# Package 'rTCRBCRr'

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Type Package

Title Repertoire Analysis of the Detected Clonotype

Version 0.1.1
Description The 'TRUST4' or 'MiXCR' is used to identify the clonotypes.  The goal of 'rTCRBCRr' is to process the results from these clonotyping tools, and analyze the clonotype repertoire metrics based on chain names and IGH isotypes. The manuscript is still under preparation for publication for now.  The references describing the methods in this package will be added later.
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Author Lizhong Ding [aut, cre]
Maintainer Lizhong Ding <z82xh3f4r@mozmail.com></z82xh3f4r@mozmail.com>
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all\_sample\_all\_chain\_all\_metrics\_wide\_format\_dataframe a dataframe of wide format

# Description

The data frame must be generated from combine\_all\_sample\_repertoire\_metrics function

# Usage

 $\verb|all_sample_all_chain_all_metrics_wide_format_data frame|\\$ 

# **Format**

A data frame with m rows and n variables:

#### **Source**

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

 $all\_sample\_all\_chain\_all\_metrics\_wide\_format\_dataframe\_list\\ a \ dataframe \ list \ to \ combine \ rowwise.$ 

# **Description**

The element data frame must be generated from compute\_repertoire\_metrics\_by\_chain\_name function

# Usage

```
all_sample_all_chain_all_metrics_wide_format_dataframe_list
```

#### **Format**

A list of data frames with m rows and n variables:

#### **Source**

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

```
all\_sample\_all\_chain\_individual\_metrics\_dataframe\_list a\ \textit{list of dataframes}
```

# **Description**

The list of data frames generated from get\_item\_name\_x\_sample\_name\_for\_each\_metric

#### **Usage**

```
\verb|all_sample_all_chain_individual_metrics_data frame_list|
```

#### **Format**

A list of data frame with m rows and n variables:

#### **Source**

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

all\_sample\_IGH\_chain\_all\_metrics\_wide\_format\_dataframe a dataframe of wide format

# **Description**

The data frame must be generated from combine\_all\_sample\_repertoire\_metrics function

# Usage

```
all_sample_IGH_chain_all_metrics_wide_format_dataframe
```

#### **Format**

A data frame with m rows and n variables:

#### Source

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

```
all_sample_IGH_chain_all_metrics_wide_format_dataframe_list 
 a dataframe list to combine rowwise.
```

# **Description**

The element data frame must be generated from calculate\_IGH\_isotype\_proportion function

# Usage

```
\verb|all_sample_IGH_chain_all_metrics_wide_format_data frame_list|
```

#### **Format**

A list of data frames with m rows and n variables:

#### **Source**

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

all\_sample\_IGH\_chain\_individual\_metrics\_dataframe\_list  $a\ list\ of\ dataframes$ 

#### **Description**

The list of data frames generated from get\_item\_name\_x\_sample\_name\_for\_each\_metric

# Usage

```
all\_sample\_IGH\_chain\_individual\_metrics\_dataframe\_list
```

#### **Format**

A list of data frame with m rows and n variables:

#### **Source**

https://www.rdocumentation.org/packages/base/versions/3.6.2/topics/list

annotate\_chain\_name\_and\_isotype\_name

Annotate chain name and isotype name

#### **Description**

Annotate chain name and isotype name

# Usage

```
annotate_chain_name_and_isotype_name(input_dataframe)
```

#### **Arguments**

input\_dataframe

A dataframe of clonotype of immunarch format

#### Value

A dataframe of clonotype of immunarch format

#### **Examples**

annotate\_chain\_name\_and\_isotype\_name(nonproductive\_CDR3aa\_removed\_dataframe)

```
calculate\_IGH\_isotype\_proportion \\ calculate\_IGH\_isotype\_proportion
```

# **Description**

```
calculate_IGH_isotype_proportion
```

# Usage

```
calculate_IGH_isotype_proportion(input_dataframe)
```

# **Arguments**

input\_dataframe

a clonotype dataframe with chain name and isotype name information

#### Value

a dataframe of count and proportion of isotypes of IGH chain.

# **Examples**

calculate\_IGH\_isotype\_proportion(convergent\_clonotype\_merged\_dataframe)

# Description

```
calculate_repertoire_metrics
```

# Usage

```
calculate_repertoire_metrics(named_species_vector)
```

# Arguments

```
named_species_vector
a named vector
```

#### Value

a named vector

# **Examples**

```
a_named_vector <- c("apple" = 1, "banana" = 3, "cranberry" = 7, "date" = 10)
calculate_repertoire_metrics(a_named_vector)</pre>
```

```
calibrate\_proportion\_by\_clones \\ calibrate\_proportion\_by\_clones
```

# Description

calibrate\_proportion\_by\_clones

#### **Usage**

calibrate\_proportion\_by\_clones(a\_dataframe)

# **Arguments**

```
a_dataframe a dataframe
```

#### Value

a dataframe

```
chain_separated_dataframe
```

a dataframe separated by chain name

# Description

an example dataframe separated by chain name

# Usage

```
chain_separated_dataframe
```

#### **Format**

A data frame with m rows and n variables:

#### Source

```
https://en.wikipedia.org/wiki/T-cell_receptor
```

# **Description**

```
combine_all_sample_repertoire_metrics
```

#### Usage

```
combine_all_sample_repertoire_metrics(input_dataframe_list)
```

#### **Arguments**

# Value

a wide table with all samples' all chains' all repertoire metrics

# **Examples**

```
combine_all_sample_repertoire_metrics(all_sample_all_chain_all_metrics_wide_format_dataframe_list)
combine_all_sample_repertoire_metrics(all_sample_IGH_chain_all_metrics_wide_format_dataframe_list)
```

# Description

```
compute_repertoire_metrics_by_chain_name
```

#### **Usage**

```
\verb|compute_repertoire_metrics_by_chain_name(input_dataframe)|\\
```

# **Arguments**

```
input_dataframe
```

a clonotype dataframe with chain name information

# Value

repertoire metrics by chain name.

