortal

Description

Retrieve and parse mutation data from cBioPortal by the given cBioPortal cancer study ID and the gene symbol.

Usage

```
getMutationsFromCbioportal(
  study.id,
  gene.symbol,
  output.file = NA,
  mutation.type.to.class.df = NA,
  cgds.url = "http://www.cbioportal.org/",
  test.cgds = FALSE
)
```


Arguments

<code>study.id</code>	cbioprotal study ID
<code>gene.symbol</code>	HGNC gene symbol.
<code>output.file</code>	if specified, output to a file in CSV format. Default is NA.
<code>mutation.type.to.class.df</code>	mapping table from mutation type to class. See mapMutationTypeToMutationClass for details. Default NA, which indicates to use default mappings.
<code>cgds.url</code>	the URL for the public CGDS server (Cancer Genomic Data Server). Default is http://www.cbioportal.org/ . Check <i>cgdsr</i> R-package for details.
<code>test.cgds</code>	if test CGDS connection. Default is FALSE

Value

a data frame with columns

Hugo_Symbol Hugo gene symbol

Protein_Change Protein change information (cBioportal uses *HGVS*p format)

Sample_ID Sample ID

Mutation_Type mutation type, aka, variant classification.

Chromosome chromosome

Start_Position start position

End_Position end position

Reference_Allele reference allele

Variant_Allele variant allele

Mutation_Class mutation class (e.g., Truncating/Missense/Inframe/Other)

AA_Position amino-acid position of the variant; if the variant is not in protein-coding region, NA

Examples

```
## Not run:
# Usage:
# Connection to CGDS (Cange Genomic Data Server). Internet access required.
# Note: this may need more than 10 seconds, and sometimes it may fail.
library(cgdsr)
cgds <- CGDS("http://www.cbioportal.org/")

# test if connection is OK (warning: sometimes it may fail)
test(cgds)

# list all studies of cBioPortal
all.studies <- getCancerStudies(cgds)

# First, select a cancer study that contains mutation data set ("caner_study_id")
# then, query genomic mutation data using a HGNC gene symbol,
# for example
mutation.dat <- getMutationsFromCbioportal("msk_impact_2017", "TP53")
```

```
mutation.dat <- getMutationsFromCbioportal("all_stjude_2016", "TP53")
## End(Not run)
```

guessMAFColumnName *Guess column name for MAF file*

Description

Guess column name for MAF file

Usage

```
guessMAFColumnName(maf.df, alt.column.names)
```

Arguments

maf.df MAF data frame
alt.column.names a vector of alternative column names

Value

if hit one alternative column name, return the name; otherwise, return NA

hgnc2pfam *Map from Hugo symbol to Pfam domains*

Description

Mapping from Hugo symbol to Pfam-A domain composition. If the given Hugo symbol has multiple UniProt ID mappings, and guess == TRUE, the longest UniProt protein is selected. Return is either a list of a JSON.

Usage

```
hgnc2pfam(hgnc.symbol, guess = TRUE, uniprot.id = NA, output.format = "json")
```

Arguments

hgnc.symbol primary Hugo symbol
guess if the given Hugo symbol links to multiple UniProt IDs, choose the longest one (guess == TRUE); otherwise NA (guess == FALSE). Default TRUE.
uniprot.id UniProt ID, in case that gene symbol maps to multiple UniProt entries.
output.format output format: JSON or list

Value

A list or a JSON with attributes: *symbol*, *uniprot*, *length*, and a list of *Pfam* entries, including *hmm.acc*, *hmm.name*, *start*, *end*, and *type*.

Examples

```
# general usage
hgnc2pfam("TP53")
hgnc2pfam("TP53", output.format = "json")
hgnc2pfam("TP53", output.format = "list")
hgnc2pfam("TP53", output.format = "json", uniprot.id = "P04637") # OK

# for gene mapping to multiple UniProt enties
hgnc2pfam("GNAS", guess = TRUE)
hgnc2pfam("GNAS", guess = FALSE)
hgnc2pfam("GNAS", output.format = "list")
hgnc2pfam("GNAS", output.format = "list", uniprot.id = "P84996")
## Not run:
hgnc2pfam("GNAS", output.format = "list", uniprot.id = "P84997") # not exists, returns FALSE

## End(Not run)

hgnc2pfam("PRAMEF9", output.format = "list") # no Pfam mappings
```

hgnc2pfam.df

*Mapping table between gene.symbol, uniprot.id, and pfam***Description**

A dataset containing the mapping table between Hugo symbol, UniProt ID, and Pfam ACC.