#### **Examples**

```
compute_repertoire_metrics_by_chain_name(convergent_clonotype_merged_dataframe)
```

```
convergent_clonotype_merged_dataframe

a dataframe where convergent clonotypes are merged
```

# **Description**

an example dataframe where convergent clonotypes are merged

#### Usage

```
convergent_clonotype_merged_dataframe
```

#### **Format**

A data frame with m rows and n variables:

#### Source

```
https://www.frontiersin.org/articles/10.3389/fimmu.2019.02985/full
```

```
format_clonotype_to_immunarch_style

*Convert clonotype dataframe to immunarch format*
```

# **Description**

Convert clonotype dataframe to immunarch format

# Usage

```
format_clonotype_to_immunarch_style(input_dataframe, clonotyping_tool)
```

# **Arguments**

# Value

a clonotype dataframe of immunarch format

#### **Examples**

```
format_clonotype_to_immunarch_style(raw_input_clonotype_dataframe, "trust")
```

# **Description**

```
get_item_name_x_sample_name_for_each_metric
```

#### **Usage**

```
get_item_name_x_sample_name_for_each_metric(input_dataframe)
```

#### **Arguments**

input\_dataframe

a dataframe with repertoire metrics calculated for each item name

#### Value

a list of dataframes that is one single repertoire metric for all items in all samples

#### **Examples**

```
{\tt get\_item\_name\_x\_sample\_name\_for\_each\_metric(all\_sample\_all\_chain\_all\_metrics\_wide\_format\_dataframe)}
```

immunarch\_style\_dataframe

An converted dataframe to immunarch format

# Description

An example dataframe converted to immunarch format

#### **Usage**

```
immunarch_style_dataframe
```

#### **Format**

A data frame with m rows and n variables:

#### **Source**

https://github.com/immunomind/immunarch/

merge\_convergent\_clonotype

Merge convergent clonotype

# **Description**

Merge convergent clonotype

# Usage

merge\_convergent\_clonotype(input\_dataframe)

# Arguments

input\_dataframe

A dataframe of clonotype of immunarch format

#### Value

A dataframe of clonotype of immunarch format merge\_convergent\_clonotype\_function(an\_immunarch\_format\_clonotype\_d

#### **Examples**

merge\_convergent\_clonotype(chain\_separated\_dataframe)

 $nonproductive\_CDR3aa\_removed\_dataframe \\ a \ dataframe \ with \ nonproductive \ CDR3aa \ removed$ 

# Description

an example dataframe with nonproductive CDR3aa removed

# Usage

nonproductive\_CDR3aa\_removed\_dataframe

#### **Format**

A data frame with m rows and n variables:

#### **Source**

https://mixcr.readthedocs.io/en/master/analyze.html

parse\_trust4

parse\_mixcr

Parse mixer format

# Description

Parse mixer format

# Usage

```
parse_mixcr(input_clone_dataframe)
```

# **Arguments**

input\_clone\_dataframe

Generated from mixer program

# Value

A dataframe of immunarch format

parse\_trust4

Parse trust4 format

# Description

Parse trust4 format

# Usage

```
parse_trust4(input_clone_dataframe)
```

# Arguments

```
input_clone_dataframe
```

Generated from trust4 program

# Value

A dataframe of immunarch format

raw\_input\_clonotype\_dataframe

Raw clonotype output from trust4

# **Description**

An example dataframe for raw clonotype identified by trust4

# Usage

```
raw_input_clonotype_dataframe
```

# **Format**

A data frame with m rows and n variables:

#### Source

```
https://github.com/liulab-dfci/TRUST4
```

```
remove_nonproductive_CDR3aa
```

Clean the clonotype dataframe

# Description

Clean the clonotype dataframe

# Usage

```
remove_nonproductive_CDR3aa(input_dataframe)
```

# **Arguments**

input\_dataframe

A dataframe of clonotype of immunarch format

#### Value

A dataframe of clonotype of immunarch format

# **Examples**

```
remove_nonproductive_CDR3aa(immunarch_style_dataframe)
```

```
summarize\_convergence\_function \\ summarize\_convergence\_function
```

# Description

```
summarize_convergence_function
```

# Usage

```
summarize_convergence_function(a_dataframe)
```

# **Arguments**

```
a_dataframe a dataframe
```

# Value

a dataframe

```
summarize_repertoire_metrics
summarize_repertoire_metrics
```

# Description

```
summarize_repertoire_metrics
```

# Usage

```
summarize_repertoire_metrics(one_sample_dataframe)
```

# **Arguments**

```
\label{eq:cone_sample_data} one\_sample\_data frame \\ a \ data frame \ of \ clonotype \ in \ immunarch \ format
```

# Value

a named vector

the global variables 15

the global variables Make variable names global to avoid CHECK notes

# Description

Global variables

 $the\_divergent\_clonotype\_dataframe\_list\\ a \ \textit{list of dataframe where convergent clonotypes are merged}$ 

# **Description**

an list of example dataframes where convergent clonotypes are merged

# Usage

 $the\_divergent\_clonotype\_dataframe\_list$ 

#### **Format**

A list of data frame with m rows and n variables:

#### **Source**

https://www.frontiersin.org/articles/10.3389/fimmu.2019.02985/full

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