Usage

```
hgnc2pfam.df
```

Format

A data frame with columns:

symbol Gene symbol

uniprot UniProt ID

length protein length

start starting position of Pfam domain

end ending position of Pfam domain

hmm.acc Pfam accession number

hmm.name Pfam name

type Pfam type, i.e., domain/family/motif/repeat/disordered/coiled-coil

Source

Pfam (v31.0) and UniProt

Examples

```
hgnc2pfam.df
```

hgnc2uniprot

Mapping from Hugo symbol to UniProt IDs

Description

Mapping from Hugo Symbol to UniProt ID using internal mapping table. Return a data frame with columns *symbol* (Hugo symbol), *uniprot* (UniProt ID), and *length* (protein length).

Usage

```
hgnc2uniprot(hgnc.symbol)
```

Arguments

hgnc.symbol primary HUGO symbol

Value

a data frame with columns *symbol* (Hugo symbol), *uniprot* (UniProt ID), and *length* (protein length).

Examples

```
# maps to single UniProt entry
hgnc2uniprot("TP53")

# maps to multiple UniProt entries
hgnc2uniprot("GNAS")
hgnc2uniprot("AKAP7")
```

`mapMutationTypeToMutationClass`*Map from mutation type (aka, variant classification) to mutation class*

Description

Map from mutation type (aka, variant classification) to mutation class. Default mappings are as follows,

- Missense
 - *Missense_Mutation* — a point mutation in which a single nucleotide change results in a codon that codes for a different amino acid See https://en.wikipedia.org/wiki/Missense_mutation.
- Inframe
 - *In_Frame_Del* — a deletion that keeps the sequence in frame
 - *In_Frame_Ins* — an insertion that keeps the sequence in frame
 - *Silent* — variant is in coding region of the chosen transcript, but protein structure is identical (i.e., a synonymous mutation)
 - *Targeted_Region* — targeted region
- Truncating
 - *Frame_Shift* — a variant caused by indels of a number of nucleotides in a DNA sequence that is not divisible by three. See https://en.wikipedia.org/wiki/Frameshift_mutation.
 - *Frame_Shift_Ins* — a variant caused by insertion that moves the coding sequence out of frame. See https://en.wikipedia.org/wiki/Frameshift_mutation.
 - *Frame_Shift_Del* — a variant caused by deletion that moves the coding sequence out of frame. See https://en.wikipedia.org/wiki/Frameshift_mutation.
 - *Nonsense_Mutation* — a premature stop codon that is created by the variant. See https://en.wikipedia.org/wiki/Nonsense_mutation.
 - *Nonstop_Mutation* — a variant that removes stop codon.
 - *Splice_Site* — a variant that is within two bases of a splice site.
 - *Splice_Region* — a variant that is within splice region.
- Other
 - *5'UTR* — a variant that is on the 5'UTR for the chosen transcript.
 - *3'UTR* — a variant that is on the 3'UTR for the chosen transcript.
 - *5'Flank* — a variant that is upstream of the chosen transcript (generally within 3kb).
 - *3'Flank* — a variant that is downstream of the chosen transcript (generally within 3kb).
 - *Fusion* — a gene fusion.
 - *IGR* — an intergenic region. Does not overlap any transcript.
 - *Intron* — a variant that lies between exons within the bounds of the chosen transcript.
 - *Translation_Start_Site* — a variant that is in translation start site.

- *De_novo_Start_InFrame* — a novel start codon that is created by the given variant using the chosen transcript. However, it is in frame relative to the coded protein.
- *De_novo_Start_OutOfFrame* — a novel start codon that is created by the given variant using the chosen transcript. However, it is out of frame relative to the coded protein.
- *Start_Codon_SNP* — a point mutation that overlaps the start codon.
- *Start_Codon_Ins* — an insertion that overlaps the start codon.
- *Start_Codon_Del* — a deletion that overlaps the start codon.
- *RNA* — a variant that lies on one of the RNA transcripts.
- *lincRNA* — a variant that lies on one of the lincRNAs.
- *Unknown* — Unknown

Usage

```
mapMutationTypeToMutationClass(
  mutation.type.vec,
  mutation.type.to.class.df = NA
)
```

Arguments

`mutation.type.vec`
a vector of mutation type information

`mutation.type.to.class.df`
A mapping table from mutation type (header *Mutation_Type*) to mutation class (header *Mutation_Class*). Default NA, which indicates to use default mappings.

Value

a vector of mapped mutation class information

<code>mutation.table.df</code>	<i>Default mapping table between mutation type (aka, variant classification) to mutation class</i>
--------------------------------	--

Description

A dataset containing the mapping table between genomic mutation type (aka, variant classification) to mutation class. See [mapMutationTypeToMutationClass](#) for details.

Usage

```
mutation.table.df
```

Format

A data frame with three columns:

Mutation_Type Mutation type, aka, variant classification

Mutation_Class mutation class

Short_Name short name of mutation type

Examples

```
mutation.table.df
```

parseProteinChange	<i>Extract amino_acid_position from Protein_Change</i>
--------------------	--

Description

Parse *amino_acid_position* according to HGVS_{short} format.

For example, *p.Q16Rfs*28*, amino-acid position is 16. See <http://varnomen.hgvs.org/recommendations/protein/> or <https://www.hgvs.org/mutnomen/recs-prot.html>.

Usage

```
parseProteinChange(protein.change.vec, mutation.class.vec)
```

Arguments

`protein.change.vec`

a vector of strings with protein change information, usually in HGVS_{short} format.

`mutation.class.vec`

a vector of strings with mutation class (or so-called variant classification) information.

Value

a vector of parsed amino-acid position

readMAF	<i>Read MAF file</i>
---------	----------------------

Description

Read mutation information from MAF file. For MAF format specification, see https://docs.gdc.cancer.gov/Data/File_Formats/MAF_Format/.

Usage

```
readMAF(
  maf.file,
  gene.symbol.col = "Hugo_Symbol",
  variant.class.col = c("Variant_Classification", "Mutation_Type"),
  protein.change.col = c("Protein_Change", "HGVS_Short"),
  if.parse.aa.pos = TRUE,
  if.parse.mutation.class = TRUE,
  mutation.class.col = "Mutation_Class",
  aa.pos.col = "AA_Position",
  mutation.type.to.class.df = NA,
  sep = "\t",
  quote = "",
  ...
)
```

Arguments

<code>maf.file</code>	MAF file name. Gzipped input file allowed, with ".gz" file extension.
<code>gene.symbol.col</code>	Column name of Hugo gene symbols (e.g., TP53). Default <i>Hugo_Symbol</i> .
<code>variant.class.col</code>	Column name for variant class information (e.g., <i>Missense_Mutation</i> , <i>Nonsense_Mutation</i>). Default is the first match of <i>Variant_Classification</i> or <i>Mutation_Type</i> .
<code>protein.change.col</code>	Column name for protein change information (e.g., p.K960R, G658S, L14Sfs*15). Default is the first match of <i>Protein_Change</i> or <i>HGVS_Short</i> .
<code>if.parse.aa.pos</code>	if parse amino-acid position of mutations. Default is TRUE.
<code>if.parse.mutation.class</code>	if parse mutation class from mutation type (variant classification) information. Default is TRUE.
<code>mutation.class.col</code>	Column name of the parsed mutation class. Default <i>Mutation_Class</i> .
<code>aa.pos.col</code>	Column name of the parsed amino-acid change position. Default <i>AA_Position</i> .


```

mutation.type.to.class.df
    mapping table from mutation type to class. mapMutationTypeToMutationClass
    for details. Default NA, which indicates to use default mappings.
sep
    separator of columns. Default sep = "\t".
quote
    the set of quoting characters. To disable quoting altogether, use quote = "".
    Default quote = "".
...
    additional parameters pass to read.table.

```

Value

a data frame containing MAF information, plus optional columns of the parsed *Mutation_Class* and *Protein_Position*.

```
uniprot2pfam      From UniProt ID to Pfam-A domain composition
```

Description

Map from UniProt ID to Pfam-A domain composition.

Usage

```
uniprot2pfam(uniprot.id)
```

Arguments

```
uniprot.id      UniProt ID
```

Value

a data frame with columns

- *uniprot* — UniProt ID
- *length* — protein length
- *hmm.acc* — accession number of Pfam HMM model, e.g., PF08563
- *hmm.name* — Pfam name, e.g., P53_TAD
- *start* — Pfam domain start position
- *end* — Pfam domain end position
- *type* — Pfam type, including domain/motif/family

Examples

```
uniprot2pfam("Q5VWM5") # PRAMEF9; PRAMEF15
uniprot2pfam("P04637")
```

Index

* datasets

hgnc2pfam.df, [11](#)
mutation.table.df, [14](#)

g3Lollipop, [2](#), [6](#), [8](#)
g3Lollipop-shiny, [3](#)
g3Lollipop.options, [3](#), [4](#)
g3Lollipop.theme, [8](#)
g3LollipopOutput (g3Lollipop-shiny), [3](#)
getMutationsFromCbioportal, [8](#)
guessMAFColumnName, [10](#)

hgnc2pfam, [10](#)
hgnc2pfam.df, [11](#)
hgnc2uniprot, [12](#)

mapMutationTypeToMutationClass, [9](#), [13](#),
[14](#), [17](#)
mutation.table.df, [14](#)

parseProteinChange, [15](#)

read.table, [17](#)
readMAF, [16](#)
renderG3Lollipop (g3Lollipop-shiny), [3](#)
uniprot2pfam, [17](#)